DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

42 CFR Parts 405, 412, 413, 414, 416, 486, 488, 489, and 495

[CMS–1677–P]

RIN 0938–AS98

Medicare Program; Hospital Inpatient Prospective Payment Systems for Acute Care Hospitals and the Long-Term Care Hospital Prospective Payment System and Proposed Policy Changes and Fiscal Year 2018 Rates; Quality Reporting Requirements for Specific Providers; Medicare and Medicaid Electronic Health Record (EHR) Incentive Program Requirements for Eligible Hospitals, Critical Access Hospitals, and Eligible Professionals; Provider-Based Status of Indian Health Service and Tribal Facilities and Organizations; Costs Reporting and Provider Requirements; Agreement Termination Notices

AGENCY: Centers for Medicare and Medicaid Services (CMS), HHS.

ACTION: Proposed rule.

SUMMARY: We are proposing to revise the Medicare hospital inpatient prospective payment systems (IPPS) for operating and capital-related costs of acute care hospitals to implement changes arising from our continuing experience with these systems for FY 2018. Some of these proposed changes would implement certain statutory provisions contained in the Pathway for Sustainable Growth Rate (SGR) Reform Act of 2013, the Improving Medicare Post-Acute Care Transformation Act of 2014, the Medicare Access and CHIP Reauthorization Act of 2015, the 21st Century Cures Act, and other legislation. We also are making proposals relating to the provider-based status of Indian Health Service (IHS) and Tribal facilities and organizations and to the low-volume hospital payment adjustment for hospitals operated by the IHS or a Tribe. In addition, we are providing the proposed estimated market basket update that would apply to the rate-of-increase limits for certain hospitals excluded from the IPPS that are paid on a reasonable cost basis subject to these limits for FY 2018. We are proposing to update the payment policies and the annual payment rates for the Medicare prospective payment system (PPS) for inpatient hospital services provided by long-term care hospitals (LTCHs) for FY 2018.

In addition, we are proposing to establish new requirements or revise existing requirements for quality reporting by specific Medicare providers (acute care hospitals, PPS-exempt cancer hospitals, LTCHs, and inpatient psychiatric facilities). We also are proposing to establish new requirements or revise existing requirements for eligible professionals (EPs), eligible hospitals, and critical access hospitals (CAHs) participating in the Medicare and Medicaid Electronic Health Record (EHR) Incentive Programs. We are proposing to update policies relating to the Hospital Value-Based Purchasing (VBP) Program, the Hospital Readmissions Reduction Program, and the Hospital-Acquired Condition (HAC) Reduction Program.

We also are proposing changes relating to transparency of accrediting organization survey reports and plans of correction of providers and suppliers; electronic signature and electronic submission of the Certification and Settlement Summary page of the Medicare cost reports; and clarification of provider disposal of assets.

DATES: Comment Period: To be assured consideration, comments must be received at one of the addresses provided in the ADDRESS section, no later than 5 p.m. EDT on June 13, 2017.

ADDRESSES: In commenting, please refer to file code CMS–1677–P. Because of staff and resource limitations, we cannot accept comments by facsimile (FAX) transmission.

You may submit comments in one of four ways (no duplicates, please):

1. Electronically. You may (and we encourage you to) submit electronic comments on this regulation to http://www.regulations.gov. Follow the instructions under the “submit a comment” tab.

2. By regular mail. You may mail written comments to the following address ONLY: Centers for Medicare & Medicaid Services, Department of Health and Human Services, Attention: CMS–1677–P, P.O. Box 8011, Baltimore, MD 21244–1850.

   Please allow sufficient time for mailed comments to be received before the close of the comment period.

3. By express or overnight mail. You may submit comments via express or overnight mail to the following address ONLY: Centers for Medicare & Medicaid Services, Department of Health and Human Services, Attention: CMS–1677–P, Mail Stop C4–26–05, 7500 Security Boulevard, Baltimore, MD 21244–1850.

4. By hand or courier. If you prefer, you may deliver (by hand or courier) your written comments before the close of the comment period to either of the following addresses:

   a. For delivery in Washington, DC—Centers for Medicare & Medicaid Services, Department of Health and Human Services, Room 445–G, Hubert H. Humphrey Building, 200 Independence Avenue SW., Washington, DC 20201. (Because access to the interior of the Hubert H. Humphrey Building is not readily available to persons without Federal Government identification, commenters are encouraged to leave their comments in the CMS drop slots located in the main lobby of the building. A stamp-in clock is available for persons wishing to retain a proof of filing by stamping in and retaining an extra copy of the comments being filed.)

   b. For delivery in Baltimore, MD—Centers for Medicare & Medicaid Services, Department of Health and Human Services, 7500 Security Boulevard, Baltimore, MD 21244–1850.

If you intend to deliver your comments to the Baltimore address, please call the telephone number (410) 786–7195 in advance to schedule your arrival with one of our staff members.

Comments mailed to the addresses indicated as appropriate for hand or courier delivery may be delayed and received after the comment period.

For information on viewing public comments, we refer readers to the beginning of the SUPPLEMENTARY INFORMATION section.

FOR FURTHER INFORMATION CONTACT: Donald Thompson, (410) 786–4487, and Michele Hudson, (410) 786–4487, Operating Prospective Payment, MS–DRGs, Wage Index, New Medical Service and Technology Add-On Payments, Hospital Geographic Reclassifications, Graduate Medical Education, Capital Prospective Payment, Excluded Hospitals, Sole Community Hospitals, Medicare Disproportionate Share Hospital (DSH) Payment Adjustment, Medicare-Dependent Small Rural Hospital (MDH) Program, and Low-Volume Hospital Payment Adjustment Issues.

Michele Hudson, (410) 786–4487, Mark Luxton, (410) 786–4530, and Emily Lipkin, (410) 786–3633, Long-Term Care Hospital Prospective Payment System and MS–LTC–DRG Relative Weights Issues.

Mollie Knight, (410) 786–7948, and Bridget Dickensheets, (410) 786–8670, Rebasing and Revising the Hospital Market Basket Issues.

Siddhartha Maity, (410) 786–6673, Rural Community Hospital Demonstration Program Issues.
Supplementary Information:

Inspection of Public Comments: All comments received before the close of the comment period are available for viewing by the public, including any personally identifiable or confidential business information that is included in a comment. We post all comments received before the close of the comment period on the following Web site as soon as possible after they have been received: http://www.regulations.gov. Follow the search instructions on that Web site to view public comments.

Comments received timely will also be available for public inspection, generally beginning approximately 3 weeks after publication of the rule, at the headquarters of the Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Baltimore, MD 21244, on Monday through Friday of each week from 8:30 a.m. to 4:00 p.m. EST. To schedule an appointment to view public comments, phone 1-800-743-3951.

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In the past, a majority of the tables referred to throughout this preamble and in the Addendum to the proposed rule and the final rule were published in the Federal Register as part of the annual proposed and final rules. However, beginning in FY 2012, some of the IPPS tables and LTCH PPS tables are no longer published in the Federal Register. Instead, these tables generally will be available only through the Internet. The IPPS tables for this proposed rule are available through the Internet on the CMS Web site at: http://www.cms.hhs.gov/Medicare-Medicaid-Fee-for-Service-Payment/AcuteInpatientPPS/index.html. Click on the link on the left side of the screen titled, “FY 2018 IPPS Proposed Rule Home Page” and “Acute Inpatient—Files for Download”. The LTCH PPS tables for this FY 2018 proposed rule are available through the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/LongTermCareHospitalPPS/index.html under the list item for Regulation Number CMS-1677-P. For further details on the contents of the tables referenced in this proposed rule, we refer readers to section VI. of the Addendum to this proposed rule.

Readers who experience any problems accessing any of the tables that are posted on the CMS Web sites identified above should contact Michael Treitel at (410) 786-4552.

Acronyms

3M 3M Health Information System
AAMC Association of American Medical Colleges
ACGME Accreditation Council for Graduate Medical Education
ACoS American College of Surgeons
AHA American Hospital Association
AHIC American Health Information Community
AIHA American Health Information Association
AMCA American Medical Association
AMGA American Medical Group Association
AMI Acute myocardial infarction
AO Accrediting Organizations
AOA American Osteopathic Association
APDRG All Patient Refined Diagnosis Related Group System
APRN Advanced practice registered nurse
ASITN American Society of Interventional and Therapeutic Neuroradiology
ASPE Assistant Secretary for Planning and Evaluation (DEHIS)
ATRA American Taxpayer Relief Act of 2012, Public Law 112–240
BBRA Medicare, Medicaid, and SCHIP [State Children’s Health Insurance Program] Balanced Budget Refinement Act of 1999, Public Law 106–113
BLS Bureau of Labor Statistics
CABG Coronary artery bypass graft [surgery]
CAH Critical access hospital
CARE [Medicare] Continuity Assessment and Record Evaluation [Instrument]
CART CMS Abstraction & Reporting Tool
CAUTI Gathered-associated urinary tract infection
CBSAs Core-based statistical areas
CC Complication or comorbidity
CCN CMS Certification Number
CCR Cost-to-charge ratio
CDAC [Medicare] Clinical Data Abstraction Center
CDAD Clostridium difficile-associated disease
CDC Centers for Disease Control and Prevention

Jeri Smith, (410) 786–0110, Frontier Community Health Integration Project Demonstration Issues.
Lein Han, (617) 879–0129, Hospital Readmissions Reduction Program—Readmission Measures for Hospitals Issues.
Delia Houseal, (410) 786–2724, Hospital Readmissions Reduction Program—Administration Issues.
Elizabeth Bainger, (410) 786–0529, Hospital Acquired Condition Reduction Program Issues.
Graco Im, (410) 786–0700 and James Poyer, (410) 786–2261, Hospital Inpatient Quality Reporting and Hospital Value-Based Purchasing—Program Administration, Validation, and Reconsideration Issues.
Kim Spaulding Bush, (410) 786–3232, Hospital Value-Based Purchasing—Efficiency Measures Issues.
Elizabeth Goldstein, (410) 786–6665, Hospital Inpatient Quality Reporting—Hospital Consumer Assessment of Healthcare Providers and Systems Measures Issues.
James Poyer, (410) 786–2261, PPS-Exempt Cancer Hospital Quality Reporting Issues.
Mary Pratt, (410) 786–6867, Long-Term Care Hospital Quality Data Reporting Issues.
Kathleen Johnson, (410) 786–3295 and Steven Johnson (410) 786–3332, EHR Incentive Program Nonclinical Quality Measure Related Issues.
PSF  Provider-Specific File
PSI  Patient safety indicator
PS&R  Provider Statistical and Reimbursement [System]
PQRS  Physician Quality Reporting System
PUF  Public use file
QDM  Quality data model
QES ASAP Quality Improvement Evaluation System Assessment Submission and Processing
QIG  Quality Improvement Group [CMS]
QIO  Quality Improvement Organization
QM  Quality measure
QPP  Quality Payment Program
QRDA  Quality Reporting Document Architecture
RFA  Regulatory Flexibility Act, Public Law 96–254
RHC  Rural health clinic
RHQDAPU  Reporting hospital quality data for annual payment update
RIM  Reference information model
RNHCI  Religious nonmedical health care institution
RPL  Rehabilitation psychiatric long-term care hospital
RRC  Rural referral center
RSR  Risk-standard mortality rate
RSP  Risk-standardized payment
RSDR  Risk-standard readmission rate
RTI  Research Triangle Institute, International
RUCA  Rural-urban commuting area codes
RY  Rate year
SAF  Standard Analytic File
SCA  Sole community hospital
SCHI  State Child Health Insurance Program
SCIP  Surgical Care Improvement Project
SFY  State fiscal year
SGR  Sustainable Growth Rate
SIC  Standard Industrial Classification
SIR  Standardized infection ratio
SNF  Skilled nursing facility
SNF-QRP  Skilled Nursing Facility Quality Reporting Program
SNF-VBP  Skilled Nursing Facility Value-Based Purchasing
SOCs  Standard occupational classifications
SOM  State Operations Manual
SRR  Standardized risk ratio
SSI  Supplemental Security Income
SRO  Short-stay outlier
SUD  Substance use disorder
TEP  Technical expert panel
THA/TKA  Total hip arthroplasty/total knee arthroplasty
TMA TMA [Transitional Medical Assistance], Abstinence Education, and QI [Qualifying Individuals] Programs
Extension Act of 2007, Public Law 110–90
TPS  Total Performance Score
UHDDS Uniform hospital discharge data set
UA  Utilization data set
VBP  [Hospital] Value Based Purchasing [Program]
VTE  Venous thromboembolism

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Under various statutory authorities, we are proposing to make changes to the Medicare IPPS, to the LTCH PPS, and to other related payment methodologies and programs for FY 2018 and subsequent fiscal years. These statutory authorities include, but are not limited to, the following:

- Section 1806(d) of the Social Security Act (the Act), which sets forth a system of payment for the operating costs of acute care hospital inpatient stays under Medicare Part A (Hospital Insurance) based on prospectively set rates. Section 1886(g) of the Act requires that, instead of paying for capital-related costs of inpatient hospital services on a reasonable cost basis, the Secretary use a prospective payment system (PPS).
  - Section 1886(d)(1)(B) of the Act, which specifies that certain hospitals and hospital units are excluded from the IPPS. These hospitals and units are: Rehabilitation hospitals and units; LTCHs; psychiatric hospitals and units; children’s hospitals; cancer hospitals; long-term care neoplastic disease hospitals, and hospitals located outside the 50 States, the District of Columbia, and Puerto Rico (that is, hospitals located in the U.S. Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa). Religious nonmedical health care institutions (RNHCIs) are also excluded from the IPPS.
  - Sections 123(a) and (c) of the BBRA (Pub. L. 106–113) and section 307(b)(1) of the BIPA (Pub. L. 106–554) (as codified under section 1886(m)(1) of the Act), which provide for the development and implementation of a prospective payment system for payment for inpatient hospital services of long-term care hospitals (LTCHs) described in section 1886(d)(1)(B)(iv) of the Act.
  - Sections 1814(l), 1820, and 1834(g) of the Act, which specify that payments are made to critical access hospitals (CAHs) (that is, rural hospitals or facilities that meet certain statutory requirements) for inpatient and outpatient services and that these payments are generally based on 101 percent of reasonable cost.
  - Section 1866(k) of the Act, as added by section 3005 of the Affordable Care Act, which establishes a quality reporting program for hospitals described in section 1886(d)(1)(B)(v) of the Act, referred to as “PPS-exempt cancer hospitals.”
  - Section 1886(a)(4) of the Act, which specifies that costs of approved
educational activities are excluded from the operating costs of inpatient hospital services. Hospitals with approved graduate medical education (GME) programs are paid for the direct costs of GME in accordance with section 1886(h) of the Act.

- Section 1886(b)(3)(B)(viii) of the Act, which requires the Secretary to reduce the applicable percentage increase that would otherwise apply to the standardized amount applicable to a subsection (d) hospital for discharges occurring in a fiscal year if the hospital does not submit data on measures in a form and manner, and at a time, specified by the Secretary.

- Section 1886(o) of the Act, which requires the Secretary to establish a Hospital Value-Based Purchasing (VBP) Program under which value-based incentive payments are made in a fiscal year to hospitals meeting performance standards established for a performance period for such fiscal year.

- Section 1886(p) of the Act, as added by section 3008 of the Affordable Care Act, which establishes a Hospital-Acquired Condition (HAC) Reduction Program, under which payments to applicable hospitals are adjusted to provide an incentive to reduce hospital-acquired conditions.

- Section 1886(q) of the Act, as added by section 3025 of the Affordable Care Act and amended by section 10309 of the Affordable Care Act and section 15002 of the 21st Century Cures Act, which establishes the “Hospital Readmissions Reduction Program.” Under the program, payments for discharges from an “applicable hospital” under section 1886(d) of the Act will be reduced to account for certain excess readmissions. Section 15002 of the 21st Century Cures Act requires the Secretary to compare cohorts of hospitals to each other in determining the extent of excess readmissions.

- Section 1886(r) of the Act, as added by section 3133 of the Affordable Care Act, which provides for a reduction to disproportionate share hospital (DSH) payments under section 1886(d)(5)(F) of the Act and for a new uncompensated care payment to eligible hospitals. Specifically, section 1886(r) of the Act requires that, for fiscal year 2014 and each subsequent fiscal year, subsection (d) hospitals that would otherwise receive a DSH payment made under section 1886(d)(5)(F) of the Act will receive two separate payments: (1) 25 percent of the amount they previously would have received under section 1886(d)(5)(F) of the Act for DSH (“the empirically justified amount”), and (2) an additional payment for the DSH hospital’s proportion of uncompensated care, determined as the product of three factors. These three factors are: (1) 75 percent of the payments that would otherwise be made under section 1886(d)(5)(F) of the Act; (2) 1 minus the percent increase in the percent of individuals who are uninsured (minus 0.2 percentage points for FY 2018 through FY 2019); and (3) a hospital’s uncompensated care amount relative to the uncompensated care amount of all DSH hospitals expressed as a percentage.

- Section 1886(m)(6) of the Act, as amended by section 1206(a)(1) of the Pathway for Sustainable Growth Rate (SGR) Reform Act of 2013 (Pub. L. 113–67), which provides for the establishment of site neutral payment rate criteria under the LTCH PPS with implementation beginning in FY 2016.

- Section 1886(m)(6) of the Act, as amended by section 15009 of the 21st Century Cures Act (Pub. L. 114–255), which provides for a temporary exception to the application of the site neutral payment rate under the LTCH PPS for certain spinal cord specialty hospitals for discharges in cost reporting periods beginning during FYs 2018 and 2019.

- Section 1886(m)(6) of the Act, as amended by section 15010 of the 21st Century Cures Act (Pub. L. 114–255), which provides for a temporary exception to the application of the site neutral payment rate under the LTCH PPS for certain LTCHs with certain discharges with severe wounds occurring in cost reporting periods beginning during FY 2018.

- Section 1886(m)(5)(D)(iv) of the Act, as added by section 1206(c) of the Pathway for Sustainable Growth Rate (SGR) Reform Act of 2013 (Pub. L. 113–67), which provides for the establishment of a functional status quality measure under the LTCH QRP for change in mobility among inpatients requiring ventilator support.

- Section 1899B of the Act, as added by the Improving Medicare Post-Acute Care Transformation Act of 2014 (the IMPACT Act, Pub. L. 113–185), which imposes data reporting requirements for certain post-acute care providers, including LTCHs.


a. MS–DRG Documentation and Coding Adjustment

Section 631 of the American Taxpayer Relief Act (ATRA, Pub. L. 112–240) amended section 7(b)(1)(B) of Public Law 110–90 to require the Secretary to make a recoupment adjustment to the standardized amount of Medicare payments to acute care hospitals to account for changes in MS–DRG documentation and coding that do not reflect real changes in case-mix, totaling $11 billion over a 4-year period of FYs 2014, 2015, 2016, and 2017. The FY 2014 through FY 2017 adjustments represented the amount of the increase in aggregate payments as a result of not completing the prospective adjustment authorized under section 7(b)(1)(A) of Public Law 110–90 until FY 2013. Prior to the ATRA, this amount could not have been recovered under Public Law 110–90. Section 414 of the Medicare Access and CHIP Reauthorization Act (MACRA) of 2015 (Pub. L. 114–10) replaced the single positive adjustment we intended to make in FY 2018 with a 0.5 percent positive adjustment to the standardized amount of Medicare payments to acute care hospitals for FYs 2018 through 2023. The FY 2018 adjustment was subsequently adjusted to 0.4588 percent by section 15005 of the 21st Century Cures Act.

For FY 2018, we are proposing to make the 0.4588 percent positive adjustment to the standardized amount as required by section 414 of Public Law 114–10, as amended by section 15005 of the 21st Century Cures Act.

b. Adjustment to IPPS Rates Resulting From 2-Midnight Policy

In FY 2017, we made a permanent adjustment to the standardized amount, the hospital-specific payment rates, and the national capital Federal rate to prospectively remove the 0.2 percent reduction to the rates put in place in FY 2014 to offset the estimated increase in IPPS expenditures as a result of the 2-midnight policy. In addition, we made a temporary one-time prospective increase to the FY 2017 standardized amount, the hospital-specific payment rates, and the national capital Federal rate of 0.6 percent by including a temporary one-time factor of 1.006 in the calculation of the standardized amount, the hospital-specific payment rates, and the national capital Federal rate to address the effects of the 0.2 percent reduction to the rate for the 2-midnight policy in effect for FYs 2014, 2015, and 2016.

For FY 2018, we are including a factor of (1/1.006) in the calculation of the FY 2018 standardized amount, the hospital-specific payment rates, and the national capital Federal rate to remove the temporary one-time factor of 1.006, as established in the FY 2017 IPPS/LTCH PPS final rule.
c. Reduction of Hospital Payments for Excess Readmissions

We are proposing to make changes to policies for the Hospital Readmissions Reduction Program, which is established under section 1886(q) of the Act, as added by section 3025 of the Affordable Care Act, as amended by section 10309 of the Affordable Care Act. The Hospital Readmissions Reduction Program requires a reduction to a hospital’s base operating DRG payment to account for excess readmissions of selected applicable conditions. For FY 2018 and subsequent years, the reduction is based on a hospital’s risk-adjusted readmission rate during a 3-year period for acute myocardial infarction (AMI), heart failure (HF), pneumonia, chronic obstructive pulmonary disease (COPD), total hip arthroplasty/total knee arthroplasty (THA/TKA), and coronary artery bypass graft (CABG). In this proposed rule, we are proposing the following policies: (1) Specify applicable time period for FY 2018; (2) specify the calculation of aggregate payments for excess readmissions for FY 2018; (3) propose changes to the payment adjustment factor in accordance with the 21st Century Cures Act for FY 2019; and (4) update the Extraordinary Circumstances Exception policy.

d. Hospital Value-Based Purchasing (VBP) Program

Section 1886(o) of the Act requires the Secretary to establish a Hospital VBP Program under which value-based incentive payments are made in a fiscal year to hospitals based on their performance on measures established for a performance period for such fiscal year. In this proposed rule, we are proposing to remove one previously adopted measure, the PSI 90: Patient Safety for Selected Indicators measure, from the Hospital VBP Program beginning with the FY 2019 program year. We also are proposing to adopt one new measure, Hospital-Level, Risk-Standardized Payment Associated with a 30-Day Episode of Care for Pneumonia, beginning with the FY 2022 program year, and to adopt a modified version of a previously adopted measure, Patient Safety and Adverse Events Composite (NQF #0531), beginning with the FY 2023 program year. In addition, we are proposing two modifications to our domain scoring policies beginning with the FY 2019 program year, and further proposing a new weighting methodology for the Efficiency and Cost Reduction domain. We also are inviting public comment on the appropriateness of accounting for social risk factors in the Hospital VBP Program, including which social risk factors should be included; and how to account for these social risk factors in the Hospital VBP Program.

e. Hospital-Acquired Condition (HAC) Reduction Program

Section 1886(p) of the Act, as added under section 3008(a) of the Affordable Care Act, establishes an incentive to hospitals to reduce the incidence of hospital-acquired conditions by requiring the Secretary to make an adjustment to payments to applicable hospitals effective for discharges beginning on October 1, 2014. This 1-percent payment reduction applies to a hospital whose ranking is in the top 25 percent of all applicable hospitals, relative to the national average, of conditions acquired during the applicable period and on all of the hospital’s discharges for the specified fiscal year. In this proposed rule, we are proposing the following policies: (1) Specifying the dates of the time period used to calculate hospital performance for the FY 2020 HAC Reduction Program; (2) requesting comments on additional measures for potential future adoption; (3) requesting comments on social risk factors; (4) requesting comments on accounting for disability and medical complexity in the CDC NHSN measures in Domain 2; and (5) updating the HAC Reduction Program’s Extraordinary Circumstances Exception policy.

f. DSH Payment Adjustment and Additional Payment for Uncompensated Care

Section 3133 of the Affordable Care Act modified the Medicare disproportionate share hospital (DSH) payment methodology beginning in FY 2014. Under section 1886(r) of the Act, which was added by section 3133 of the Affordable Care Act, starting in FY 2014, DSHs receive 25 percent of the amount they previously would have received under the statutory formula for Medicare DSH payments in section 1886(d)(5)(F) of the Act. The remaining amount, equal to 75 percent of the amount that otherwise would have been paid as Medicare DSH payments, is paid as additional payments after the amount is reduced for changes in the percentage of individuals that are uninsured. Each Medicare DSH will receive an additional payment based on its share of the total amount of uncompensated care for all Medicare DSHs for a given time period.

In this proposed rule, we are proposing to update our estimates of the three factors used to determine uncompensated care payments for FY 2018. The statute permits the use of a data source other than the CBO estimates to determine the percent change in the rate of uninsurance as part of the calculation of Factor 2 beginning in FY 2018. We are proposing to use uninsured estimates produced by CMS’ Office of the Actuary (OACT) as part of the development of the National Health Expenditure Accounts (NHEA) in the calculation of Factor 2. We also are proposing to begin incorporating data from Worksheet S–10 in the calculation of hospitals’ share of uncompensated care by combining data on uncompensated care costs from the Worksheet S–10 for FY 2014 with proxy data regarding a hospital’s share of low-income insured days for FYs 2012 and 2013 to determine Factor 3 for FY 2018. The proposal to continue to use data from three cost reporting periods to calculate Factor 3 would have the effect of transitioning from the use of proxy data on low-income insured days toward use of uncompensated care data from Worksheet S–10. As part of this proposal, we are proposing a definition of uncompensated care costs consisting of the sum of charity care and bad debt and a trim methodology to address anomalous charges. We also are proposing that, for Puerto Rico hospitals and Indian Health Service and Tribal hospitals, we would substitute data regarding low-income insured days for FY 2013 for the Worksheet S–10 data from FY 2014 cost reports.

We are proposing to continue the policies that were finalized in FY 2015 to address several specific issues concerning the process and data to be employed in determining hospitals’ share of uncompensated care in the case of hospital mergers. We also are proposing to continue the policies finalized in FY 2017 concerning the methodology for calculating each hospital’s relative share of uncompensated care, such as combining data from multiple cost reports beginning in the same fiscal year and averaging the sum of three individual Factor 3s by the number of cost reporting periods with data. In addition, we are proposing to annualize hospital cost reports that do not span 12 months. We also are proposing to apply a scaling factor to each hospital’s uncompensated care amount so that total uncompensated care payments will be consistent with the estimated amount available to make uncompensated care payments for FY 2018.
g. Proposed Changes to the LTCH PPS

In this proposed rule, we set forth proposed changes to the LTCH PPS Federal payment rates, factors, and other payment rate policies under the LTCH PPS for FY 2018; proposed changes to the payment methodology under the short-stay outlier (SSO) policy; proposals to implement several provisions of the 21st Century Cures Act; and a proposal to adopt a 1-year regulatory delay on the full implementation of the 25-percent threshold policy for discharges occurring in FY 2018 (that is, for the fiscal year after expiration of the current statutory moratoria under the 21st Century Cures Act, which is set to expire September 30, 2017).

h. Hospital Inpatient Quality Reporting (IQR) Program

Under section 1886(b)(3)(B)(viii) of the Act, subsection (d) hospitals are required to report data on measures selected by the Secretary for a fiscal year in order to receive the full annual percentage increase that would otherwise apply to the standardized amount applicable to discharges occurring in that fiscal year. In past years, we have established measures on which hospitals must report data and the process for submittal and validation of the data.

In this proposed rule, we are proposing to make several changes. First, we are proposing to refine two previously adopted measures. Specifically, we are proposing to update the Hospital Consumer Assessment of Healthcare Providers and Systems (HCAPHS) Survey measure by replacing the three existing questions about Pain Management with three new questions that address Communication About Pain During the Hospital Stay, beginning with the FY 2020 payment determination. In addition, we are proposing to update the stroke mortality measure to include the use of NIH Stroke Scale claims data for risk adjustment, beginning with the FY 2023 payment determination.

Second, we are proposing to adopt the Hospital-Wide All-Cause Unplanned Readmission Hybrid Measure as a voluntary measure for the CY 2018 reporting period and note that we are considering proposing this measure as a required measure as early as the CY 2021 reporting period/FY 2023 payment determination and requiring hospitals to submit the core clinical data elements and linking variables used in the measure as early as CY 2020 to support a dry run of the measure during which hospitals would receive a confidential preview of their results in 2021.

Third, we are proposing modifications of our previously finalized eCQMs reporting requirements. For the CY 2017 reporting period/FY 2019 payment determination, we are proposing that hospitals would be required to select and submit six of the available eCQMs included in the Hospital IQR Program measure set and provide two, self-selected, calendar year quarters of data. For the CY 2018 reporting period/FY 2020 payment determination, we are proposing that hospitals would be required to select and submit six of the available eCQMs and provide data for the first three calendar quarters (Q1–Q3). These modifications are being proposed in alignment with proposals for the Medicare and Medicaid EHR Incentive Programs, and would decrease the required number of eCQMs and quarters of reporting as compared with the previously finalized requirements in the FY 2017 IPPS/LTCH PPS final rule. Fourth, we are proposing modifications to the eCQM validation process if our proposals to modify the eCQM reporting requirements for the CY 2017 reporting period/FY 2019 payment determination and CY 2018 reporting period/FY 2020 payment determination are finalized as proposed, whereby hospitals would be required to submit a reduced number of cases for eCQM data validation for the FY 2020 and FY 2021 payment determinations. In addition, we are proposing policies related to the exclusion criteria for hospital selection and the data submission requirements for participating hospitals.

Fifth, we are proposing to modify our educational review process for chart-abstracted measures for the FY 2020 payment determination and subsequent years, such that educational reviews would be offered quarterly for the first three quarters of validation. Hospitals would be allowed 30 calendar days following the date the results of validation are posted to request an educational review. Also, we are proposing that if an educational review demonstrates that the abstraction score calculated by CMS is incorrect, we would use the corrected quarterly score to compute the final confidence interval.

Sixth, we are making proposals related to our Hospital IQR Program Extraordinary Circumstances Extension or Exemptions (ECE) policy, including a change to the name of the policy to Extraordinary Circumstances Exceptions policy.

Finally, we are inviting public comment on accounting for social risk factors in the Hospital IQR Program, the confidential and potential future public reporting of clinical quality measure data stratified by patients’ dual-eligible status, and the following clinical quality measures that we are considering for future inclusion in the Hospital IQR Program: (1) Quality of Informed Consent Documents for Hospital-Performed, Elective Procedures measure; (2) four End-of-Life process and outcome measures for cancer patients; (3) two nurse staffing measures; and (4) eleven newly specified electronic clinical quality measures (eCQMs).

i. Long-Term Care Hospital Quality Reporting Program (LTCH QRP)

Section 1886(m)(5) of the Act requires LTCHs to report certain quality data to CMS in order to receive their full annual update under the LTCH PPS. In this proposed rule, we are proposing to adopt one new outcome measure related to pressure ulcers and two new measures (one process and one outcome) related to ventilator weaning. We also are proposing to define the standardized patient assessment data that LTCHs must report to comply with section 1886(m)(5)(F)(ii) of the Act, as well as the requirements for the reporting of these data. Finally, we are proposing to publicly report data on four assessment-based measures and three claims-based measures.

j. Inpatient Psychiatric Facility Quality Reporting (IPFQR) Program

For the Inpatient Psychiatric Facility Quality Reporting (IPFQR) Program, we are making several proposals. First, beginning with the FY 2020 payment determination, we are proposing the Medication Continuation following Inpatient Psychiatric Discharge measure. Second, beginning with the FY 2019 payment determination (that is, for extraordinary circumstances occurring during CY 2018), we are proposing to update the IPFQR Program’s extraordinary circumstances exception (ECE) policy by: (1) Allowing designated personnel to provide their contact information and sign the ECE request in lieu of the Chief Executive Officer (CEO); (2) allowing up to 90 days after the extraordinary circumstance to submit the request; and (3) stating that we will strive to respond to requests for ECEs within 90 days of receiving these requests. Third, we are proposing to change the annual data submission period from a specific date range to a 45-day period that begins at least 30 days following the end of the collection period. Fourth, we are proposing to align our deadline for submission of a Notice of Participation (NOP) or
program withdrawal with this proposed data submission timeframe. Finally, we are proposing factors by which we will evaluate measures for removal from the IPFQR Program. These factors align with those in use in other quality reporting programs.

3. Summary of Costs and Benefits

- **Adjustment for MS–DRG Documentation and Coding Changes**. Section 414 of the MACRA replaced the single positive adjustment we intended to make in FY 2018 once the recoupment required by section 631 of the ATRA was complete with a 0.5 percent positive adjustment to the standardized amount of Medicare payments to acute care hospitals for FYs 2018 through 2023. The FY 2018 adjustment was subsequently adjusted to 0.4588 percent by section 15005 of the 21st Century Cures Act (Pub. L. 114–255). For FY 2018, we are proposing to make the 0.4588 percent positive adjustment to the standardized amount as required by these provisions.

- **Adjustment to IPPS Payment Rates as a Result of the 2-Midnight Policy**. The removal of the adjustment to IPPS rates resulting from the 2-midnight policy will decrease IPPS payment rates by (1/1.006) for FY 2018. The (1/1.006) is a one-time factor that will be applied to the standardized amount, the hospital-specific rates, and the national capital Federal rate for FY 2018 only.

- **Medicare DSH Payment Adjustment and Additional Payment for Uncompensated Care**. Under section 1886(r) of the Act (as added by section 3133 of the Affordable Care Act), DSH payments to hospitals under section 1886(d)(5)(F) of the Act are reduced and an additional payment for uncompensated care is made to eligible hospitals beginning in FY 2014. Hospitals that receive Medicare DSH payments receive 25 percent of the amount they previously would have received under the statutory formula for Medicare DSH payments in section 1886(d)(5)(F) of the Act. The remainder, equal to an estimate of 75 percent of what otherwise would have been paid as Medicare DSH payments, is the basis for determining the additional payments for uncompensated care after the amount is reduced for changes in the percentage of individuals that are uninsured and additional statutory adjustments. Each hospital that receives Medicare DSH payments will receive an additional payment for uncompensated care based on its share of the total uncompensated care amount reported by Medicare DSHs. The reduction to Medicare DSH payments is not budget neutral.

For FY 2018, we are proposing that the 75 percent of what otherwise would have been paid for Medicare DSH will be adjusted to approximately 58.01 percent of the amount to reflect changes in the percentage of individuals that are uninsured and additional statutory adjustments. In other words, approximately 43.51 percent (the product of 75 percent and 58.01 percent) of our estimate of Medicare DSH payments, prior to the application of section 3133 of the Affordable Care Act, would be available to make additional payments to hospitals for their relative share of the total amount of uncompensated care.

We project that estimated Medicare DSH payments, and additional payments for uncompensated care made for FY 2018, will increase payments overall by approximately 0.8 percent as compared to the estimate of overall payments, including Medicare DSH payments and uncompensated care payments, that will be distributed in FY 2017. The additional payments have redistributive effects based on a hospital’s uncompensated care amount relative to the uncompensated care amount for all hospitals that are estimated to receive Medicare DSH payments, and the calculated payment amount is not directly tied to a hospital’s number of discharges.

- **Proposed Changes to the Hospital Readmissions Reduction Program**. For FY 2018 and subsequent years, the reduction is based on a hospital’s risk-adjusted readmission rate during a 3-year period for acute myocardial infarction (AMI), heart failure (HF), pneumonia, chronic obstructive pulmonary disease (COPD), total hip arthroplasty/total knee arthroplasty (THA/TKA), and coronary artery bypass graft (CABG). Overall, in this proposed rule, we estimate that 2,591 hospitals would have their base operating DRG payments reduced by their determined proxy FY 2018 hospital-specific readmission adjustment. As a result, we estimate that the Hospital Readmissions Reduction Program would save approximately $564 million in FY 2018, an increase of approximately $27 million over the estimated FY 2017 savings.

- **Value-Based Incentive Payments Under the Hospital VBP Program**. We estimate that there would be no net financial impact to the Hospital VBP Program for the FY 2018 program year in the aggregate because, by law, the amount available for value-based incentive payments under the program in a given year must equal the total amount of base operating MS–DRG payment amount reductions for that year, as estimated by the Secretary. The estimated amount of base operating MS–DRG payment amount reductions for the FY 2018 program year and, therefore, the estimated amount available for value-based incentive payments for FY 2018 discharges is approximately $1.9 billion.

- **Proposed Changes to the HAC Reduction Program**. A hospital’s Total HAC score and its ranking in comparison to other hospitals in any given year depends on several different factors. Any significant impact due to the proposed HAC Reduction Program changes for FY 2018, including which hospitals will receive the adjustment, will depend on actual experience.

- **Update to the LTCH PPS Payment Rates and Other Payment Factors**. Based on the best available data for the 415 LTCHs in our database, we estimate that the proposed changes to the payment rates and factors that we are presenting in the preamble and Addendum of this proposed rule, which reflects the rolling end to the transition of the statutory application of the site neutral payment rate required by section 1886(m)(6)(A) of the Act, the proposed update to the LTCH PPS standard Federal payment rate for FY 2018, and estimated changes to the site neutral payment rate and haz-cost-outlier (HCO) payments would result in an estimated decrease in payments from FY 2017 of approximately $238 million.

- **Proposed Changes to the 25-Percent Threshold Policy**. In this proposed rule, we estimate our proposal to adopt a 1-year regulatory delay of the full implementation of the 25-percent threshold policy for discharges occurring in FY 2018 would increase payments to LTCHs in FY 2018 by $50 million.

- **Proposed Changes to the Hospital Inpatient Quality Reporting (IQR) Program**. Across 3,300 IPPS hospitals, we estimate that our policy proposals would result in the following changes to costs and benefits in the Hospital IQR Program compared to previously finalized requirements: (1) A cost reduction of $361,240 for the FY 2019 payment determination due to the proposed updates to the eCQM reporting requirements; (2) a total net cost reduction of $392,963 for the FY 2020 payment determination due to the proposed updates to the eCQM reporting requirements, the proposed updates to the eCQM validation procedures, and the proposed voluntary reporting of the new Hybrid Hospital-Wide Readmissions measure; and (3) a total cost reduction of $70,048 for the FY 2021 payment determination due to...
the proposed updates to the eCQM validation procedures.

- **Proposed Changes Related to the LTCH QRP.** In this proposed rule, we are proposing one outcome measure related to pressure ulcers and two new measures (one process and one outcome) related to ventilator weaning. We also are proposing to specify the use of the standardized patient assessment data as required under section 1899B(b)(1)(B) of the Act and policies regarding public display of measure data. Overall, the cost associated with the proposed changes to the LTCH QRP is estimated at an additional $3,187.15 per LTCH annually, or $1,357,726 for all LTCHs annually.

- **Proposed Changes to the IPFQR Program.** In this proposed rule, we are proposing to adopt one claims based measure, update our ECE process, change the specification of the data submission period, align the timeframe for submission of the NOP or program withdrawal with the data submission period, and establish criteria to evaluate measures for retention or removal. We do not believe that these policies will have any impact on the IPFQR program burden.

**B. Summary**

1. **Acute Care Hospital Inpatient Prospective Payment System (IPPS)**

   Section 1886(d) of the Social Security Act (the Act) sets forth a system of payment for the operating costs of acute care hospital inpatient stays under Medicare Part A (Hospital Insurance) based on prospectively set rates. Section 1886(g) of the Act requires the Secretary to use a prospective payment system (PPS) to pay for the capital-related costs of inpatient hospital services for these “subsection (d) hospitals.” Under these PPSs, Medicare payment for hospital inpatient operating and capital-related costs is made at predetermined, specific rates for each hospital discharge.

   Discharges are classified according to a list of diagnosis-related groups (DRGs).

   The basic payment rate is comprised of a standardized amount that is divided into a labor-related share and a nonlabor-related share. The labor-related share is adjusted by the wage index applicable to the area where the hospital is located. If the hospital is located in Alaska or Hawaii, the nonlabor-related share is adjusted by a cost-of-living adjustment factor. This base payment rate is multiplied by the DRG relative weight.

   If the hospital treats a high percentage of certain low-income patients, it receives a percentage add-on payment applied to the DRG-adjusted base payment rate. This add-on payment, known as the disproportionate share hospital (DSH) adjustment, provides for a percentage increase in Medicare payments to hospitals that qualify under either of two statutory formulas designed to identify hospitals that serve a disproportionate share of low-income patients. For qualifying hospitals, the amount of this adjustment varies based on the outcome of the statutory calculations. The Affordable Care Act revised the Medicare DSH payment methodology and provides for a new additional Medicare payment that considers the amount of uncompensated care beginning on October 1, 2013.

   If the hospital is training residents in an approved residency program(s), it receives a percentage add-on payment for each case paid under the IPPS, known as the indirect medical education (IME) adjustment. This percentage varies, depending on the ratio of residents to beds.

   Additional payments may be made for cases that involve new technologies or medical services that have been approved for special add-on payments. To qualify, a new technology or medical service must demonstrate that it is a substantial clinical improvement over technologies or services otherwise available, and that, absent an add-on payment, it would be inadequately paid under the regular DRG payment.

   The costs incurred by the hospital for a case are evaluated to determine whether the hospital is eligible for an additional payment as an outlier case. This additional payment is designed to protect the hospital from large financial losses due to unusually expensive cases. Any eligible outlier payment is added to the DRG-adjusted base payment rate, plus any DSH, IME, and new technology or medical service add-on adjustments.

   Although payments to most hospitals under the IPPS are made on the basis of the standardized amounts, some categories of hospitals are paid in whole or in part based on their hospital-specific rate, which is determined from their costs in a base year. For example, sole community hospitals (SCHs) receive the higher of a hospital-specific rate based on their costs in a base year (the highest of FY 1982, FY 1987, FY 1996, or FY 2006) or the IPPS Federal rate based on the standardized amount. SCHs are the sole source of care in their areas. Specifically, section 1886(d)(5)(D)(iii) of the Act defines an SCH as a hospital that is located more than 35 road miles from another hospital or that, by reason of factors such as location, weather conditions, travel conditions, or absence of other like hospitals (as determined by the Secretary), is the sole source of hospital inpatient services reasonably available to Medicare beneficiaries. In addition, certain rural hospitals previously designated by the Secretary as essential access community hospitals are considered SCHs.

   Under current law, the Medicare-dependent, small rural hospital (MDH) program is effective through FY 2017. Through and including FY 2006, an MDH received the higher of the Federal rate or the Federal rate plus 50 percent of the amount by which the Federal rate was exceeded by the higher of its FY 1982 or FY 1987 hospital-specific rate. For discharges occurring on or after October 1, 2007, but before October 1, 2017, an MDH receives the higher of the Federal rate or the Federal rate plus 75 percent of the amount by which the Federal rate is exceeded by the highest of its FY 1982, FY 1987, or FY 2002 hospital-specific rate. MDHs are a major source of care for Medicare beneficiaries in their areas. Section 1886(d)(5)(G)(iv) of the Act defines an MDH as a hospital that is located in a rural area, has not more than 100 beds, is not an SCH, and has a high percentage of Medicare discharges (not less than 60 percent of its inpatient days or discharges in its cost reporting year beginning in FY 1987 or in two of its three most recently settled Medicare cost reporting years).

   Section 1886(g) of the Act requires the Secretary to pay for the capital-related costs of inpatient hospital services in accordance with a prospective payment system established by the Secretary. The basic methodology for determining capital prospective payments is set forth in our regulations at 42 CFR 412.308 and 412.312. Under the capital IPPS, payments are adjusted by the same DRG for the case as they are under the operating IPPS. Capital IPPS payments are also adjusted for IME and DSH, similar to the adjustments made under the operating IPPS. In addition, hospitals may receive outlier payments for those cases that have unusually high costs.

   The existing regulations governing payments to hospitals under the IPPS are located in 42 CFR part 412, subparts A through M.

2. **Hospitals and Hospital Units Excluded From the IPPS**

   Under section 1886(d)(1)(B) of the Act, as amended, certain hospitals and hospital units are excluded from the IPPS. These hospitals and units are: Inpatient rehabilitation facility (IRF) hospitals and units; long-term care hospitals (LTCHs); psychiatric hospitals and units; children’s hospitals; cancer hospitals; long-term care neoplastic
1886(d)(3)(B)(iv)(II) of the Act and redesignated by section 15008 of Pub. L. 114–255 and hospitals located outside the 50 States, the District of Columbia, and Puerto Rico (that is, hospitals located in the U.S. Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa). Religious nonmedical health care institutions (RNHCIs) are also excluded from the IPPS. Various sections of the Balanced Budget Act of 1997 (BBA, Pub. L. 105–33), the Medicare, Medicaid and SCHIP Benefits Improvement and Protection Act of 2000 (BIPA, Pub. L. 106–554) provide for the implementation of PPSs for IRF hospitals and units, LTCHs, and psychiatric hospitals and units (referred to as inpatient psychiatric facilities (IPFs)). (We note that the annual updates to the LTCH PPS are now included as part of the IPPS annual update document. Updates to the IRF PPS and IPF PPS are issued as separate documents.) Children’s hospitals, cancer hospitals, hospitals located outside the 50 States, the District of Columbia, and Puerto Rico (that is, hospitals located in the U.S. Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa), and RNHCIs continue to be paid solely under a reasonable cost-based system subject to a rate-of-increase ceiling on inpatient operating costs. The existing regulations governing payments to excluded hospitals and hospital units are located in 42 CFR parts 412 and 413.

3. Long-Term Care Hospital Prospective Payment System (LTCH PPS)

The Medicare prospective payment system (PPS) for LTCHs applies to hospitals described in section 1886(d)(3)(B)(iv) of the Act effective for cost reporting periods beginning on or after October 1, 2002. The LTCH PPS was established under the authority of sections 123 of the BBRA and section 307(b) of the BIPA (as codified under section 1886(m)(1) of the Act). During the 5-year (optional) transition period, a LTCH’s payment under the PPS was based on an increasing proportion of the LTCH Federal rate with a corresponding decreasing proportion based on reasonable cost principles. Effective for cost reporting periods beginning on or after October 1, 2006, all LTCHs are paid 100 percent of the Federal rate. Section 1206(a) of the Pathway for SGR Reform Act of 2013 (Pub. L. 113–67) established the site neutral payment rate under the LTCH PPS, which made the LTCH PPS a dual rate payment system beginning in FY 2016. Under this statute, based on a rolling effective date that is linked to the date on which a given LTCH’s Federal FY 2016 cost reporting period begins, LTCHs are paid for LTCH discharges at the site neutral payment rate unless the discharge meets the patient criteria for payment at the LTCH PPS standard Federal payment rate. The existing regulations governing payment under the LTCH PPS are located in 42 CFR part 412, subpart O. Beginning October 1, 2009, we issue the annual updates to the LTCH PPS in the same documents that update the IPPS (73 FR 26797 through 26798).

4. Critical Access Hospitals (CAHs)

Under sections 1814(l), 1820, and 1834(g) of the Act, payments made to critical access hospitals (CAHs) (that is, rural hospitals or facilities that meet certain statutory requirements) for inpatient and outpatient services are generally based on 101 percent of reasonable cost. Reasonable cost is determined under the provisions of section 1861(v) of the Act and existing regulations under 42 CFR part 413.

5. Payments for Graduate Medical Education (GME)

Under section 1886(a)(4) of the Act, costs of approved educational activities are excluded from the operating costs of inpatient hospital services. Hospitals with approved graduate medical education (GME) programs are paid for the direct costs of GME in accordance with section 1886(h) of the Act. The amount of payment for direct GME costs for a cost reporting period is based on the hospital’s number of residents in that period and the hospital’s costs per resident in a base year. The existing regulations governing payments to the various types of hospitals are located in 42 CFR part 413.

C. Summary of Provisions of Recent Legislation Proposed To Be Implemented In This Proposed Rule


Section 631 of the American Taxpayer Relief Act of 2012 (ATRA) (Pub. L. 112–240) amended section 7(b)(1)(B) of Public Law 110–90 to require CMS to make a recoupment adjustment to the standardized amounts under section 1886(d) of the Act based upon the Secretary’s estimates for discharges occurring from FYs 2014 through FY 2017 to fully offset $11 billion. Once the recoupment required under section 631 of the ATRA was completed, CMS had anticipated making a single positive adjustment in FY 2018 to offset the reductions required to recoup the $11 billion under section 631 of the ATRA. However, section 414 of the MACRA (enacted on April 16, 2015) replaced the single positive adjustment CMS intended to make in FY 2018 with a 0.5 percent positive adjustment for each of FYs 2018 through 2023. Section 15005 of the 21st Century Cures Act (Pub. L. 114–255, enacted December 13, 2016) further amended Public Law 110–90 to reduce the adjustment for FY 2018 from 0.5 percent point to 0.4588 percentage point.


The Pathway for SGR Reform Act of 2013 (Pub. L. 113–67) introduced new payment rules in the LTCH PPS. Under section 1206 of this law, discharges in cost reporting periods beginning on or after October 1, 2015 under the LTCH PPS will receive payment under a site neutral rate unless the discharge meets certain patient-specific criteria. In this proposed rule, we are continuing to provide clarifications to prior policy changes that implemented provisions under section 1206 of the Pathway for SGR Reform Act.


The Improving Medicare Post-Acute Care Transformation Act of 2014 (IMPACT Act) (Pub. L. 113–185), enacted on October 6, 2014, made a number of changes that affect the Long-Term Care Quality Reporting Program (LTCH QRP). In this proposed rule, we are proposing to continue to implement portions of section 1899B of the Act, as added by section 2 of the IMPACT Act, which, in part, requires LTCHs, among other postacute care providers, to report standardized patient assessment data, data on quality measures, and data on resource use and other measures.


Section 411(g) of the Medicare Access and CHIP Reauthorization Act of 2015 (MACRA, Pub. L. 114–10) sets the annual update under the LTCH PPS to 1.0 percent for FY 2018. In this proposed rule, consistent with this requirement, we are proposing to update
the LTCH standard Federal payment rate by 1.0 percent for FY 2018.

The MACRA also extended the MDH program and changes to the payment adjustment for low-volume hospitals through FY 2017. In this proposed rule, we discuss the expiration of the MDH program and the expiration of the temporary changes to the low-volume hospital payment adjustment under current law.


The 21st Century Cures Act (Pub. L. 114–255), enacted on December 13, 2016, contains a number of provisions affecting payments under the LTCH PPS and the Hospital Readmissions Reduction Program and the Medicare EHR Incentive Program, which we are proposing to implement in this proposed rule:

• Section 4002(b)(1)(A) amended section 1848(a)(7)(B) of the Act to provide that the Secretary shall exempt an eligible professional from the application of the payment adjustment under section 1848(a)(7)(A) of the Act with respect to a year, subject to annual renewal, if the Secretary determines that compliance with the requirement for being a meaningful EHR user is not possible because the certified EHR technology used by such eligible professional has been decertified under the Office of the National Coordinator for Health Information Technology’s (ONC) Health IT Certification Program.

• Section 4002(b)(2) amended section 1886(b)(3)(B)(ix)(II) of the Act to provide that the Secretary shall exempt a hospital from the application of the payment adjustment under section 1886(b)(3)(B)(ix)(I) with respect to a fiscal year, subject to annual renewal, if the Secretary determines that compliance with the requirement for being a meaningful EHR user is not possible because the certified EHR technology used by the hospital is decertified under ONC’s Health IT Certification Program.

• Section 5002, which amended section 1886(g)(3) of the Act by adding subparagraphs (D) and (E), which requires the Secretary to develop a methodology for the calculating the excess readmissions adjustment factor for the Hospital Readmissions Reduction Program based on cohorts defined by the percentage of dual eligible patients (that is, patients who are eligible for both Medicare and full-benefit Medicaid coverage) cared for by a hospital. In this proposed rule, we are proposing to implement changes to the payment adjustment factor to assess penalties based on a hospital’s performance relative to other hospitals treating a similar proportion of dual-eligible patients.

• Section 15004(a), which further amended section 114(d)(7) of the MMSEA (as amended) by striking “The moratorium under paragraph (1)(A)” and inserting “[a]ny moratorium under paragraph (1)” and specified that such amendment shall take effect as if included in the enactment of section 112 of the PAMA. We are proposing to implement the exceptions to the current statutory moratorium, which is in effect through September 30, 2017, on increasing beds in an existing LTCH or an existing LTCH satellite as provided by Section 15004(a).

• Section 15004(b), which modifies high cost outlier payments to LTCH standard Federal rate cases beginning in FY 2018.

• Section 15006, which further amended section 114(c)(1)(A) of the MMSEA (as amended) by extending the moratorium on the full implementation of the 25-percent threshold policy through June 30, 2016, and for discharges occurring on or after October 1, 2016, and before October 1, 2017. In this proposed rule, we are implementing the moratorium on the full implementation of the 25-percent threshold policy for discharges occurring on or after October 1, 2016, through September 30, 2017, as provided by section 15006.

• Section 15007, which amended section 1206(a)(3) of the Pathway for SGR Reform Act by extending the exclusion of Medicare Advantage plans’ and site neutral payment rate discharges from the calculation of the average length-of-stay to all LTCHs, for discharges occurring in cost reporting periods beginning on or after October 1, 2015.

• Section 15008, which provided for a change in Medicare classification for “subclause (II)” LTCHs by redesignating such hospitals from section 1886(d)(1)(B)(iv)(II) to section 1886(d)(1)(B)(vi) of the Act. In this proposed rule, we are proposing to implement the reclassification of hospitals which had previously been classified as “subclause (II)” LTCHs as their own category of IPPS-excluded hospitals as provided by the provisions of section 15008.

• Section 15009 of Public Law 114–255, which added new subparagraph (F) to section 1886(m)(6) of the Act, providing for a temporary exception to the site neutral payment rate for certain spinal cord specialty hospitals for all discharges occurring during FYs 2018 and 2019.

• Section 15010, which added a new subparagraph (G) to section 1886(m)(6) of the Act, to create a temporary exception to the site neutral payment rate for certain severe wound discharges from certain LTCHs during such LTCH’s cost reporting period beginning during FY 2018.

Public Law 114–255 also amended section 1886(q)(3) of the Act by adding subparagraphs (D) and (E), which requires the Secretary to develop a methodology for the Hospital Readmissions Reduction Program that accounts for the percentage of dual-eligible patients (that is, patients who are eligible for both Medicare and full-benefit Medicaid coverage) cared for by a hospital. In this proposed rule, we are proposing to implement changes to the payment adjustment factor to assess penalties based on a hospital’s performance relative to other hospitals treating a similar proportion of dual-eligible patients.

• Section 16003 amended section 1848(a)(7)(D) of the Act to provide that no payment adjustment may be made under section 1848(a)(7)(A) of the Act for 2017 and 2018 in the case of an eligible professional who furnishes substantially all of his or her covered professional services in an ambulatory surgical center (ASC). Section 1848(a)(7)(D)(iii) of the Act provides that determinations of whether an eligible professional is ASC-based may be made based on the site of service as defined by the Secretary or an attestation, but shall be made without regard to any employment or billing arrangement between the eligible professional and any other supplier or provider of services. Section 1848(a)(7)(D)(iv) of the Act provides that the ASC-based exception shall no longer apply as of the first year that begins more than 3 years after the date on which the Secretary determines, through notice-and-comment rulemaking, that certified EHR technology applicable to the ASC setting is available.

D. Summary of Provisions of This Proposed Rule

In this proposed rule, we are setting forth proposed payment and policy changes to the Medicare IPPS for FY 2018 operating costs and for capital-related costs of acute care hospitals and certain hospitals and hospital units that are excluded from IPPS. In addition, we are setting forth proposed changes to the payment rates, factors, and other payment and policy-related changes to programs associated with payment rate policies under the LTCH PPS for FY 2018.
Below is a summary of the major changes that we are proposing to make:

1. Proposed Changes to MS–DRG Classifications and Recalibrations of Relative Weights

In section II. of the preamble of this proposed rule, we include—
• Proposed changes to MS–DRG classifications based on our yearly review for FY 2018.
• Proposed adjustment to the standardized amounts under section 1886(d) of the Act for FY 2018 in accordance with the amendments made to section 7(b)(1)(B) of Public Law 110–90 by section 414 of the MACRA and section 15005 of the 21st Century Cures Act.
• Proposed recalibrations of the MS–DRG relative weights.
• A discussion of the FY 2018 status of new technologies approved for add-on payments for FY 2017 and a presentation of our evaluation and analysis of the FY 2018 applicants for add-on payments for high-cost new medical services and technologies (including public input, as directed by Pub. L. 108–173, obtained in a town hall meeting).

2. Proposed Changes to the Hospital Wage Index for Acute Care Hospitals

In section III. of the preamble to this proposed rule, we are proposing to make revisions to the wage index for acute care hospitals and the annual update of the wage data. Specific issues addressed include, but are not limited to, the following:
• The proposed FY 2018 wage index update using wage data from cost reporting periods beginning in FY 2014.
• Clarification of other wage-related costs in the wage index.
• Calculation of the proposed occupational mix adjustment for FY 2018 based on the 2013 Occupational Mix Survey.
• Analysis and implementation of the proposed FY 2018 occupational mix adjustment to the wage index for acute care hospitals.
• Proposed application of the rural floor and the frontier State floor and the proposed expiration of the imputed floor.
• Proposed revisions to the wage index for acute care hospitals based on hospital redesignations and reclassifications under sections 1886(d)(8)(B), (d)(8)(E), and (d)(10) of the Act.
• Proposal to require documentation of SCH and RRC classification status approval to be submitted to the MGCRB by the first business day after January 1.

3. Proposed Revising and Rebasining of Hospital Market Basket

In section IV. of this proposed rule, we are proposing to revise and rebase the hospital market baskets for acute care hospitals and update the labor-related share.

4. Other Decisions and Proposed Changes to the IPPS for Operating Costs

In section V. of the preamble of this proposed rule, we discuss proposed changes or clarifications of a number of the provisions of the regulations in 42 CFR parts 412 and 413, including the following:
• Proposed changes to MS–DRGs subject to the postacute care transfer policy.
• Proposed changes to the inpatient hospital update for FY 2018.
• Proposed changes to the volume decrease adjustment for SCHs.
• Proposed updated national and regional case-mix values and discharges for purposes of determining RRC status.
• Expiration of the MDH program and the temporary changes to the payment adjustment for low-volume hospitals at the end of FY 2017.
• Proposed parallel low-volume hospital payment adjustment concerning hospitals operated by the Indian Health Service (IHS) or a Tribe.
• The statutorily required IME adjustment factor for FY 2018.
• Proposed changes to the methodologies for determining Medicare DSH payments and the additional payments for uncompensated care.
• Discussion of expiration of the MDH program at the end of FY 2017 and our policy to allow MDHs to apply for SCH status in advance of the expiration of the MDH program and be paid as such under certain conditions.
• Proposed changes to the rules for payment adjustments under the Hospital Readmissions Reduction Program based on hospital readmission measures and the process for hospital review and correction of those rates for FY 2018.

5. Proposed Changes to the LTCH PPS

In section VI. of the preamble to this proposed rule, we discuss the proposed payment policy requirements for capital-related costs and capital payments to hospitals for FY 2018.

6. Proposed Changes to the Payment Rates for Certain Excluded Hospitals: Rate-of-Increase Percentages

In section VII. of the preamble of this proposed rule, we discuss—
• Proposed changes to payments to certain excluded hospitals for FY 2018.
• Proposed policy changes relating to payments to hospitals-within-hospitals.
• Proposed continued implementation of the Frontier Community Health Integration Project (FCHIP) Demonstration.

7. Proposed Changes to the LTCH PPS

In section VIII. of the preamble of this proposed rule, we set forth—
• Proposed changes to the LTCH PPS Federal payment rates, factors, and other payment rate policies under the LTCH PPS for FY 2018.
• Proposed changes to the short-stay outlier (SSO) policy.
• Proposed 1-year regulatory delay of the full implementation of the 25-percent threshold policy for discharges occurring in FY 2018.
• Proposed changes to implement the temporary exception to the site neutral payment rate for certain spinal cord specialty hospitals and for certain discharges with severe wounds from certain LTCHs, as provided under sections 15009 and 15010 of Public Law 114–255, respectively.
8. Proposed Changes Relating to Quality Data Reporting for Specific Providers and Suppliers

In section IX. of the preamble of the proposed rule, we address—

• Proposed requirements for the Hospital Inpatient Quality Reporting (IQR) Program.
• Proposed changes to the requirements for the quality reporting program for PPS-exempt cancer hospitals (PCHQR Program).
• Proposed changes to the requirements under the LTCH Quality Reporting Program (LTCH QRP).
• Proposed changes to the requirements under the Inpatient Psychiatric Facility Quality Reporting (IPFQR) Program.
• Proposed changes to requirements pertaining to the clinical quality measurement of eligible hospitals and CAHs as well as EPs participating in the Medicare and Medicaid Electronic Health Record (EHR) Incentive Programs.


In section X. of the preamble of this proposed rule, we present our proposals to revise the regulations to allow providers to use an electronic signature to sign the Certification and Settlement Summary page of the Medicare cost report and submit this page electronically, and clarify the rules relating to the sale or scrapping of depreciable assets disposed of on or after December 1, 1997.

10. Proposed Changes Relating to Survey and Certification Requirements

In section XI. of the preamble of this proposed rule, we present our proposals for allowing transparency in accrediting organization survey reports and plans of correction and for changing the requirement for providers to publish self-termination notices in newspapers.

11. Determining Prospective Payment Operating and Capital Rates and Rate-of-Increase Limits for Acute Care Hospitals

In section V. of the Addendum to this proposed rule, we set forth proposed changes to the amounts and factors for determining the proposed FY 2018 prospective payment rates for operating costs and capital-related costs for acute care hospitals. We are proposing to establish the threshold amounts for outlier cases. In addition, we are addressing the update factors for determining the rate-of-increase limits for cost reporting periods beginning in FY 2018 for certain hospitals excluded from the IPPS.

12. Determining Prospective Payment Rates for LTCHs

In the Addendum to this proposed rule, we set forth proposed changes to the amounts and factors for determining the proposed FY 2018 LTCH PPS standard Federal payment rate and other factors used to determine LTCH PPS payments under both the LTCH PPS standard Federal payment rate and the site neutral payment rate in FY 2018. We are proposing to establish the adjustments for wage levels, the labor-related share, the cost-of-living adjustment, and high-cost outliers, including the applicable fixed-loss amounts and the LTCH cost-to-charge ratios (CCRs) for both payment rates.

13. Impact Analysis

In Appendix A of this proposed rule, we set forth an analysis of the impact that the proposed changes would have on affected acute care hospitals, CAHs, LTCHs, PCHs, and IPFs.

14. Recommendation of Update Factors for Operating Cost Rates of Payment for Hospital Inpatient Services

In Appendix B of this proposed rule, as required by sections 1886(e)(4) and (e)(5) of the Act, we are providing our recommendations of the appropriate percentage changes for FY 2018 for the following:

• A single average standardized amount for all areas for hospital inpatient services paid under the IPPS for operating costs of acute care hospitals (and hospital-specific rates applicable to SCHs).
• Target rate-of-increase limits to the allowable operating costs of hospital inpatient services furnished by certain hospitals excluded from the IPPS.
• The LTCH PPS standard Federal payment rate and the site neutral payment rate for hospital inpatient services provided for LTCH PPS discharges.

15. Discussion of Medicare Payment Advisory Commission Recommendations

Under section 1805(b) of the Act, MedPAC is required to submit a report to Congress, no later than March 15 of each year, in which MedPAC reviews and makes recommendations on Medicare payment policies. MedPAC’s March 2017 recommendations concerning hospital inpatient payment policies address the update factor for hospital inpatient operating costs and capital-related costs for hospitals under the IPPS. We address these recommendations in Appendix B of this proposed rule. For further information relating specifically to the MedPAC March 2017 report or to obtain a copy of the report, contact MedPAC at (202) 220–3700 or visit MedPAC’s Web site at: http://www.medpac.gov.

II. Proposed Changes to Medicare Severity Severity Diagnosis-Related Group (MS–DRG) Classifications and Relative Weights

A. Background

Section 1886(d) of the Act specifies that the Secretary shall establish a classification system (referred to as diagnosis-related groups (DRGs)) for inpatient discharges and adjust payments under the IPPS based on appropriate weighting factors assigned to each DRG. Therefore, under the IPPS, Medicare pays for inpatient hospital services on a rate per discharge basis that varies according to the DRG to which a beneficiary’s stay is assigned. The formula used to calculate payment for a specific case multiplies an individual hospital’s payment rate per case by the weight of the DRG to which the case is assigned. Each DRG weight represents the average resources required to care for cases in that particular DRG, relative to the average resources used to treat cases in all DRGs.

Section 1886(d)(4)(C) of the Act requires that the Secretary adjust the DRG classifications and relative weights at least annually to account for changes in resource consumption. These adjustments are made to reflect changes in treatment patterns, technology, and any other factors that may change the relative use of hospital resources.

B. MS–DRG Reclassifications

For general information about the MS–DRG system, including yearly reviews and changes to the MS–DRGs, we refer readers to the previous discussions in the FY 2010 IPPS/RY 2010 LTCH PPS final rule (74 FR 43764 through 43766) and the FYs 2011 through 2017 IPPS/LTCH PPS final rules (75 FR 50053 through 50055; 76 FR 51485 through 51487; 77 FR 53273; 78 FR 50512; 79 FR 49871; 80 FR 49342; and 81 FR 56787 through 56872, respectively).
C. Adoption of the MS–DRGs in FY 2008

For information on the adoption of the MS–DRGs in FY 2008, we refer readers to the FY 2008 IPPS final rule with comment period (72 FR 47140 through 47189).

D. Proposed FY 2018 MS–DRG Documentation and Coding Adjustment

1. Background on the Prospective MS–DRG Documentation and Coding Adjustments for FY 2008 and FY 2009

Authorized by Public Law 110–90

In the FY 2008 IPPS final rule with comment period (72 FR 47140 through 47189), we adopted the MS–DRG patient classification system for the IPPS, effective October 1, 2007, to better recognize severity of illness in Medicare payment rates for acute care hospitals. The adoption of the MS–DRG system resulted in the expansion of the number of DRGs from 538 in FY 2007 to 745 in FY 2008. By increasing the number of MS–DRGs and more fully taking into account patient severity of illness in Medicare payment rates for acute care hospitals, MS–DRGs encourage hospitals to improve their documentation and coding of patient diagnoses.

In the FY 2008 IPPS final rule with comment period (72 FR 47175 through 47186), we indicated that the adoption of the MS–DRGs had the potential to lead to increases in aggregate payments without a corresponding increase in actual patient severity of illness due to the incentives for additional documentation and coding. In that final rule with comment period, we exercised our authority under section 1886(d)(3)(A)(vi) of the Act, which authorizes us to maintain budget neutrality by adjusting the national standardized amount, to eliminate the estimated effect of changes in coding or classification that do not reflect real changes in case-mix. Our actuaries estimated that maintaining budget neutrality required an adjustment of −4.8 percentage points to the national standardized amount. We provided for phasing in this −4.8 percentage point adjustment over 3 years. Specifically, we established prospective documentation and coding adjustments of −1.2 percentage points for FY 2008, −1.8 percentage points for FY 2009, and −1.8 percentage points for FY 2010.

On September 29, 2007, Congress enacted the TMA [Transitional Medical Assistance], Abstinence Education, and QI [Qualifying Individuals] Programs Extension Act of 2007 (Pub. L. 110–90). Section 7(a) of Public Law 110–90 reduced the documentation and coding adjustment made as a result of the MS–DRG system that we adopted in the FY 2008 IPPS final rule with comment period to −0.6 percentage point for FY 2008 and −0.9 percentage point for FY 2009.

As discussed in prior year rulemaking, and most recently in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56780 through 56782), we implemented a series of adjustments required under sections 7(b)(1)(A) and 7(b)(1)(B) of Public Law 110–90, based on a retrospective review of FY 2008 and FY 2009 claims data. We completed those adjustments in FY 2013, but indicated in the FY 2013 IPPS/LTCH PPS final rule (77 FR 53274 through 53275) that delaying full implementation of the adjustment required under section 7(b)(1)(A) of Public Law 110–90 until FY 2013 resulted in payments in FY 2010 through FY 2012 being overstated, and that these overpayments could not be recovered.

2. Recoupment or Repayment Adjustment Authorized by Section 631 of the American Taxpayer Relief Act of 2012 (ATRA)

Section 631 of the ATRA amended section 7(b)(1)(B) of Public Law 110–90 to require the Secretary to make a recoupment adjustment or adjustments totaling $11 billion by FY 2017. This adjustment represented the amount of the increase in aggregate payments as a result of not completing the prospective adjustment authorized under section 7(b)(1)(A) of Public Law 110–90 until FY 2013. As discussed earlier, this delay in implementation resulted in overstated payment rates in FYs 2010, 2011, and 2012. The resulting overpayments could not have been recovered under Public Law 110–90. Similar to the adjustments authorized under section 7(b)(1)(B) of Public Law 110–90, the adjustment required under section 631 of the ATRA was a one-time recoupment of a prior overpayment, not a permanent reduction to payment rates.

Therefore, we anticipated that any adjustment made to reduce payment rates in one year would eventually be offset by a positive adjustment in 2018, once the necessary amount of overpayment was recovered. However, section 414 of the Medicare Access and CHIP Reauthorization Act (MACRA) of 2015, Public Law 114–10, enacted on April 16, 2015, replaced the single positive adjustment we intended to make in FY 2018 with a 0.5 percentage point positive adjustment for each of FYs 2016 through 2023. We stated in the FY 2016 IPPS/LTCH PPS final rule (80 FR 49345) that we would address this MACRA provision in future rulemaking.

However, section 15005 of the 21st Century Cures Act (Pub. L. 114–255), enacted on December 13, 2016, reduced the adjustment for FY 2018 from 0.5 percentage points to 0.4588 percentage points. We are addressing these provisions of MACRA and the 21st Century Cures Act in section I.D.3. of the preamble of this proposed rule.

As we stated in the FY 2014 IPPS/LTCH PPS final rule (78 FR 50515 through 50517), our actuaries estimated that a −9.3 percentage point adjustment to the standardized amount would be necessary if CMS were to fully recover the $11 billion recoupment required by section 631 of the ATRA in FY 2014. It is often our practice to phase in payment rate adjustments over more than one year, in order to moderate the effect on payment rates in any one year. Therefore, consistent with the policies that we have adopted in many similar cases, and after consideration of the public comments we received, in the FY 2014 IPPS/LTCH PPS final rule (78 FR 50515 through 50517), we implemented a −0.8 percentage point recoupment adjustment to the standardized amount in FY 2014. We estimated that if adjustments of approximately −0.8 percentage point were implemented in FYs 2014, 2015, 2016, and 2017, using standard inflation factors, the entire $11 billion would be accounted for by the end of the statutory 4-year timeline. As estimates of any future adjustments are subject to variations in total savings, we did not provide for specific adjustments for FYs 2015, 2016, or 2017 at that time.

Consistent with this approach discussed in the FY 2014 rulemaking for recouping the $11 billion required by section 631 of the ATRA, in the FY 2015 IPPS/LTCH PPS final rule (79 FR 49874) and the FY 2016 IPPS/LTCH PPS final rule (80 FR 49345), we implemented additional −0.8 percentage point recoupment adjustments to the standardized amount in FY 2015 and FY 2016, respectively. We estimated that these adjustments, combined with leaving the prior −0.8 percentage point adjustments in place, would recover up to $2 billion in FY 2015 and another $3 billion in FY 2016. When combined with the approximately $1 billion adjustment made in FY 2014, we estimated that approximately $5 to $6 billion would be left to recover under section 631 of the ATRA by the end of FY 2016.

As indicated in the FY 2017 IPPS/LTCH PPS proposed rule (81 FR 24966), due to lower than previously estimated inpatient spending, we determined that an adjustment of −0.8 percentage point in FY 2017 would not recoup the $11 billion under section 631 of the ATRA.
For the FY 2017 IPPS/LTCH PPS final rule (81 FR 56785), based on the Midsession Review of the President’s FY 2017 Budget, our actuaries estimated that, to the nearest tenth of a percentage point, the FY 2017 documentation and coding adjustment factor that will recoup as closely as possible $11 billion from FY 2014 through FY 2017 without exceeding this amount is −1.5 percentage points. Based on those updated estimates by the Office of the Actuary using the Midsession Review of the President’s FY 2017 Budget, we made a −1.5 percentage point adjustment for FY 2017 as the final adjustment required under section 631 of the ATRA. The estimates by our actuaries related to this finalized adjustment were included in a memorandum that we made publicly available on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/FY2017-IPPS-Final-Rule-Home-Page-Items/FY2017-IPPS-Final-Rule-OACT.html.

3. Proposed Adjustment for FY 2018 Required Under Section 414 of Public Law 114–10 (MACRA) and Section 15005 of Public Law 114–255

As stated in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56785), once the recoupment required under section 631 of the ATRA was complete, we had anticipated making a single positive adjustment in FY 2018 to offset the reductions required to recoup the $11 billion under section 631 of the ATRA. However, section 414 of the MACRA (which was enacted on April 16, 2015) replaced the single positive adjustment we intended to make in FY 2018 with a 0.5 percentage point positive adjustment for each of FYs 2018 through 2023. In the FY 2017 rulemaking, we indicated that we would address the adjustments for FY 2018 and later fiscal years in future rulemaking. As noted previously, section 15005 of the 21st Century Cures Act (Pub. L. 114–255), which was enacted on December 13, 2016, amended section 7(b)(1)(B) of the TMA, as amended by section 631 of the ATRA and section 414 of the MACRA, to reduce the adjustment for FY 2018 from a 0.5 percentage point to a 0.4588 percentage point. We believe the directive under section 15005 of Public Law 114–255 is clear. Therefore, for FY 2018, we are proposing to implement the required +0.4588 percentage point adjustment to the standardized amount. This is a permanent adjustment to payment rates. While we are not proposing future adjustments required under section 414 of the MACRA and section 15005 of Public Law 114–255 at this time, we expect to propose positive 0.5 percentage point adjustments to the standardized amounts for FYs 2019 through 2023.

E. Refinement of the MS–DRG Relative Weight Calculation

1. Background

Beginning in FY 2007, we implemented relative weights for DRGs based on cost report data instead of charge information. We refer readers to the FY 2007 IPPS final rule (71 FR 47882) for a detailed discussion of our final policy for calculating the cost-based DRG relative weights and to the FY 2008 IPPS final rule with comment period (72 FR 47199) for information on how we blended relative weights based on the CMS DRGs and MS–DRGs. We also refer readers to the FY 2017 IPPS/LTCH PPS final rule (81 FR 56785 through 56787) for a detailed discussion of the history of changes to the number of cost centers used in calculating the DRG relative weights. Since FY 2014, we calculate the IPPS MS–DRG relative weights using 19 CCRs, which now include distinct CCRs for implantable devices, MRIs, CT scans, and cardiac catheterization.

2. Discussion of Policy for FY 2018

Consistent with our established policy, we calculated the proposed MS–DRG relative weights for FY 2018 using two data sources: The MedPAR file as the claims data source and the HCRIS as the cost report data source. We adjusted the charges from the claims to costs by applying the 19 national average CCRs developed from the cost reports. The description of the calculation of the proposed 19 CCRs and the proposed MS–DRG relative weights for FY 2018 is included in section II.G. of the preamble to this FY 2018 IPPS/LTCH PPS proposed rule. As we did with the FY 2017 IPPS/LTCH PPS final rule, for this proposed rule, we are providing the version of the HCRIS from which we calculated these proposed 19 CCRs on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html. Click on the link on the left side of the screen titled, “FY 2018 IPPS Proposed Rule Home Page” or “Acute Inpatient Files for Download.”

F. Proposed Changes to Specific MS–DRG Classifications

1. Discussion of Changes to Coding System and Basis for Proposed FY 2018 MS–DRG Updates

a. Conversion of MS–DRGs to the International Classification of Diseases, 10th Revision (ICD–10)

As of October 1, 2015, providers use the International Classification of Diseases, 10th Revision (ICD–10) coding system to report diagnoses and procedures for Medicare hospital inpatient services under the MS–DRG system instead of the ICD–9–CM coding system, which was used through September 30, 2015. The ICD–10 coding system includes the International Classification of Diseases, 10th Revision, Clinical Modification (ICD–10–CM) for diagnosis coding and the International Classification of Diseases, 10th Revision, Procedure Coding System (ICD–10–PCS) for inpatient hospital procedure coding, as well as the Official ICD–10–CM and ICD–10–PCS Guidelines for Coding and Reporting. For a detailed discussion of the conversion of the MS–DRGs to ICD–10, we refer readers to the FY 2017 IPPS/LTCH PPS final rule (81 FR 56787 through 56789).

b. Basis for FY 2018 Proposed MS–DRG Updates

CMS has previously encouraged input from our stakeholders concerning the annual IPPS updates when that input is made available to us by December 7 of the year prior to the next annual proposed rule update. For example, to be considered for any updates or changes in FY 2018, comments and suggestions should have been submitted by December 7, 2016. The comments that were submitted in a timely manner for FY 2018 are discussed in this section of the preamble of this proposed rule. As CMS works with the public to examine the ICD–10 claims data used for updates to the ICD–10 MS–DRGs, we would like to examine areas where the MS–DRGs can be improved. This will require additional time for us to review requests from the public to make specific updates, analyze claims data, and consider any proposed updates. Given the need for more time to carefully evaluate requests and propose updates, we are changing the deadline to request updates to MS–DRGs to November 1 of each year. This will provide an additional 5 weeks for the data analysis and review process. Interested parties should submit any comments and suggestions for FY 2019 by November 1, 2017, via the CMS MS–
As shown in the table above, for MS–DRG 052, there were a total of 865 cases with an average length of stay of 5.4 days and average costs of $10,247. Of the 865 cases in MS–DRG 052, there were 63 cases that reported a principal diagnosis of functional quadriplegia, with an average length of stay of 4.9 days and average costs of $6,420. For MS–DRG 053, there were a total of 239 cases, with an average length of stay of 3.3 days and average costs of $6,326. Of the 239 cases in MS–DRG 053, there were 16 cases that reported a principal diagnosis of functional quadriplegia, with an average length of stay of 3.3 days and average costs of $2,318.

To address the request to reassign cases reporting a diagnosis of functional quadriplegia to a different MS–DRG, we reviewed the data for a total of 79 cases (63 cases in MS–DRG 052 and 16 cases in MS–DRG 053) that reported a principal diagnosis of functional quadriplegia in MS–DRGs 052 and 053. As shown in the table above, our data analysis demonstrates that the average costs for these 79 cases are lower than the average costs of all cases in MS–DRGs 052 and 053 ($6,420 compared to $10,247 for all cases in MS–DRG 052, and $2,318 compared to $6,326 for all cases in MS–DRG 053), and the average
lengths of stay are shorter for cases reporting a diagnosis of functional quadriplegia in MS–DRG 052 (4.9 days compared to 5.4 days for all cases in MS–DRG 052), but equal for cases in MS–DRG 053 (3.3 days for cases reporting a diagnosis of functional quadriplegia and for all cases).

Our clinical advisors reviewed this issue and agreed that a diagnosis of functional quadriplegia does not involve a spinal disorder or injury, and may be associated with, or the result of, a variety of underlying conditions. Our clinical advisors also agreed that it is not clinically appropriate to include cases reporting a diagnosis of functional quadriplegia within MS–DRGs 052 and 053 because these cases do not involve a spinal disorder or injury. Therefore, given the fact that functional quadriplegia can be the result of a variety of other conditions, we reviewed the MS–DRGs in order to identify a more appropriate placement for cases reporting this diagnosis. Our clinical advisors recommended assigning cases representing a diagnosis of functional quadriplegia from MS–DRGs 052 and 053 to MS–DRGs 091, 092, and 093 (Other Disorders of Nervous System with MCC, with CC, and without CC/MCC, respectively). Within each MDC, there are MS–DRGs that describe a variety of other conditions that do not have the clinical characteristics of the more specific MS–DRGs. In this case, MS–DRGs 091, 092, and 093 describe a variety of other disorders of the nervous system that are not clinically similar in characteristics to the disorders described by MS–DRGs 052 and 053. Our clinical advisors believe that MS–DRGs 091, 092, and 093 are more appropriate MS–DRG assignments for cases representing a diagnosis of functional quadriplegia.

We examined claims data from the December 2016 update of the FY 2016 MedPAR file on cases in MS–DRGs 091, 092, and 093. Our findings are shown in the table below.

### Cases in MS–DRGs 091, 092, and 093

<table>
<thead>
<tr>
<th>MS–DRG</th>
<th>Number of cases</th>
<th>Average length of stay</th>
<th>Average costs</th>
</tr>
</thead>
<tbody>
<tr>
<td>MS–DRG 091—All cases</td>
<td>12,607</td>
<td>5.6</td>
<td>$10,815</td>
</tr>
<tr>
<td>MS–DRG 092—All cases</td>
<td>19,392</td>
<td>3.9</td>
<td>6,706</td>
</tr>
<tr>
<td>MS–DRG 093—All cases</td>
<td>8,120</td>
<td>2.7</td>
<td>5,253</td>
</tr>
</tbody>
</table>

As shown in the table above, for MS–DRG 091, there were a total of 12,607 cases, with an average length of stay of 5.6 days and average costs of $10,815. For MS–DRG 092, there were a total of 19,392 cases, with an average length of stay of 3.9 days and average costs of $6,706. For MS–DRG 093, there were a total of 8,120 cases, with an average length of stay of 2.7 days and average costs of $5,253. As stated earlier, of the 865 total cases in MS–DRG 052, there were 63 cases that reported a principal diagnosis of functional quadriplegia, with an average length of stay of 4.9 days and average costs of $6,420. Of the 239 total cases in MS–DRG 053, there were 16 cases that reported a principal diagnosis of functional quadriplegia, with an average length of stay of 3.3 days and average costs of $2,318. The average lengths-of-stay for cases reporting a diagnosis of functional quadriplegia in MS–DRGs 052 and 053 are similar to the average lengths of stay for cases found in MS–DRGs 091, 092 and 093 (4.9 days and 3.3 days for cases in MS–DRGs 052 and 053, respectively, compared to 5.6 days, 3.9 days, and 2.7 days, respectively, for cases in MS–DRGs 091, 092, and 093). The average costs for cases reporting a diagnosis of functional quadriplegia in MS–DRGs 052 and 053 are $6,420 and $2,318, respectively, compared to $10,815, $6,706, and $5,253 for all cases in MS–DRGs 091, 092, and 093. The average costs for cases reporting a diagnosis of functional quadriplegia in MS–DRG 053 are lower than the average costs for all cases in MS–DRG 093 without a CC or MCC ($2,318 compared to $5,253, respectively). The average costs for cases reporting a diagnosis of functional quadriplegia in MS–DRG 052 are $6,420, which is lower than the average costs of $10,815 for all cases in MS–DRG 091, but close to the average costs of $6,706 for all cases in MS–DRG 092. While we acknowledge that the average costs for cases reporting a diagnosis of functional quadriplegia are lower than those cases within MS–DRGs 091, 092, and 093, as stated earlier, the average costs of cases reporting a diagnosis of functional quadriplegia also are lower than the average costs of all cases in MS–DRGs 052 and 053 where these cases are currently assigned.

Our clinical advisors reviewed the clinical issues as well as the claims data for MS–DRGs 052, 053, 091, 092, and 093. As a result of this review, they recommended that cases reporting a diagnosis of functional quadriplegia be reassigned from MS–DRGs 052 and 053 to MS–DRGs 091, 092, and 093 because the current MS–DRG assignment is not clinically appropriate. Our clinical advisors stated that reassigning these cases to MS–DRGs 091, 092, and 093 is more appropriate because this set of MS–DRGs includes a variety of nervous system disorders that are not appropriately classified to more specific MS–DRGs within MDC 1. Therefore, we are proposing to reassign cases identified by diagnosis code R53.2 from MS–DRGs 052 and 053 to MS–DRGs 091, 092, and 093 for FY 2018.

We are inviting public comments on our proposal.

b. Responsive Neurostimulator (RNS©) System

We received a request to modify the MS–DRG assignment for cases involving the use of the RNS® neurostimulator, a cranially implanted neurostimulator that is a treatment option for persons diagnosed with medically intractable epilepsy. Cases involving the use of the RNS® neurostimulator are assigned to MS–DRG 023 (Craniotomy with Major Device Implant or Acute Complex Central Nervous System (CNS) Principal Diagnosis (PDX) with MCC or Chemo Implant) and MS–DRG 024 (Craniotomy with Major Device Implant or Acute Complex Central Nervous System (CNS) Principal Diagnosis (PDX) without MCC). Cases involving the use of the RNS® neurostimulator generator and leads are captured within the descriptions of four ICD–10–PCS codes. ICD–10–PCS code 0NH00NZ (Insertion of neurostimulator generator into skull, open approach) captures the use of the neurostimulator generator, and the other three ICD–10–PCS codes, 0H00MZ (Insertion of neurostimulator lead into brain, open approach), 0H03MZ (Insertion of neurostimulator lead into brain, percutaneous approach), and 0H04MZ (Insertion of neurostimulator lead into brain, percutaneous endoscopic approach) describe the insertions of the leads, depending on the approach used. The combination of an ICD–10–PCS...
code capturing the use of the generator and another ICD–10–PCS code describing the specific approach used to insert the leads would capture the performance of the entire procedure.

The requestor stated that the RNS<sup>®</sup> neurostimulator received FDA premarket approval on November 14, 2013, and is the first and only FDA-approved device used to provide responsive stimulation directly to the seizure onset zone in the brain. The RNS<sup>®</sup> neurostimulator includes a cranially implanted programmable neurostimulator connected to one or two depth and/or subdural cortical strip leads that are surgically placed in or on the brain at the seizure focus. The neurostimulator and leads are typically implanted during a single acute inpatient hospital procedure at a Comprehensive Epilepsy Center (CEC). The implanted neurostimulator continuously monitors brain electrical activity and is programmed by a physician to detect abnormal patterns of electrical activity that the physician believes may lead to seizures (epileptiform activity). In response to the detection of epileptiform activity, the device delivers brief, mild electrical pulses (responsive stimulation) to one or two epileptic foci. Detection and stimulation parameters are adjusted noninvasively by the physician to optimize control of epileptic seizures for each patient.

As the neurostimulator monitors brain activity, electrocorticograms (ECoGs) recorded immediately before and after certain events are stored for later review by the physician. The physician reviews the stored recordings to see the detections and the effects of stimulation. The physician can reprogram the neurostimulator at an in-office appointment to change detection and stimulation settings based on this information, as well as review the patient’s seizures.

The RNS<sup>®</sup> neurostimulator was approved for new technology add-on payments for FY 2015 and FY 2016, and new technology add-on payments were discontinued for FY 2017. The new technology add-on payment application was discussed in the FY 2015 IPPS/LTCH PPS proposed and final rules (79 FR 28051 through 28054 and 79 FR 49946 through 49950, respectively), the FY 2016 IPPS/LTCH PPS proposed and final rules (80 FR 24427 through 24448 and 80 FR 49442 through 49443, respectively), and the FY 2017 IPPS/LTCH PPS proposed and final rules (81 FR 25036 through 25037 and 81 FR 56882 through 56884, respectively).

The requestor suggested the following three options for MS–DRG assignment updates for cases involving the RNS<sup>®</sup> neurostimulator:

- Creating new MS–DRGs for cases involving the use of the RNS<sup>®</sup> neurostimulator. The requestor suggested MS–DRG XXX (Cranially Implanted Neurostimulators with MCC) and MS–DRG XXX (Cranially Implanted Neurostimulators without MCC) as possible MS–DRG titles. The requestor acknowledged that the number of cases assigned to this MS–DRG would be low, but anticipated that the number of cases would increase in the future.
- Reassign cases involving the use of the RNS<sup>®</sup> neurostimulator to MS–DRGs 020 and 021 (Intracranial Vascular Procedures with Principal Diagnosis of Hemorrhage with MCC, with CC, respectively) and update the MS–DRG logic and titles. The requestor asked CMS to reassign all cases involving the use of the RNS<sup>®</sup> neurostimulator that currently map to MS–DRG 023 (Craniotomy with Major Device Implant/Acute Complex CNS Principal Diagnosis without MCC) to MS–DRG 021, and change the title of MS–DRG 021 to “Intracranial Vascular Procedures with Principal Diagnosis of Hemorrhage with CC or Cranially Implanted Neurostimulator without MCC''. The requestor believed that the majority of cases involving the use of the RNS<sup>®</sup> neurostimulator that map to MS–DRG 024 do not include a secondary diagnosis that is classified as a CC, and the average cost of cases involving the use of the RNS<sup>®</sup> neurostimulator without a CC is significantly higher than the average cost of all cases in MS–DRG 022 (Intracranial Vascular Procedures with Principal Diagnosis of Hemorrhage without CC/MCC). Therefore, the requestor stated that it would not be adequate to assign cases involving the use of the RNS<sup>®</sup> neurostimulator without a CC to MS–DRG 022.
- Reassign cases involving the use of the RNS<sup>®</sup> neurostimulator to other higher paying MS–DRGs that would provide adequate payment.

The requestor stated that it had analyzed data from two sources, which demonstrated that the average cost of cases involving the use of the RNS<sup>®</sup> neurostimulator was higher than the average cost of all cases in MS–DRGs 023 and 024 (the current MS–DRGs for cases involving the use of the RNS<sup>®</sup> neurostimulator). The requestor indicated that the data used for its analysis was obtained from hospitals performing the procedure, as well as from the FY 2015 MedPAR file.

The requestor also asked that CMS examine the cases representing cranially implanted neurostimulators and leads that were inserted for the treatment of epilepsy. The requestor pointed out that neurostimulators also are used in the treatment of movement disorders such as Parkinson’s disease, essential tremor, or dystonia. The requestor asked that CMS identify those cases with a principal diagnosis of epilepsy, and identified the following ICD–10–CM codes that it believed were representative of potential epilepsy cases.

<table>
<thead>
<tr>
<th>ICD–10–CM code</th>
<th>ICD–10–CM code title</th>
</tr>
</thead>
<tbody>
<tr>
<td>G40.001</td>
<td>Localization-related (focal) (partial) idiopathic epilepsy and epileptic syndromes with seizures of localized onset, not intractable, with status epilepticus.</td>
</tr>
<tr>
<td>G40.009</td>
<td>Localization-related (focal) (partial) idiopathic epilepsy and epileptic syndromes with seizures of localized onset, intractable, without status epilepticus.</td>
</tr>
<tr>
<td>G40.011</td>
<td>Localization-related (focal) (partial) idiopathic epilepsy and epileptic syndromes with seizures of localized onset, intractable, with status epilepticus.</td>
</tr>
<tr>
<td>G40.019</td>
<td>Localization-related (focal) (partial) idiopathic epilepsy and epileptic syndromes with seizures of localized onset, intractable, without status epilepticus.</td>
</tr>
<tr>
<td>G40.101</td>
<td>Localization-related (focal) (partial) symptomatic epilepsy and epileptic syndromes with simple partial seizures, not intractable, with status epilepticus.</td>
</tr>
<tr>
<td>G40.119</td>
<td>Localization-related (focal) (partial) symptomatic epilepsy and epileptic syndromes with simple partial seizures, intractable, without status epilepticus.</td>
</tr>
</tbody>
</table>
The following three ICD–10–PCS code combinations capture the use of the RNS generators are inserted into the skull.

MS–DRGs 023 and 024 contain a number of cases representing neurostimulator generator and lead code combinations that are captured under a list referred to as “Major Device Implant.” The neurostimulator generators on this list are inserted into the skull, as well as into the subcutaneous areas of the chest, back, or abdomen. The leads are all inserted into the brain. The RNS® neurostimulator generators are inserted into the skull and the leads are inserted into the brain. The following three ICD–10–PCS code combinations capture the use of the RNS® neurostimulator and leads that would determine an assignment of a case to MS–DRGs 023 and 024, as shown in the “Major Device Implant” list:

- **0NH00NZ** (Insertion of neurostimulator generator into skull, open approach), in combination with 00H00MZ (Insertion of neurostimulator lead into brain, open approach);
- **0NH00NZ** (Insertion of neurostimulator generator into skull, open approach), in combination with 00M03MZ (Insertion of neurostimulator lead into brain, percutaneous approach); and
- **0NH00NZ** (Insertion of neurostimulator generator into skull, open approach), in combination with 00H04MZ (Insertion of neurostimulator lead into brain, percutaneous endoscopic approach).

We examined claims data from the December 2016 update of the FY 2016 MedPAR file for all cases representing the use of a neurostimulator in MS–DRGs 023 and 024 listed under the “Major Device Implant” list. As requested, we also examined the cases represented by the three neurostimulator code combinations, which capture the use of the RNS® neurostimulator that are a subset of the cases listed on the “Major Device Implant” list using the code combinations listed above, and that had a principal diagnosis of epilepsy from the list supplied by the requestor. The following tables show our findings for those cases in MS–DRGs 023 and 024 as well as findings for cases in MS–DRGs 020 and 021.

### MS–DRGs 023 AND 024 [Neurostimulator Cases]

<table>
<thead>
<tr>
<th>MS–DRG</th>
<th>Number of cases</th>
<th>Average length of stay</th>
<th>Average costs</th>
</tr>
</thead>
<tbody>
<tr>
<td>MS–DRG 023—All cases</td>
<td>6,723</td>
<td>10.9</td>
<td>$39,014</td>
</tr>
<tr>
<td>MS–DRG 023—Cases with neurostimulators (Major Device Implant list cases)</td>
<td>21</td>
<td>6.7</td>
<td>48,821</td>
</tr>
<tr>
<td>MS–DRG 023—Cases with neurostimulator generators inserted into skull (includes cases involving the use of the RNS® neurostimulator) and cases with a principal diagnosis of epilepsy</td>
<td>7</td>
<td>8.0</td>
<td>63,365</td>
</tr>
<tr>
<td>MS–DRG 024—All cases</td>
<td>2,275</td>
<td>5.5</td>
<td>27,574</td>
</tr>
</tbody>
</table>
As shown by the table above, for MS–DRG 023, we identified a total of 6,723 cases, with an average length of stay of 10.9 days and average costs of $39,014. Of the 6,723 cases in MS–DRG 023, there were 21 cases representing the implantation of any type of neurostimulator generator with an average length of stay of 6.7 days, and average costs of $48,821. Of the 21 neurostimulator generator cases, there were 7 cases with the neurostimulator generators inserted into skull (including cases involving the use of the RNS© neurostimulator) and a principal diagnosis of epilepsy with an average length of stay of 4.3 days and average costs of $31,669. Of the 6,723 cases in MS–DRG 023, we identified a total of 394 cases representing the implantation of any type of neurostimulator generator with an average length of stay of 6.7 days and average costs of $39,014. Of the 394 neurostimulator generator cases, there were 54 cases with the neurostimulator generators inserted into skull (including cases involving the use of the RNS© neurostimulator) and a principal diagnosis of epilepsy with an average length of stay of 4.3 days and average costs of $31,669. The RNS© neurostimulator generators are not used to treat patients with diagnosis of a hemorrhage. Therefore, our clinical advisors stated that it was inappropriate to reassign cases representing a principal diagnosis of epilepsy to an MS–DRG that contains cases that represent the treatment of intracranial hemorrhage. They also stated that the differences in average length of stay and average costs support this recommendation.

We then explored alternative MS–DRG assignments, as was requested. We noted that the 7 cases with the neurostimulator generators inserted into the skull (including cases involving the use of the RNS© neurostimulator) and a principal diagnosis of epilepsy to MS–DRG 020. While these neurostimulator cases had average costs that were $24,351 higher than the average costs of all cases in MS–DRG 020, there were only a total of 7 cases. There may have been other factors contributing to the higher costs. We noted that the 54 cases with the neurostimulator generators inserted into skull (including cases involving the use of the RNS© neurostimulator) and a principal diagnosis of epilepsy to an MS–DRG that contains cases that represent the treatment of intracranial hemorrhage. They also stated that the differences in average length of stay and average costs support this recommendation.
present, the cases would receive higher payment. The average costs of MS–DRG 023 were $39,014, compared to the average costs of $51,041 for the cases with the neurostimulator generators inserted into skull (including cases involving the use of the RNS© neurostimulator) and a principal diagnosis of epilepsy in MS–DRG 024. Our clinical advisors reviewed the clinical issues and the claims data, and supported the recommendation to reassign the cases with the neurostimulator generators inserted into skull (including cases involving the use of the RNS© neurostimulator) and a principal diagnosis of epilepsy to MS–DRG 023, even if there is no MCC reported. Therefore, we are proposing to reassign all cases with a principal diagnosis of epilepsy from the epilepsy diagnosis list provided earlier, and one of the following ICD–10–PCS code combinations capturing cases with the neurostimulator generators inserted into the skull (including cases involving the use of the RNS© neurostimulator), to MS–DRG 023, even if there is no MCC reported:

- 0NH00NZ (Insertion of neurostimulator generator into skull, open approach), in combination with 00H00MZ (Insertion of neurostimulator lead into brain, open approach);
- 0NH00NZ (Insertion of neurostimulator generator into skull, open approach), in combination with 00H03MZ (Insertion of neurostimulator lead into brain, percutaneous approach); and
- 0NH00NZ (Insertion of neurostimulator generator into skull, open approach), in combination with 00H04MZ (Insertion of neurostimulator lead into brain, percutaneous endoscopic approach).

We also are proposing to change the title of MS–DRG 023 from “Craniotomy with Major Device Implant or Acute Complex Central Nervous System (CNS) Principal Diagnosis (PDX) with MCC or Chemotherapy Implant or Epilepsy with Neurostimulator”) to “Craniotomy with Major Device Implant or Acute Complex Central Nervous System (CNS) Principal Diagnosis (PDX) with MCC or Chemotherapy Implant or Epilepsy with Neurostimulator” to reflect the proposed modifications to MS–DRG assignments.

We are inviting public comments on our proposals.

We received a request to add the ICD–10–CM diagnosis codes currently assigned to MS–DRGs 067 and 068 (Nonspecific CVA and Precerebral Occlusion without Infarction with MCC and without MCC, respectively) and the ICD–10–CM diagnosis codes currently assigned to MS–DRG 069 (Transient Ischemia) to the GROUPER logic for MS–DRGs 061, 062, and 063 (Acute Ischemic Stroke with Use of Thrombolytic Agent with MCC, with CC, and without CC/MCC, respectively) when those conditions are sequenced as the principal diagnosis and reported with an ICD–10–PCS procedure code describing use of a thrombolytic agent (for example, tPA).

The ICD–10–CM diagnosis codes displayed in the table below identify the conditions that are assigned to MS–DRGs 067 and 068 when reported as a principal diagnosis.

<table>
<thead>
<tr>
<th>ICD–10–CM code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>I65.01</td>
<td>Occlusion and stenosis of right vertebral artery.</td>
</tr>
<tr>
<td>I65.02</td>
<td>Occlusion and stenosis of left vertebral artery.</td>
</tr>
<tr>
<td>I65.03</td>
<td>Occlusion and stenosis of bilateral vertebral arteries.</td>
</tr>
<tr>
<td>I65.09</td>
<td>Occlusion and stenosis of unspecified vertebral artery.</td>
</tr>
<tr>
<td>I65.1</td>
<td>Occlusion and stenosis of basilar artery.</td>
</tr>
<tr>
<td>I65.21</td>
<td>Occlusion and stenosis of right carotid artery.</td>
</tr>
<tr>
<td>I65.22</td>
<td>Occlusion and stenosis of left carotid artery.</td>
</tr>
<tr>
<td>I65.23</td>
<td>Occlusion and stenosis of bilateral carotid arteries.</td>
</tr>
<tr>
<td>I65.29</td>
<td>Occlusion and stenosis of unspecified carotid artery.</td>
</tr>
<tr>
<td>I65.8</td>
<td>Occlusion and stenosis of other precerebral arteries.</td>
</tr>
<tr>
<td>I65.9</td>
<td>Occlusion and stenosis of unspecified precerebral artery.</td>
</tr>
<tr>
<td>I66.01</td>
<td>Occlusion and stenosis of right middle cerebral artery.</td>
</tr>
<tr>
<td>I66.02</td>
<td>Occlusion and stenosis of left middle cerebral artery.</td>
</tr>
<tr>
<td>I66.03</td>
<td>Occlusion and stenosis of bilateral middle cerebral arteries.</td>
</tr>
<tr>
<td>I66.09</td>
<td>Occlusion and stenosis of unspecified middle cerebral artery.</td>
</tr>
<tr>
<td>I66.11</td>
<td>Occlusion and stenosis of right anterior cerebral artery.</td>
</tr>
<tr>
<td>I66.12</td>
<td>Occlusion and stenosis of left anterior cerebral artery.</td>
</tr>
<tr>
<td>I66.13</td>
<td>Occlusion and stenosis of bilateral anterior cerebral arteries.</td>
</tr>
<tr>
<td>I66.19</td>
<td>Occlusion and stenosis of unspecified anterior cerebral artery.</td>
</tr>
<tr>
<td>I66.21</td>
<td>Occlusion and stenosis of right posterior cerebral artery.</td>
</tr>
<tr>
<td>I66.22</td>
<td>Occlusion and stenosis of left posterior cerebral artery.</td>
</tr>
<tr>
<td>I66.23</td>
<td>Occlusion and stenosis of bilateral posterior cerebral arteries.</td>
</tr>
<tr>
<td>I66.29</td>
<td>Occlusion and stenosis of unspecified posterior cerebral artery.</td>
</tr>
<tr>
<td>I66.3</td>
<td>Occlusion and stenosis of cerebellar arteries.</td>
</tr>
<tr>
<td>I66.8</td>
<td>Occlusion and stenosis of other cerebral arteries.</td>
</tr>
<tr>
<td>I66.9</td>
<td>Occlusion and stenosis of unspecified cerebral artery.</td>
</tr>
</tbody>
</table>

The ICD–10–CM diagnosis codes displayed in the table below identify the conditions that are assigned to MS–DRG 069 when reported as a principal diagnosis.

<table>
<thead>
<tr>
<th>ICD–10–CM code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>G45.0</td>
<td>Vertebro-basilar artery syndrome.</td>
</tr>
<tr>
<td>G45.1</td>
<td>Carotid artery syndrome (hemispheric).</td>
</tr>
<tr>
<td>G45.2</td>
<td>Multiple and bilateral precerebral artery syndromes.</td>
</tr>
<tr>
<td>G45.8</td>
<td>Other transient cerebral ischemic attacks and related syndromes.</td>
</tr>
<tr>
<td>G45.9</td>
<td>Transient cerebral ischemic attack, unspecified.</td>
</tr>
</tbody>
</table>
The ICD–10–PCS procedure codes displayed in the table below describe the use of a thrombolytic agent. These procedure codes are designated as non-O.R. procedure codes affecting the MS–DRG assignment for MS–DRGs 061, 062, and 063.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>3E03017</td>
<td>Introduction of other thrombolytic into peripheral vein, open approach.</td>
</tr>
<tr>
<td>3E03317</td>
<td>Introduction of other thrombolytic into peripheral vein, percutaneous approach.</td>
</tr>
<tr>
<td>3E04017</td>
<td>Introduction of other thrombolytic into central vein, open approach.</td>
</tr>
<tr>
<td>3E04317</td>
<td>Introduction of other thrombolytic into central vein, percutaneous approach.</td>
</tr>
<tr>
<td>3E05017</td>
<td>Introduction of other thrombolytic into peripheral artery, open approach.</td>
</tr>
<tr>
<td>3E05317</td>
<td>Introduction of other thrombolytic into peripheral artery, percutaneous approach.</td>
</tr>
<tr>
<td>3E06017</td>
<td>Introduction of other thrombolytic into central artery, open approach.</td>
</tr>
<tr>
<td>3E06317</td>
<td>Introduction of other thrombolytic into central artery, percutaneous approach.</td>
</tr>
<tr>
<td>3E08017</td>
<td>Introduction of other thrombolytic into heart, open approach.</td>
</tr>
<tr>
<td>3E08317</td>
<td>Introduction of other thrombolytic into heart, percutaneous approach.</td>
</tr>
</tbody>
</table>

At the onset of stroke symptoms, tPA must be given within 3 hours (or up to 4.5 hours for certain eligible patients) in an attempt to dissolve a clot and improve blood flow to the specific area affected in the brain. If, upon receiving the tPA, the stroke symptoms completely resolve within 24 hours and imaging studies (if performed) are negative, the patient has suffered what is clinically defined as a transient ischemic attack, not a stroke. According to the requestor, the current MS–DRG assignments do not account for this subset of patients who were successfully treated with tPA to prevent a stroke.

In addition, the requestor expressed concerns regarding documentation and quality of the data. For example, the requestor noted that the terms “stroke-in-evolution” and “aborted stroke” may be documented as a “workaround” for a patient exhibiting symptoms of a stroke who receives tPA and, regardless of the outcome, would result in assignment to MS–DRG 061, 062, or 063. Therefore, in cases where the patient’s stroke symptoms completely resolved upon receiving tPA and the patient clinically suffered a precerebral occlusion or transient ischemia, this documentation practice is incorrectly labeling these patients as having had a stroke and ultimately leading to inaccurate data.

We analyzed claims data from the December 2016 update of the FY 2016 MedPAR file for MS–DRGs 061, 062, and 063. Our findings are shown in the tables below.
We recognize that while the volume of cases for patients with a diagnosis of precerebral occlusion receiving tPA in MS–DRGs 067 and 068 is relatively low, the average length of stay is longer, and the average costs for this subset of patients is approximately twice the amount of the average costs in comparison to all cases in MS–DRGs 067 and 068. We then analyzed claims data for cases in MS–DRG 069 when reported with a procedure code describing the use of tPA. As shown in the table below, the total number of cases reported in MS–DRG 069 was 50,633, with an average length of stay of 2.5 days and average costs of $5,518. There were 554 cases of transient ischemia receiving tPA, with an average length of stay of 3.2 days and average costs of $12,481.

### MS–DRG FOR PRECEREBRAL OCCLUSION WITH USE OF THROMBOLYTIC AGENT—Continued

<table>
<thead>
<tr>
<th>MS–DRG</th>
<th>Number of cases</th>
<th>Average length of stay</th>
<th>Average costs</th>
</tr>
</thead>
<tbody>
<tr>
<td>MS–DRG 068—Cases with tPA</td>
<td>33</td>
<td>4.3</td>
<td>13,814</td>
</tr>
</tbody>
</table>

Similar to the findings for MS–DRGs 067 and 068, the number of cases for transient ischemia receiving tPA in MS–DRG 069 was relatively low in comparison to all the cases in the MS–DRG, with a longer average length of stay and approximately twice the amount of average costs in comparison to all cases in MS–DRG 069.

The results of analysis of the data and the advice of our clinical advisors support adding the ICD–10–CM diagnosis codes in MS–DRGs 067, 068, and 069 to the list of principal diagnoses in MS–DRGs 061, 062, and 063 to better account for this subset of patients who were successfully treated with tPA to prevent a stroke, to identify the increasing use of thrombolytics at the onset of symptoms of a stroke, to further encourage appropriate physician documentation for a precerebral occlusion or transient ischemic attack when patients are treated with tPA, and to reflect more appropriate payment for the resources involved in evaluating and treating these patients. We believe this approach will improve accuracy of the data and assist in addressing the concern that facilities may be reporting incorrect diagnoses for this subset of patients.

Therefore, for FY 2018, we are proposing to add the ICD–10–CM diagnosis codes listed earlier in this section that are currently assigned to MS–DRGs 067 and 068 and the ICD–10–CM diagnosis codes currently assigned to MS–DRG 069 to the GROUPER logic for MS–DRGs 061, 062, and 063 when those conditions are sequenced as the principal diagnosis and reported with an ICD–10–PCS procedure code describing use of a thrombolytic agent (for example, tPA). We are inviting public comments on our proposal.

We also are proposing to retitle MS–DRGs 061, 062, and 063 as “Ischemic Stroke, Precerebral Occlusion or Transient Ischemia with Thrombolytic Agent with MCC, without CC/MCC”, respectively, and to retitle MS–DRG 069 as “Transient Ischemia without Thrombolytic”. We are inviting public comments on our proposals.

3. MDC 2 (Diseases and Disorders of the Eye: Swallowing Eye Drops (Tetrahydrozoline))

We received a request to reassign the following ICD–10–CM diagnosis codes that capture swallowing eye drops from MS–DRGs 124 and 125 (Other Disorders of the Eye with and without MCC, respectively) to MS–DRGs 917 and 918 (Poisoning and Toxic Effects of Drugs with and without MCC, respectively).

The requestor described a case where a patient was treated following swallowing eye drops, specifically Tetrahydrozoline, which the provider considers to be a poisoning, not a disorder of the eye.

- T49.5X1A (Poisoning by ophthalmological drugs and preparations, accidental (unintentional), initial encounter);
- T49.5X2A (Poisoning by ophthalmological drugs and preparations, intentional self-harm, initial encounter);
- T49.5X3A (Poisoning by ophthalmological drugs and preparations, assault, initial encounter); and
- T49.5X4A (Poisoning by ophthalmological drugs and preparations, undetermined, initial encounter).

We agree with the requestor that the four diagnosis codes describe a poisoning, not a disorder of the eye. We examined claims data for cases in MS–DRGs 124 and 125 from the December 2016 update of the FY 2016 MedPAR file. Our findings are shown in the table below.

### MS–DRG FOR TRANSIENT ISCHEMIA WITH USE OF THROMBOLYTIC AGENT

<table>
<thead>
<tr>
<th>MS–DRG</th>
<th>Number of cases</th>
<th>Average length of stay</th>
<th>Average costs</th>
</tr>
</thead>
<tbody>
<tr>
<td>MS–DRG 069—All cases</td>
<td>50,633</td>
<td>2.5</td>
<td>$5,518</td>
</tr>
<tr>
<td>MS–DRG 069—Cases with tPA</td>
<td>554</td>
<td>3.2</td>
<td>12,481</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>MS–DRG 124 AND 125 CASES</th>
<th>Number of cases</th>
<th>Average length of stay</th>
<th>Average costs</th>
</tr>
</thead>
<tbody>
<tr>
<td>MS–DRG 124—All cases</td>
<td>874</td>
<td>4.8</td>
<td>$8,826</td>
</tr>
<tr>
<td>MS–DRG 124—Cases reporting poisoning by ophthalmological drugs and preparations code</td>
<td>1</td>
<td>2.0</td>
<td>3,007</td>
</tr>
<tr>
<td>MS–DRG 125—All cases</td>
<td>3,205</td>
<td>3.3</td>
<td>5,565</td>
</tr>
<tr>
<td>MS–DRG 125—Cases reporting poisoning by ophthalmological drugs and preparations code</td>
<td>1</td>
<td>2.0</td>
<td>1,446</td>
</tr>
</tbody>
</table>
As shown in the table above, there were only 2 cases of poisoning by ophthalmological drugs and preparations—1 case in MS–DRG 124 with an average length of stay of 2 days and average costs of $3,007 and 1 case in MS–DRG 125 with an average length of stay of 2 days and average costs of $1,446. The case of poisoning by ophthalmological drugs and preparations in MS–DRG 124 had a shorter average length of stay than the average length of stay for all cases in MS–DRG 124 (2.0 days compared to 4.8 days) and lower average costs than the average costs for all cases in MS–DRG 124 ($3,007 compared to $8,826). The case of poisoning by ophthalmological drugs and preparations in MS–DRG 125 also had a shorter average length of stay than the average length of stay for all cases in MS–DRG 125 (2.0 days compared to 3.3 days) and lower average costs than the average costs for all cases in MS–DRG 125 ($1,446 compared to $5,565).

We also examined claims data on cases reported in MS–DRGs 917 and 918 from the December 2016 update of the FY 2016 MedPAR file. Our findings are shown in the table below.

<table>
<thead>
<tr>
<th>MS–DRGs 917 AND 918 CASES</th>
<th>Number of cases</th>
<th>Average length of stay</th>
<th>Average costs</th>
</tr>
</thead>
<tbody>
<tr>
<td>MS–DRG 917—All cases</td>
<td>32,381</td>
<td>4.8</td>
<td>$9,882</td>
</tr>
<tr>
<td>MS–DRG 918—All cases</td>
<td>24,061</td>
<td>3.0</td>
<td>5,326</td>
</tr>
</tbody>
</table>

As shown in the table above, the 2 cases of poisoning by ophthalmological drugs and preparations also had shorter average lengths of stay than the average length of stay for all cases in MS–DRGs 917 and 918 (2.0 days compared to 4.8 days in MS–DRG 917 and 2.0 days compared to 3.0 days in MS–DRG 918). The average costs also were lower for the 2 cases of poisoning by ophthalmological drugs and preparations than the average costs for all cases in MS–DRGs 917 and 918 ($3,007 compared to $9,882 for all cases in MS–DRG 917 and $1,446 compared to $5,326 for all cases in MS–DRG 918). Therefore, cases with this type of poisoning had lower average lengths of stay and lower average costs than all other cases assigned to MS–DRGs 124 and 125 and cases in MS–DRGs 917 and 918 where poisonings are assigned.

Because the codes clearly capture a poisoning and not an eye disorder, we believe that these codes are more appropriately assigned to MS–DRGs 917 and 918 where other poisonings are assigned. Our clinical advisors also reviewed this issue and agreed that the codes should be moved from MS–DRGs 124 and 125 to MS–DRGs 917 and 918 because they clearly capture a poisoning and not a disorder of the eye. Because MS–DRGs 917 and 918 contain cases with multiple types of poisonings, it is expected that some types of poisoning cases will have longer lengths of stay and greater average costs than other types of poisoning cases. Therefore, we are proposing to reassign the following ICD–10–CM diagnosis codes from MS–DRGs 124 and 125 to MS–DRGs 917 and 918 for FY 2018: T49.5X1A; T49.5X2A; T49.5X3A; and T49.5X4A.

We are inviting public comments on our proposal.

4. MDC 5 (Diseases and Disorders of the Circulatory System)
   a. Percutaneous Cardiovascular Procedures and Insertion of a Radioactive Element
      Currently, under ICD–10–PCS, the logic for MS–DRG 246 (Percutaneous Cardiovascular Procedures with Drug-Eluting Stent with MCC or 4+ Vessels or Stents), MS–DRG 247 (Percutaneous Cardiovascular Procedures with Drug-Eluting Stent without MCC), MS–DRG 248 (Percutaneous Cardiovascular Procedures with Non-Drug-Eluting Stent with MCC or 4+ Vessels or Stents), and MS–DRG 249 (Percutaneous Cardiovascular Procedures with Non-Drug-Eluting Stent without MCC) includes six procedure codes that describe the insertion of a radioactive element. When any of these six procedure codes are reported without the reporting of a percutaneous cardiovascular procedure code, they are assigned to MS–DRG 264 (Other Circulatory System O.R. Procedures). The six specific procedure codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0WHC01Z</td>
<td>Insertion of radioactive element into mediastinum, open approach.</td>
</tr>
<tr>
<td>0WHC31Z</td>
<td>Insertion of radioactive element into mediastinum, percutaneous approach.</td>
</tr>
<tr>
<td>0WHC41Z</td>
<td>Insertion of radioactive element into mediastinum, percutaneous endoscopic approach.</td>
</tr>
<tr>
<td>0WHD01Z</td>
<td>Insertion of radioactive element into pericardial cavity, open approach.</td>
</tr>
<tr>
<td>0WHD31Z</td>
<td>Insertion of radioactive element into pericardial cavity, percutaneous approach.</td>
</tr>
<tr>
<td>0WHD41Z</td>
<td>Insertion of radioactive element into pericardial cavity, percutaneous endoscopic approach.</td>
</tr>
</tbody>
</table>

Unlike procedures involving the insertion of stents, none of the procedures described by the procedure codes listed above are performed in conjunction with a percutaneous cardiovascular procedure, and two of the six procedures described by these procedure codes (ICD–10–PCS codes 0WHC01Z and 0WHD01Z) are not performed using a percutaneous approach, but rather describe an open approach to performing the specific procedure. Our clinical advisors agreed that these procedures should not be used to classify cases within MS–DRGs 246 through 249 because they are not performed in conjunction with a percutaneous cardiovascular procedure. Furthermore, the indications for the insertion of a radioactive element typically involve a diagnosis of cancer, whereas the indications for the insertion of a coronary artery stent typically involve a diagnosis of coronary artery disease.

We conducted an analysis for the six procedures described by these procedure codes by reviewing the claims data for MS–DRGs 246 through 249 from the December 2016 update of
b. Proposed Modification of the Titles for MS–DRG 246 (Percutaneous Cardiovascular Procedures With Drug-Eluting Stent With MCC or 4+ Vessels or Stents) and MS–DRG 248 (Percutaneous Cardiovascular Procedures With Non-Drug-Eluting Stent With MCC or 4+ Vessels or Stents)

We are proposing to revise the titles for MS–DRGs 246 (Percutaneous Cardiovascular Procedures with Drug-Eluting Stent with MCC or 4+ Vessels or Stents) and MS–DRG 248 (Percutaneous Cardiovascular Procedures with Non-Drug-Eluting Stent with MCC or 4+ Vessels or Stents) to better reflect the ICD–10–PCS terminology of “arteries” versus “vessels” as used in the procedure code titles within the classification. Specifically, we are proposing to revise the title of MS–DRG 246 to “Percutaneous Cardiovascular Procedures with Drug-Eluting Stent with MCC or 4+ Arteries or Stents”. We are proposing to revise the title of MS–DRG 248 to “Percutaneous Cardiovascular Procedures with Non-Drug-Eluting Stent with MCC or 4+ Arteries or Stents”. We are inviting public comments on our proposals.

c. Transcatheter Aortic Valve Replacement (TAVR) and Left Atrial Appendage Closure (LAAC)

We received a request to create new MS–DRGs for cases involving transcatheter aortic valve replacement (TAVR) and left atrial appendage closure (LAAC) procedures when performed in combination in the same operative episode. The requestor stated that there are both clinical and financial advantages for the patient when performing concomitant procedures. For example, the requestor indicated that the clinical advantages for the patient may include single exposure to anesthesia and a reduction in overall procedure time, while the financial advantages may include lower cost-sharing. The requestor further believed that a single hospitalization for these concomitant procedures could be cost-effective for various providers and payers.

TAVR is indicated and approved as a treatment option for patients diagnosed with symptomatic aortic stenosis who are not surgical candidates for traditional open surgical techniques. Cases involving TAVR procedures are assigned to MS–DRGs 266 and 267 (Endovascular Cardiac Valve Replacement with MCC and without MCC, respectively), and are identified by the following ICD–10–PCS procedure codes shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>02RF37Z ..........</td>
<td>Replacement of aortic valve with autologous tissue substitute, percutaneous approach.</td>
</tr>
<tr>
<td>02RF38Z ..........</td>
<td>Replacement of aortic valve with synthetic substitute, percutaneous approach.</td>
</tr>
<tr>
<td>02RF39Z ..........</td>
<td>Replacement of aortic valve with nonautologous tissue substitute, percutaneous approach.</td>
</tr>
<tr>
<td>02RF37H ..........</td>
<td>Replacement of aortic valve with autologous tissue substitute, transapical, percutaneous approach.</td>
</tr>
<tr>
<td>02RF38H ..........</td>
<td>Replacement of aortic valve with synthetic substitute, transapical, percutaneous approach.</td>
</tr>
<tr>
<td>02RF39H ..........</td>
<td>Replacement of aortic valve with nonautologous tissue substitute, transapical, percutaneous approach.</td>
</tr>
</tbody>
</table>

LAAC is indicated and approved as a treatment option for patients diagnosed with atrial fibrillation. Cases involving LAAC procedures are assigned to MS–DRGs 273 and 274 (Percutaneous Intracardiac Procedures with MCC and without MCC, respectively), and are identified by ICD–10–PCS procedure code 02L73DK (Occlusion of left atrial appendage with intraluminal device, percutaneous approach).

The requestor suggested that the structure of the possible new MS–DRGs for TAVR procedures performed in combination with LAAC procedures could be modeled similar to the structure of MS–DRGs 266 and 267. While contemplating creation of the new MS–DRGs, the requestor asked CMS to also consider subdividing the possible new MS–DRGs into two severity levels and title them as follows:

- Suggested MS–DRG 26x (Endovascular Cardiac Valve Replacement with LAAC with MCC);
- Suggested MS–DRG 26x (Endovascular Cardiac Valve Replacement with LAAC without MCC).

We analyzed claims data from the December 2016 update of the FY 2016 MedPAR file for MS–DRGs 266 and 267 and identified the cases reporting TAVR procedures with and without an LAAC procedure. As shown in the table below, the data findings show that the total number of cases reported in MS–DRG 266 was 9,949, with an average length of stay of 7.2 days and average costs of $56,762. There were 9,872 cases involving a TAVR procedure, with an average length of stay of 7.2 days and average costs of $56,628. There was only one case identified in MS–DRG 266 where both a TAVR and an LAAC procedure were reported. This case had an average length of stay of 21.0 days and average costs of $60,226. For MS–DRG 267, the total number of cases found was 13,245, with an average length of stay of 3.5 days and average costs of $45,297. There were 13,245 cases involving a TAVR procedure, with an average length of stay of 3.5 days and average costs of $45,302. There were no cases identified in MS–DRG 267 where both a TAVR and an LAAC procedure were reported.
We then analyzed claims data in MS–DRGs 273 and 274 for cases reporting an LAAC procedure. As shown in the table below, the data findings show that the total number of cases reported in MS–DRG 273 was 6,541, with an average length of stay of 7.7 days and average costs of $20,267. There were 179 cases involving an LAAC procedure, with an average length of stay of 3.0 days and average costs of $26,042. For MS–DRG 274, the total number of cases found was 14,441, with an average length of stay of 3.0 days and average costs of $20,267. There were 2,428 cases involving an LAAC procedure, with an average length of stay of 1.2 days and average costs of $26,213.

### MS–DRGs for LAAC Procedures

<table>
<thead>
<tr>
<th>MS–DRG</th>
<th>Number of cases</th>
<th>Average length of stay</th>
<th>Average costs</th>
</tr>
</thead>
<tbody>
<tr>
<td>MS–DRG 273—All cases</td>
<td>6,541</td>
<td>7.7</td>
<td>$26,042</td>
</tr>
<tr>
<td>MS–DRG 273—Cases with LAAC</td>
<td>179</td>
<td>3.6</td>
<td>30,131</td>
</tr>
<tr>
<td>MS–DRG 274—All cases</td>
<td>14,441</td>
<td>3.0</td>
<td>20,267</td>
</tr>
<tr>
<td>MS–DRG 274—Cases with LAAC</td>
<td>2,428</td>
<td>1.2</td>
<td>26,213</td>
</tr>
</tbody>
</table>

The analysis of claims data for MS–DRGs 266, 267, 273, and 274 and input from our clinical advisors do not support creating new MS–DRGs for TAVR and LAAC procedures when performed in combination in the same operative episode. We found only one case in MS–DRG 266 where both a TAVR and an LAAC procedure were performed, and the claims data for cases reporting an LAAC procedure in MS–DRGs 273 and 274 support their current assignment. Our clinical advisors agreed the current MS–DRG assignments are appropriate for each respective procedure.

Therefore, we are not proposing to create new MS–DRGs for cases involving TAVR and LAAC procedures when performed in combination in the same operative episode. We are inviting public comments on our proposal to maintain the current MS–DRG structure for TAVR procedures in MS–DRGs 266 and 267, as well as the current MS–DRG structure for LAAC procedures in MS–DRGs 273 and 274.

d. Percutaneous Mitral Valve Replacement Procedures

We received a request to reassign four ICD–10–PCS procedure codes that describe percutaneous mitral valve replacement procedures from MS–DRGs 216 through 221 (Cardiac Valve and Other Major Cardiothoracic Procedures with and without Cardiac Catheterization with MCC, with CC and without CC/MCC, respectively) to MS–DRGs 266 and 267 (Endovascular Cardiac Valve Replacement with MCC and without MCC, respectively). The requestor indicated that there are inconsistencies in the current GROUPER logic for endovascular cardiac valve replacement procedures. Specifically, the requestor stated that the procedure codes that describe both the percutaneous approach and the transapical, percutaneous approach for the aortic and pulmonary valves are included in MS–DRGs 266 and 267. However, for the mitral valve, the GROUPER logic only includes the procedure codes that describe the transapical, percutaneous approach.

The requestor also stated that when MS–DRGs 266 and 267 were created, the intent was to include percutaneous replacement procedures for all cardiac valves. Therefore, the requestor recommended that CMS reassign the four ICD–10–PCS procedure codes shown in the table below that describe mitral valve replacement procedures, performed with the percutaneous approach from MS–DRGs 216 through 221 to MS–DRGs 266 and 267 to more appropriately group these procedures within the MS–DRG structure.

<table>
<thead>
<tr>
<th>ICD–10–PCS procedure code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>02RG37Z</td>
<td>Replacement of mitral valve with autologous tissue substitute, percutaneous approach.</td>
</tr>
<tr>
<td>02RG38Z</td>
<td>Replacement of mitral valve with zooplastic tissue, percutaneous approach.</td>
</tr>
<tr>
<td>02RG39Z</td>
<td>Replacement of mitral valve with synthetic substitute, percutaneous approach.</td>
</tr>
<tr>
<td>02RG9KZ</td>
<td>Replacement of mitral valve with nonautologous tissue substitute, percutaneous approach.</td>
</tr>
</tbody>
</table>

We agree with the requestor regarding the intent of the creation of MS–DRGs 266 and 267. As discussed in the FY 2015 IPPS/LTCH PPS final rule (79 FR 49890 through 49893), MS–DRGs 266 and 267 were created to uniquely classify the subset of high-risk cases representing patients who undergo a cardiac valve replacement procedure.
performed by a percutaneous (endovascular) approach. As such, we agree that all cardiac valve replacement procedures should be grouped within the same MS–DRG. In FY 2015, under the ICD–9–CM classification, there was not a specific procedure code for a percutaneous mitral valve replacement procedure. Therefore, when we converted from the ICD–9 based MS–DRGs to the ICD–10 MS–DRGs, there was not a code available from which to replicate. We refer the reader to the FY 2015 IPPS/LTCH PPS final rule (79 FR 49890 through 49893) for a detailed discussion on the initial request to create new MS–DRGs for endovascular cardiac valve replacement procedures, as well as the FY 2016 IPPS/LTCH PPS final rule (80 FR 49354 through 49358) and the FY 2017 IPPS/LTCH PPS final rule (81 FR 56787 through 56790) for a detailed discussion of the conversion to ICD–10 MS–DRGs, including our analysis of claims data and the need to accurately replicate the ICD–9–CM based MS–DRGs.

The requestor also noted that a proposal was discussed at the September 13–14, 2016 ICD–10 Coordination and Maintenance Committee meeting involving the creation of procedure codes that describe percutaneous tricuspid valve replacement procedures and, if finalized, these new procedure codes would also be assigned to MS–DRGs 266 and 267.

As shown in the table below and in Table 6B—New Procedure Codes, which is associated with this proposed rule and available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html, there are eight new procedure codes that describe tricuspid valve replacement procedures performed with percutaneous and transapical types of percutaneous approaches that will be effective October 1, 2017.

<table>
<thead>
<tr>
<th>ICD–10–PCS procedure code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>02RJ37H</td>
<td>Replacement of tricuspid valve with autologous tissue substitute, transapical, percutaneous approach.</td>
</tr>
<tr>
<td>02RJ37Z</td>
<td>Replacement of tricuspid valve with autologous tissue substitute, percutaneous approach.</td>
</tr>
<tr>
<td>02RJ38H</td>
<td>Replacement of tricuspid valve with zooplastic tissue, transapical, percutaneous approach.</td>
</tr>
<tr>
<td>02RJ38Z</td>
<td>Replacement of tricuspid valve with zooplastic tissue, percutaneous approach.</td>
</tr>
<tr>
<td>02RJ37H</td>
<td>Replacement of tricuspid valve with synthetic substitute, transapical, percutaneous approach.</td>
</tr>
<tr>
<td>02RJ37Z</td>
<td>Replacement of tricuspid valve with synthetic substitute, percutaneous approach.</td>
</tr>
<tr>
<td>02RJ3KZ</td>
<td>Replacement of tricuspid valve with nonautologous tissue substitute, percutaneous approach.</td>
</tr>
</tbody>
</table>

We agree with the requestor and believe that, in addition to the four procedure codes that describe the percutaneous mitral valve replacement procedures listed earlier in this section, the eight codes that describe percutaneous and transapical types of percutaneous tricuspid valve replacement procedures also should be grouped with the other endovascular cardiac valve replacement procedures. Therefore, we are proposing to reassign the four percutaneous mitral valve replacement procedures described by the procedure codes listed in the table above from MS–DRGs 216 through 221 to MS–DRGs 266 and 267. In addition, we are proposing to assign the eight new procedure codes (also listed in a separate table above) that describe percutaneous and transapical, percutaneous tricuspid valve replacement procedures to MS–DRGs 266 and 267.

We are inviting public comments on our proposals.

e. Percutaneous Tricuspid Valve Repair

We received a request to reassign cases reporting ICD–10–PCS procedure code 02UJ3JZ (Supplement tricuspid valve with synthetic substitute, percutaneous approach) from MS–DRGs 216 through 221 (Cardiac Valve and Other Major Cardiothoracic Procedures with and without Cardiac Catheterization with MCC, with CC and without CC/MCC, respectively) to MS–DRGs 228 and 229 (Other Cardiothoracic Procedures with MCC and without MCC, respectively). According to the requestor, reassigning cases involving these procedures would more appropriately cohesiveness with other clinically similar procedures, such as percutaneous mitral valve repair (for example, procedures involving the Mitraclip) described by procedure code 02UG3JZ (Supplement mitral valve with synthetic substitute, percutaneous approach), which are assigned to MS–DRGs 228 and 229.

The requestor noted that the FORMA Tricuspid Transcatheter Repair System (herein referred to as the FORMA system) is currently in clinical trials in the United States, Europe, and Canada, but has not received FDA approval. However, the FORMA system is presently available for compassionate use purposes. The FORMA system technology is indicated for use in the treatment of patients diagnosed with tricuspid regurgitation and occupies the regurgitant area of the affected valve, providing a surface for native leaflet coaptation. The requestor stated that the technology offers a viable alternative treatment using traditional tricuspid valve surgery. According to the requestor, the technology consists of a rail and a spacer, and the procedure to insert the device involves fluoroscopic imaging guidance.

We analyzed claims data from the December 2016 update of the FY 2016 MedPAR file for MS–DRGs 216 through 221 for cases reporting procedure code 02UJ3JZ (Supplement tricuspid valve with synthetic substitute, percutaneous approach). Our findings are shown in the following table.

<table>
<thead>
<tr>
<th>MS–DRG</th>
<th>Number of cases</th>
<th>Average length of stay</th>
<th>Average costs</th>
</tr>
</thead>
<tbody>
<tr>
<td>MS–DRG 216—All cases</td>
<td>9,139</td>
<td>14.4</td>
<td>$68,304</td>
</tr>
<tr>
<td>MS–DRG 216—Cases with percutaneous tricuspid valve repair</td>
<td>1</td>
<td>5.0</td>
<td>14,954</td>
</tr>
<tr>
<td>MS–DRG 217—All cases</td>
<td>3,536</td>
<td>8.9</td>
<td>45,857</td>
</tr>
<tr>
<td>MS–DRG 217—Cases with percutaneous tricuspid valve repair</td>
<td>1</td>
<td>3.0</td>
<td>16,234</td>
</tr>
</tbody>
</table>
We also analyzed claims data for MS–DRGs 228 and 229. Our findings are shown in the following table below.

### MS–DRGs for Other Cardiotoracic Procedures

<table>
<thead>
<tr>
<th>MS–DRG</th>
<th>Number of cases</th>
<th>Average length of stay</th>
<th>Average costs</th>
</tr>
</thead>
<tbody>
<tr>
<td>MS–DRG 228—All cases</td>
<td>3,466</td>
<td>9.8</td>
<td>$47,435</td>
</tr>
<tr>
<td>MS–DRG 229—All cases</td>
<td>4,553</td>
<td>4.9</td>
<td>$33,347</td>
</tr>
</tbody>
</table>

The claims data show that there were very few cases reported for performing a percutaneous tricuspid valve repair procedure in MS–DRGs 216 through 221. Of the 6 cases found in MS–DRG 219, with average costs of $58,075, the average cost of these cases aligned with the average cost of all cases in the MS–DRG assignment ($54,519). The data analysis and our clinical advisors do not support reassigning cases reporting procedure code 02UJ3JZ to MS–DRGs 228 and 229. The current MS–DRG assignment for percutaneous tricuspid valve repair procedures to MS–DRGs 216 through 221 is clinically coherent with the other percutaneous procedures performed on the heart valves that are currently assigned to these MS–DRGs. Percutaneous repair of the aortic, pulmonary and tricuspid valves utilizing various tissue substitutes (autologous, nonautologous, zooplastics, and synthetic) are assigned to MS–DRGs 216 through 221. The exception is the percutaneous mitral valve repair, which, as the requestor pointed out, is assigned to MS–DRGs 228 and 229 as discussed in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56809 through 56813). Our clinical advisors also agreed that the limited number of cases reported in MS–DRGs 216 through 221 does not warrant reassignment.

As a result of our review and the input from our clinical advisors, we are not proposing to reassign cases reporting procedure code 02UJ3JZ from MS–DRGs 216 through 221 to MS–DRGs 228 and 229.

We are inviting public comments on our proposal to maintain the current MS–DRG assignment for cases reporting procedure code 02UJ3JZ.

5. MDC 8 (Diseases and Disorders of the Musculoskeletal System and Connective Tissue)
   a. Total Ankle Replacement (TAR) Procedures

For FY 2018, we again received two requests for the reassignment of total ankle replacement (TAR) procedures to a different MS–DRG. TAR procedures are currently assigned to MS–DRGs 469 and 470 (Major Joint Replacement or Reattachment of Lower Extremity with and without MCC, respectively). This topic was discussed previously in the FY 2015 IPPS/LTCH PPS proposed and final rules (79 FR 28013 through 28015 and 79 FR 49896 through 49899, respectively) and in the FY 2017 IPPS/LTCH PPS proposed and final rules (81 FR 24989 through 24990 and 81 FR 56814 through 56816, respectively). For FY 2015 and FY 2017, we did not change the MS–DRG assignment for TAR procedures. The requestors indicated that TAR procedures are currently assigned to MS–DRGs 469 and 470, to which total hip replacement and total knee replacement procedures are also assigned. The requestors stated that there are significant clinical and cost differences among these procedures, which results in underpayment for TAR procedures. The requestors asked CMS to examine claims data for the following six ICD–10–PCS codes within MS–DRGs 469 and 470:

- 0SRF0J9 (Replacement of right ankle joint with synthetic substitute, cemented, open approach)
- 0SRF0JA (Replacement of right ankle joint with synthetic substitute, uncemented, open approach)
- 0SRF0JZ (Replacement of right ankle joint with synthetic substitute, open approach)
- 0SRG0J9 (Replacement of left ankle joint with synthetic substitute, cemented, open approach)
- 0SRG0JA (Replacement of left ankle joint with synthetic substitute, uncemented, open approach)
- 0SRG0JZ (Replacement of left ankle joint with synthetic substitute, open approach)

The requestors also stated that total ankle replacement is a complicated surgery that involves the replacement of the damaged parts of the three bones that comprise the ankle joint, as compared to the two bones in hip and knee replacement procedures. Furthermore, as the smallest weight-bearing large joint in the body, the requestors stated that TAR procedures demand a complexity of implant device design, engineering, and manufacture to exacting functional specifications that is vastly different from that of total hip and knee replacement devices. One of the requestors stated that the ankle region typically has poorer circulation and thinner soft tissue coverage than the
hip and knee, leading to a higher risk of wound complications and infection that may be more challenging and expensive to treat. In addition, this requestor stated that the unique anatomical characteristics and function of the ankle joint require a specialized surgical skill set, operative technique, and level of operating room resource utilization that is vastly dissimilar from that of total hip and knee replacement procedures.

We examined claims data from the December 2016 update of the FY 2016 MedPar file on reported cases of TAR procedures in MS–DRGs 469 and 470. Our findings are shown in the table below.

### Total Ankle Replacements Procedures

<table>
<thead>
<tr>
<th>MS–DRG</th>
<th>Number of cases</th>
<th>Average length of stay</th>
<th>Average costs</th>
</tr>
</thead>
<tbody>
<tr>
<td>MS–DRG 469—All cases</td>
<td>25,778</td>
<td>6.7</td>
<td>$22,139</td>
</tr>
<tr>
<td>MS–DRG 469—Cases reporting TAR procedure codes</td>
<td>31</td>
<td>4.6</td>
<td>23,828</td>
</tr>
<tr>
<td>MS–DRG 470—All cases</td>
<td>461,553</td>
<td>2.7</td>
<td>14,751</td>
</tr>
<tr>
<td>MS–DRG 470—Cases reporting TAR procedure codes</td>
<td>2,114</td>
<td>1.9</td>
<td>20,862</td>
</tr>
</tbody>
</table>

As shown in the table above, for MS–DRG 469, there were a total of 25,778 cases, with an average length of stay of 6.7 days and average costs of $22,139. Of the 25,778 cases in MS–DRG 469, there were 31 cases reporting a TAR procedure, with an average length of stay of 4.6 days and average costs of $23,828. For MS–DRG 470, there were a total of 461,553 cases, with an average length of stay of 2.7 days and average costs of $14,751. Of the 461,553 cases in MS–DRG 470, there were 2,114 cases reporting a TAR procedure, with an average length of stay of 1.9 days and average costs of $20,862. As mentioned earlier, there were only 31 TAR procedure cases in MS–DRG 469, and these cases had average costs of $1,689 higher than the average costs of all cases within MS–DRG 469. The relatively small number of cases may have been impacted by other factors. Several expensive cases could impact the average costs for a very small number of patients. We also note that the average length of stay for the TAR procedure cases was 4.6 days, as compared to 6.7 days for all cases within MS–DRG 469. The 2,114 TAR procedure cases in MS–DRG 470 had average costs that were $6,111 higher than the average costs of all cases in MS–DRG 470 ($20,862 compared to $14,751 for all cases). The data support reassigning all of the TAR procedures to MS–DRG 469, even when there is no MCC reported. However, the average costs of the TAR procedures in MS–DRG 470 are lower than the average costs for all cases in MS–DRG 469 ($20,862 compared to $22,139), the average costs are much closer to the average costs of TAR procedure cases in MS–DRG 470.

Our clinical advisors reviewed this clinical issue and the claims data, and agreed that it is clinically appropriate to reassign all of the TAR procedure cases from MS–DRG 470 to MS–DRG 469, even when there is no MCC reported. The claims data support the fact that these cases require more resources than other cases assigned to MS–DRG 470. Therefore, we are proposing to reassign the following TAR procedure codes, even if there is no MCC reported: 0SRF0J9; 0SRF0JA; 0SRF0JZ; 0SRG0J9; 0SRG0JA, and 0SRG0JZ, for FY 2018.

We are proposing to change the titles of MS–DRGs 469 and 470 to the following to reflect these proposed MS–DRG reassignments:

- Proposed retitle of MS–DRG 469: “Major Hip and Knee Joint Replacement or Reattachment of Lower Extremity with MCC or Total Ankle Replacement”;
- Proposed retitle of MS–DRG 470: “Major Hip and Knee Joint Replacement or Reattachment of Lower Extremity without MCC.”

We are inviting public comments on our proposals.

b. Revision of Total Ankle Replacement (TAR) Procedures

We received two requests to modify the MS–DRG assignment for revision of total ankle replacement (TAR) procedures, which are assigned to MS–DRGs 515, 516, and 517 (Other Musculoskeletal System and Connective Tissue O.R. Procedures with MCC, with CC, and without CC/MCC, respectively). This topic was discussed in the FY 2015 IPPS/LTCH PPS proposed and final rules (79 FR 28013 through 28015 and 79 FR 49896 through 49899, respectively) and in the FY 2017 IPPS/LTCH PPS proposed and final rules (81 FR 24992 through 24993 and 81 FR 56819 through 56820, respectively). For FY 2015 and FY 2017, we did not change the MS–DRG assignment for revision of TAR procedures.

The requestors asked that CMS examine the following eight ICD–10–PCS codes for revision of TAR procedures, which are assigned to MS–DRGs 515, 516, and 517:

- 0SWF0JZ (Revision of synthetic substitute in right ankle joint, open approach);
- 0SWF3JZ (Revision of synthetic substitute in right ankle joint, percutaneous approach);
- 0SWF4JZ (Revision of synthetic substitute in right ankle joint, percutaneous endoscopic approach);
- 0SWF5JZ (Revision of synthetic substitute in right ankle joint, external approach);
- 0SWG0JZ (Revision of synthetic substitute in left ankle joint, open approach);
- 0SWG3JZ (Revision of synthetic substitute in left ankle joint, percutaneous approach);
- 0SWG4JZ (Revision of synthetic substitute in left ankle joint, percutaneous endoscopic approach); and
- 0SWG5JZ (Revision of synthetic substitute in left ankle joint, external approach).

One requestor stated that these ICD–10–PCS codes more specifically identify the revision of TAR procedures than the prior ICD–9–CM codes. Specifically, ICD–9–CM code 81.59 (Revision of joint replacement of lower extremity, not elsewhere classified) was an unspecified code, which included toe and foot joint revision procedures in addition to revision of TAR procedures. The requestor stated that claims data reporting these ICD–10–PCS codes would allow CMS to better identify revisions of TAR procedures, and determine if the procedures are assigned to the appropriate MS–DRGs.

One requestor suggested the following three options for MS–DRG assignments:

- Assign the ICD–10–PCS ankle revision procedure codes to MS–DRGs 466, 467, and 468 (Revision of Hip or Knee Replacement with MCC, with CC, and without CC/MCC, respectively), and rename MS–DRGs 466, 467, and 468 as “Revision of Hip, Knee or Ankle with MCC, with CC, and without CC/MCC”, respectively;
• Assign the ICD–10–PCS ankle revision procedure codes to MS–DRG 469 (Major Joint Replacement or Reattachment of Lower Extremity with MCC) to more appropriately recognize higher hospital procedure costs associated with revision of TAR procedures; or

• Establish a new MS–D RG for the assignment of revision of TAR procedures.

The other requestor asked that CMS consider reassigning revision of TAR procedures to MS–DRGs that better address the cost-to-payment differential, such as MS–DRGs 466, 467, and 468.

We examined claims data from the December 2016 update of the FY 2016 MedPAR file on reported cases of revision of TAR procedures, as well as cases assigned to MS–DRGs 466, 467, 468, and MS–DRG 469. Our findings are shown in the tables below.

### Revisions of Joint Replacements Procedures

<table>
<thead>
<tr>
<th>MS–DRG</th>
<th>Number of cases</th>
<th>Average length of stay</th>
<th>Average costs</th>
</tr>
</thead>
<tbody>
<tr>
<td>MS–DRG 515—All cases</td>
<td>5,038</td>
<td>8.0</td>
<td>$20,562</td>
</tr>
<tr>
<td>MS–DRG 515—Cases reporting revision of total ankle replacement procedure codes</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>MS–DRG 516—All cases</td>
<td>13,276</td>
<td>4.8</td>
<td>13,524</td>
</tr>
<tr>
<td>MS–DRG 516—Cases reporting revision of total ankle replacement procedure codes</td>
<td>2</td>
<td>2.5</td>
<td>11,400</td>
</tr>
<tr>
<td>MS–DRG 517—All cases</td>
<td>13,330</td>
<td>2.8</td>
<td>10,003</td>
</tr>
<tr>
<td>MS–DRG 517—Cases reporting revision of total ankle replacement procedure codes</td>
<td>4</td>
<td>1.5</td>
<td>7,423</td>
</tr>
</tbody>
</table>

### Cases in MS–DRGs 466, 467, 468, and 469

<table>
<thead>
<tr>
<th>MS–DRG</th>
<th>Number of cases</th>
<th>Average length of stay</th>
<th>Average costs</th>
</tr>
</thead>
<tbody>
<tr>
<td>MS–DRG 466—All cases</td>
<td>3,886</td>
<td>8.4</td>
<td>$33,720</td>
</tr>
<tr>
<td>MS–DRG 467—All cases</td>
<td>19,145</td>
<td>4.2</td>
<td>24,609</td>
</tr>
<tr>
<td>MS–DRG 468—All cases</td>
<td>16,529</td>
<td>2.7</td>
<td>20,208</td>
</tr>
<tr>
<td>MS–DRG 469—All cases</td>
<td>25,778</td>
<td>6.7</td>
<td>22,139</td>
</tr>
</tbody>
</table>

As shown in the tables above, there were only 6 cases representing revisions of TAR procedures with no cases in MS–DRG 515, two cases in MS–DRG 516, and four cases in MS–DRG 517. The limited number of six cases does not justify the creation of a new MS–DRG for the assignment of revision of TAR procedures. Our data analysis demonstrates that the average length of stay for the revision of TAR procedures was lower than that for all cases in MS–DRG 516 (2.5 days compared to 4.8 days), and the average costs were lower ($11,400 compared to $13,524). The average length of stay for the revision of TAR procedures also was lower than that for all cases in MS–DRG 517 (1.5 days compared to 2.8 days), and the average costs were lower ($7,423 compared to $10,003). The data do not support reassigning the cases from MS–DRGs 515, 516, and 517.

Furthermore, the average length of stay and average costs of cases in MS–DRGs 466, 467, 468, and 469 are significantly higher than those for the revision of TAR procedures in MS–DRG 516 and 517. The average length of stay for all cases in MS–DRGs 466, 467, 468, and 469 is 8.4, 4.2, 2.7, and 6.7 days, respectively, compared to the average length of stay of 2.5 and 1.5 days for cases representing revision of TAR procedures in MS–DRGs 516 and 517, respectively. The average costs for all cases in MS–DRGs 466, 467, 468, and 469 are $33,720, $24,609, $20,208, and $22,139, respectively, compared to the average costs of $11,400 and $7,423 for cases representing revision of TAR procedures in MS–DRGs 516 and 517, respectively. Therefore, the data do not support reassigning the cases to MS–DRGs 466, 467, 468, or 469.

Our clinical advisors reviewed the clinical issue and the claims data and agreed that the revision of TAR procedures are appropriately assigned to MS–DRGs 515, 516, and 517, along with other procedures that describe revisions of joint replacements of the lower extremities, including the foot and toe. Our clinical advisors did not support reassigning these cases to MS–DRGs 466, 467, 468, or 469, or creating a new MS–DRG. Therefore, based on the findings of our analysis of claims data and the advice of our clinical advisors, we are proposing to maintain the current MS–DRG assignment for revision of TAR procedures within MS–DRGs 515, 516, and 517 for FY 2018.

We are inviting public comments on our proposal.

c. Magnetic Controlled Growth Rods (MAGEC® System)

We received a request to add six ICD–10–PCS procedure codes that describe the use of magnetically controlled growth rods for the treatment of early onset scoliosis (MAGEC® System) to MS–DRGs 456, 457, and 458 (Spinal Fusion Except Cervical with Spinal Curvature or Malignancy or Infection or Extensive Fusions with MCC, with CC or without CC/MCC, respectively). The MAGEC® System was discussed in the FY 2017 IPPS/LTCH PPS proposed rule (81 FR 25040 through 25042) and final rule (81 FR 56888 through 56891) as a new technology add-on payment application. The application was approved for FY 2017 new technology add-on payments, effective with discharges occurring on and after October 1, 2016. The request for new procedure codes to identify the MAGEC® System technology was discussed at the March 9–10, 2016 ICD–10 Coordination and Maintenance Committee meeting. Six new procedure codes were approved, effective October 1, 2016, and were displayed in Table 68.—New Procedure Codes associated with the FY 2017 IPPS/LTCH PPS final rule (which is available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/FY2017-IPPS-Final-Rule-Home-Page.html). These six procedure codes are currently assigned to MS–DRGs 518, 519, and 520 (Back and Neck Procedure Except Spinal Fusion with MCC or Disc Device/Neurostimulator, with CC, or without CC/MCC, respectively) and are shown in the table below.
According to the requestor, adding these six procedure codes will allow these cases to group to MS–DRGs that more accurately reflect the diagnosis of early onset scoliosis for which the MAGEC® System is indicated. In addition, the requestor stated that because this technology is utilized on a small subset of patients with approximately 300 cases per year, adding these procedure codes to MS–DRGs 456, 457, and 458 would have little impact.

Because these six procedure codes shown in the table above were effective as of October 1, 2016, there are no MedPAR claims data available to analyze. More importantly, we note that cases are assigned to MS–DRGs 456, 457, and 458 when an actual spinal fusion procedure is performed. Our clinical advisors agree that use of the MAGEC® System’s magnetically controlled growth rods technology alone does not constitute a spinal fusion.

Therefore, because there are no claims data available at this time and based on the advice of our clinical advisors, we are not proposing to add the six procedure codes to MS–DRGs 456, 457, or 458. If a spinal fusion procedure is performed along with the procedure to insert the MAGEC® System’s magnetically controlled growth rods, it would be appropriate to report that a spinal fusion was performed and the case would be assigned to one of the spinal fusion MS–DRGs.

We are inviting public comments on our proposal to maintain the current GROUPER logic for cases assigned to MS–DRGs 456, 457, and 458 and not add the six procedure codes describing the use of the MAGEC® System magnetically controlled growth rods. We also are inviting public comments on our proposal to maintain the assignment of the six procedure codes in MS–DRGs 518, 519, and 520.

The seven procedure codes currently included in the posterior spinal fusion list to MS–DRGs 453, 454, and 455 are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>XRG6092</td>
<td>Fusion of thoracic vertebral joint using nanotextured surface interbody fusion device, open approach, new technology group 2.</td>
</tr>
<tr>
<td>XRG7092</td>
<td>Fusion of 2 to 7 thoracic vertebral joints using nanotextured surface interbody fusion device, open approach, new technology group 2.</td>
</tr>
<tr>
<td>XRG8092</td>
<td>Fusion of 8 or more thoracic vertebral joints using nanotextured surface interbody fusion device, open approach, new technology group 2.</td>
</tr>
<tr>
<td>XRG9092</td>
<td>Fusion of thoracolumbar vertebral joint using nanotextured surface interbody fusion device, open approach, new technology group 2.</td>
</tr>
<tr>
<td>XRGB092</td>
<td>Fusion of lumbar vertebral joint using nanotextured surface interbody fusion device, open approach, new technology group 2.</td>
</tr>
<tr>
<td>XRG092</td>
<td>Fusion of 2 or more lumbar vertebral joints using nanotextured surface interbody fusion device, open approach, new technology group 2.</td>
</tr>
<tr>
<td>XRGDO92</td>
<td>Fusion of lumbosacral joint using nanotextured surface interbody fusion device, open approach, new technology group 2.</td>
</tr>
</tbody>
</table>

We note that the remaining three new procedure codes are accurately reflected in the anterior spinal fusion list; that is, ICD–10–PCS code XRG1092 (Fusion of cervical vertebral joint using nanotextured surface interbody fusion device, open approach, new technology group 2); ICD–10–PCS code XRG2092 (Fusion of 2 or more cervical vertebral joints using nanotextured surface interbody fusion device, open approach, new technology group 2); and ICD–10–PCS code XRG4092 (Fusion of cervicothoracic vertebral joint using nanotextured surface interbody fusion device, open approach, new technology group 2).

The seven procedure codes currently included in the posterior spinal fusion list describe an anterior spinal fusion by use of the interbody fusion device. In an interbody fusion, the anterior column of the spine is being fused. The results of our review of these procedure codes discussed below and the advice of our clinical advisors support moving the seven procedure codes from the posterior spinal fusion list to the anterior spinal fusion list in the GROUPER logic for MS–DRGs 453, 454, and 455. This will improve clinical accuracy and allow appropriate assignment to these MS–DRGs when both an anterior and posterior spinal fusion is performed.

During our review of the spinal fusion procedure codes using a nanotextured surface interbody fusion device in MS–DRGs 453, 454, and 455, we identified 149 additional procedure codes that should be moved from the posterior spinal fusion list to the anterior spinal fusion list.
We are proposing to delete these 33 procedure codes from MS–DRGs 453, 454, and 455 for FY 2018. We also note that some of the above listed codes also may be included in the logic for MS–DRGs 456, 457, and 458 (Spinal Fusion Fusions with MCC, with CC or without Exception Cervical with Spinal Curvature or Malignancy or Infection or Extensive Procedure codes listed in Table 6P.3a. associated with this proposed logic for MS–DRGs 453, 454, and 455 are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ORG00A1</td>
<td>Fusion of occipital-cervical joint with interbody fusion device, posterior approach, posterior column, open approach.</td>
</tr>
<tr>
<td>ORG03A1</td>
<td>Fusion of occipital-cervical joint with interbody fusion device, posterior approach, posterior column, percutaneous endoscopic approach.</td>
</tr>
<tr>
<td>ORG04A1</td>
<td>Fusion of occipital-cervical joint with interbody fusion device, posterior approach, posterior column, percutaneous approach.</td>
</tr>
<tr>
<td>ORG10A1</td>
<td>Fusion of cervical vertebral joint with interbody fusion device, posterior approach, posterior column, open approach.</td>
</tr>
<tr>
<td>ORG13A1</td>
<td>Fusion of cervical vertebral joint with interbody fusion device, posterior approach, posterior column, percutaneous approach.</td>
</tr>
<tr>
<td>ORG14A1</td>
<td>Fusion of cervical vertebral joint with interbody fusion device, posterior approach, posterior column, percutaneous endoscopic approach.</td>
</tr>
<tr>
<td>ORG20A1</td>
<td>Fusion of 2 or more cervical vertebral joints with interbody fusion device, posterior approach, posterior column, open approach.</td>
</tr>
<tr>
<td>ORG23A1</td>
<td>Fusion of 2 or more cervical vertebral joints with interbody fusion device, posterior approach, posterior column, percutaneous endoscopic approach.</td>
</tr>
<tr>
<td>ORG24A1</td>
<td>Fusion of 2 or more cervical vertebral joints with interbody fusion device, posterior approach, posterior column, percutaneous approach.</td>
</tr>
<tr>
<td>ORG40A1</td>
<td>Fusion of cervicothoracic vertebral joint with interbody fusion device, posterior approach, posterior column, open approach.</td>
</tr>
<tr>
<td>ORG43A1</td>
<td>Fusion of cervicothoracic vertebral joint with interbody fusion device, posterior approach, posterior column, percutaneous approach.</td>
</tr>
<tr>
<td>ORG44A1</td>
<td>Fusion of cervicothoracic vertebral joint with interbody fusion device, posterior approach, posterior column, percutaneous endoscopic approach.</td>
</tr>
<tr>
<td>ORG60A1</td>
<td>Fusion of thoracic vertebral joint with interbody fusion device, posterior approach, posterior column, open approach.</td>
</tr>
<tr>
<td>ORG63A1</td>
<td>Fusion of thoracic vertebral joint with interbody fusion device, posterior approach, posterior column, percutaneous endoscopic approach.</td>
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<tr>
<td>ORG64A1</td>
<td>Fusion of thoracic vertebral joint with interbody fusion device, posterior approach, posterior column, percutaneous approach.</td>
</tr>
<tr>
<td>ORG70A1</td>
<td>Fusion of 2 to 7 thoracic vertebral joints with interbody fusion device, posterior approach, posterior column, open approach.</td>
</tr>
<tr>
<td>ORG73A1</td>
<td>Fusion of 2 to 7 thoracic vertebral joints with interbody fusion device, posterior approach, posterior column, percutaneous endoscopic approach.</td>
</tr>
<tr>
<td>ORG74A1</td>
<td>Fusion of 2 to 7 thoracic vertebral joints with interbody fusion device, posterior approach, posterior column, percutaneous approach.</td>
</tr>
<tr>
<td>ORG80A1</td>
<td>Fusion of 8 or more thoracic vertebral joints with interbody fusion device, posterior approach, posterior column, open approach.</td>
</tr>
<tr>
<td>ORG83A1</td>
<td>Fusion of 8 or more thoracic vertebral joints with interbody fusion device, posterior approach, posterior column, percutaneous approach.</td>
</tr>
<tr>
<td>ORG84A1</td>
<td>Fusion of 8 or more thoracic vertebral joints with interbody fusion device, posterior approach, posterior column, percutaneous endoscopic approach.</td>
</tr>
<tr>
<td>ORG84A1</td>
<td>Fusion of thoracolumbar vertebral joint with interbody fusion device, posterior approach, posterior column, open approach.</td>
</tr>
<tr>
<td>ORG90A1</td>
<td>Fusion of thoracolumbar vertebral joint with interbody fusion device, posterior approach, posterior column, percutaneous approach.</td>
</tr>
<tr>
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<td>Fusion of thoracolumbar vertebral joint with interbody fusion device, posterior approach, posterior column, percutaneous endoscopic approach.</td>
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<td>Fusion of lumbar vertebral joint with interbody fusion device, posterior approach, posterior column, open approach.</td>
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<td>OS14A1</td>
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<tr>
<td>OS30A1</td>
<td>Fusion of lumbosacral joint with interbody fusion device, posterior approach, posterior column, open approach.</td>
</tr>
<tr>
<td>OS33A1</td>
<td>Fusion of lumbosacral joint with interbody fusion device, posterior approach, posterior column, percutaneous approach.</td>
</tr>
<tr>
<td>OS34A1</td>
<td>Fusion of lumbosacral joint with interbody fusion device, posterior approach, posterior column, percutaneous endoscopic approach.</td>
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We are proposing to delete these 33 procedure codes from MS–DRGs 453, 454, and 455 for FY 2018. We also note that some of the above listed codes also may be included in the logic for MS–DRGs 456, 457, and 458 (Spinal Fusion Fusions with MCC, with CC or without Exception Cervical with Spinal Curvature or Malignancy or Infection or Extensive Procedure codes listed in Table 6P.3a. associated with this proposed logic for MS–DRGs 453, 454, and 455 are shown in the table below.

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CC/MCC, respectively), MS–DRGs 459 and 460 (Spinal Fusion Except Cervical with MCC and without MCC, respectively), and MS–DRGs 471, 472, and 473 (Cervical Spinal Fusion with MCC, with CC and without CC/MCC, respectively). Therefore, we are proposing to delete the 33 procedure codes from the logic for those spinal fusion MS–DRGs as well. In addition, we are proposing to delete the 33 procedure codes from the ICD–10–PCS classification as shown in Table 6D.—Invalid Procedure Codes associated with this proposed rule (which is available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html).

In summary, we are inviting public comments on our proposal to move the seven procedure codes describing spinal fusion using a nanotextured surface interbody fusion device from the posterior spinal fusion list to the anterior spinal fusion list in the GROUPER logic for MS–DRGs 453, 454, and 455. We also are inviting public comments on our proposal to move the 149 procedure codes describing spinal fusion of the anterior column with a posterior approach from the posterior spinal fusion list to the anterior spinal fusion list in the GROUPER logic for MS–DRGs 453, 454, and 455. In addition, we are inviting public comments on our proposal to delete the 33 procedure codes describing spinal fusion of the posterior column with an interbody fusion device from MS–DRGs 453, 454, 455, 456, 457, 458, 459, 460, 471, 472, and 473, as well as from the ICD–10–PCS classification.

6. MDC 14 (Pregnancy, Childbirth and the Puerperium)

a. Vaginal Delivery and Complicating Diagnoses

In the FY 2017 IPPS/LTC PPS final rule (81 FR 56854), we noted that the code list as displayed in the ICD–10 MS–DRG Version 33 Definitions Manual for MS–DRG 774 (Vaginal Delivery with Complicating Diagnoses) required further analysis to clarify what constitutes a vaginal delivery to satisfy the ICD–10 MS–DRG logic. We stated our plans to conduct further analysis of the diagnosis code lists in MS–DRG 774 for FY 2018.

We believe that the Version 34 Definitions Manual and GROUPER logic for MS–DRG 774 continue to require additional analysis to determine how best to classify a vaginal delivery. For example, under MS–DRG 774, the Definitions Manual currently states that three conditions must be met, the first of which is a vaginal delivery. To satisfy this first condition, codes that describe conditions or circumstances from among three lists of codes must be reported. The first list is comprised of ICD–10–CM diagnosis codes that may be reported as a principal diagnosis or a secondary diagnosis. These diagnosis codes describe conditions in which it is assumed that a vaginal delivery has occurred. The second list of codes is a list of ICD–10–PCS procedure codes that also describe circumstances in which it is assumed that a vaginal delivery occurred. The third list of codes identifies diagnoses describing the outcome of the delivery. Therefore, if any code from one of those three lists is reported, the first condition (vaginal delivery) is considered to be met for assignment to MS–DRG 774.

Our continued concern with the first list of ICD–10–CM diagnosis codes as currently displayed in the Definitions Manual under the first condition is that all of the conditions necessarily reflect that a vaginal delivery occurred. Several of the diagnosis codes listed could also reflect that a cesarean delivery occurred. For example, ICD–10–CM diagnosis code O10.02 (Pre-existing essential hypertension complicating childbirth) does not specify that a vaginal delivery took place; yet it is included in the list of conditions that may be reported as a principal diagnosis or a secondary diagnosis in the GROUPER logic for a vaginal delivery. The reporting of this code also could be appropriate for a delivery that occurred by cesarean section.

As noted earlier, the second list of codes for the first condition are comprised of ICD–10–PCS procedure codes. While we agree that the current list of procedure codes in MS–DRG 774 may appropriately describe that a vaginal delivery occurred, we also believe this list could be improved and warrants closer review.

The third list of codes for the first condition in MS–DRG 774 includes conditions describing the outcome of the delivery that would be reported as secondary diagnoses. Similar to concerns with the first list of codes, we believe the conditions do not necessarily reflect that a vaginal delivery occurred because they also can be reported on claims where a cesarean delivery occurred.

For the second condition in MS–DRG 774 to be met, diagnosis codes that are identified as a complicating diagnosis from among two lists may be reported. The first list is comprised of ICD–10–CM diagnosis codes that may be reported as a principal or secondary diagnosis. The second list is comprised of ICD–10–CM diagnosis codes that may be reported as a secondary diagnosis. Currently, there is only one code listed under the secondary diagnosis list. We have concerns with these lists and what is classified as a complicating diagnosis when reviewing the code lists for this and other MS–DRGs that use that logic in MDC 14.

For the third condition in MS–DRG 774 to be met, a limited set of O.R. procedures, including both extensive and nonextensive procedures, are listed. We have concerns with this third condition as being needed to satisfy the logic for a vaginal delivery MS–DRG.

In summary, the MS–DRG logic involving a vaginal delivery under MDC 14 is technically complex as a result of the requirements that must be met to satisfy assignment to the affected MS–DRGs. Upon review and discussion, our clinical advisors recommended, and we agree, that we should solicit public comments on further refinement to the following four MS–DRGs related to vaginal delivery: MS–DRG 767 (Vaginal Delivery with Sterilization and/or D&C); MS–DRG 768 (Vaginal Delivery with O.R. Procedure Except Sterilization and/or D&C); MS–DRG 774 (Vaginal Delivery with Complicating Diagnosis); and MS–DRG 775 (Vaginal Delivery without Complicating Diagnosis).

In addition, our clinical advisors agreed that we should solicit public comments on further refinement to the conditions defined as a complicating diagnosis in MS–DRG 774 and MS–DRG 781 (Other Antepartum Diagnoses with Medical Complications).

Therefore, we are soliciting public comments on which diagnosis or procedure codes, or both, should be considered in the logic to identify a vaginal delivery and which diagnosis codes should be considered in the logic to identify a complicating diagnosis. As MS–DRGs 767, 768, 774, 775, and 781 incorporate one or both aspects (vaginal delivery or complicating diagnosis), public comments that we receive from this solicitation will be helpful in determining what proposed revisions to the current logic should be made. We will review public comments received in response to this solicitation as we continue to evaluate these areas under MDC 14 and, if warranted, we would propose refinements for FY 2019. We are requesting that all comments be directed to the CMS MS–DRG Classification Change Request Mailbox located at: MSDRGCClassificationChange@cms.hhs.gov by November 1, 2017.
b. MS–DRG 998 (Principal Diagnosis Invalid as Discharge Diagnosis)

The logic for MS–DRG 998 (Principal Diagnosis Invalid as Discharge Diagnosis) currently includes a list of diagnoses that are considered inappropriate for reporting as a principal diagnosis on an inpatient hospital claim. In other words, these conditions would reasonably be expected not to necessitate an inpatient admission. Examples of these diagnosis codes include what are referred to as the “Supervision of pregnancy” codes, as well as pregnancy, maternal care and fetal related codes with an “unspecified trimester”. We refer the reader to the ICD–10 Version 34 Definitions Manual which is available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/FY2017-IPPS-Final-Rule-Home-Page-Items/FY2017-IPPS-Final-Rule-Data-Files.html?DLPage=1&DLEntries=10&DLSort=0&DLSortDir=ascending for the complete list of diagnosis codes in MS–DRG 998 under MDC 14.

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 56840 through 56841), we was discussion regarding the supervision of “high-risk” pregnancy codes, including elderly primigravida and multigravida specifically, with regard to removing them from the Unacceptable principal diagnosis edit code list in the Medicare Code Editor (MCE). After consultation with the staff at the CDC’s NCHS, we learned that the FY 2017 ICD–10–CM Official Guidelines for Coding and Reporting were updated to explain appropriate coding for this set of codes. As a result, the codes describing supervision of high-risk pregnancy (and other supervision of pregnancy codes) remained on the Unacceptable principal diagnosis edit code list in the MCE. Therefore, the MCE code edit is consistent with the logic of MS–DRG 998 (Principal Diagnosis Invalid as Discharge Diagnosis) for these supervision of pregnancy codes.

However, as a result of our review and consultation with our clinical advisors regarding the “unspecified trimester” codes in MS–DRG 998, we have determined that there are more appropriate MS–DRG assignments for this set of codes. Although it may seem unlikely that a patient would be admitted and ultimately discharged or transferred without the caregiver or medical personnel having any further knowledge of the exact trimester, it is conceivable that a situation may present itself. For example, the pregnant patient may be from out of town or unable to communicate effectively. The fact that the specific trimester is not known or documented does not preclude the resources required to care for the patient with the particular diagnosis.

Therefore, as shown in Table 6P.3b. associated with this proposed rule (which is available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html), we are proposing to remove the 314 ICD–10–CM diagnosis codes identified with “unspecified trimester” from MS–DRG 998 and reassign them to the MS–DRGs in which their counterparts (first trimester, second trimester, or third trimester) are currently assigned as specified in Column C. This would enable more appropriate MS–DRG assignments and payment for these cases. We are inviting public comments on our proposal.

c. MS–DRG 782 (Other Antepartum Diagnoses Without Medical Complications)

The following three ICD–10–CM diagnosis codes are currently on the principal diagnosis list for the MS–DRG 782 (Other Antepartum Diagnoses without Medical Complications) logic.

<table>
<thead>
<tr>
<th>ICD–10–CM code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>O09.41 ..........</td>
<td>Supervision of pregnancy with grand multiparity, first trimester.</td>
</tr>
<tr>
<td>O09.42 ..........</td>
<td>Supervision of pregnancy with grand multiparity, second trimester.</td>
</tr>
<tr>
<td>O09.43 ..........</td>
<td>Supervision of pregnancy with grand multiparity, third trimester.</td>
</tr>
</tbody>
</table>

It was brought to our attention that these codes also are included in the MCE Unacceptable principal diagnosis code edit list. As discussed earlier in section II.F.6.b. of the preamble of this proposed rule, the supervision of pregnancy codes are accurately reflected in the MCE code edit list for Unacceptable principal diagnosis. Therefore, it is not appropriate to include the three above listed codes in MS–DRG 782.

We are proposing to remove the three codes describing supervision of pregnancy from MS–DRG 782 and reassign them to MS–DRG 998 (Principal Diagnosis Invalid as Discharge Diagnosis) to reflect a more appropriate MS–DRG assignment. We are inviting public comments on our proposal.

d. Shock During or Following Labor and Delivery

We received a request to review ICD–10–CM diagnosis code O75.1 (Shock during or following labor and delivery), which is currently assigned to MS–DRG 774 (Vaginal Delivery with Complicating Diagnosis), MS–DRG 767 (Vaginal Delivery with Sterilization and/or D&C), and MS–DRG 768 (Vaginal Delivery with O.R. Procedure Except Sterilization and/or D&C).

The requestor provided an example of a patient that delivered at Hospital A and was transferred to Hospital B for specialized care related to the diagnosis of shock. The claim for Hospital B resulted in assignment to a delivery MS–DRG, despite the fact that a delivery did not occur during that hospitalization. The requestor noted that, by not reporting the diagnosis code for shock, the claim grouped to a postpartum MS–DRG and recommended that we evaluate the issue further.

Our analysis initially involved reviewing the GROUPER logic for MS–DRGs 774, 767 and 768. As discussed earlier in section II.F.14.a. of the preamble of this proposed rule, the GROUPER logic for classification and assignment to MS–DRG 774 requires that three conditions must be met, the first of which is a vaginal delivery. Similar GROUPER logic applies for assignment to MS–DRGs 767 and 768, except that only two conditions must be met, with the first condition being a vaginal delivery. For each of these three MS–DRGs, to satisfy the first condition, one code that describes a condition or circumstance from among the three separate lists of codes must be reported. The first list is comprised of ICD–10–CM diagnosis codes that may be reported as a principal or secondary diagnosis. These diagnosis codes describe conditions in which it is assumed that a vaginal delivery has occurred. Among this first list is ICD–10–CM diagnosis code O75.1, which is included in the GROUPER logic for MS–DRGs 774, 767 and 768 (under the first condition—vaginal delivery). We refer readers to the ICD–10 MS–DRG Version 34 Definitions Manual located via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html.
PPS/FY2017-IPPS-Final-Rule-Home-Page-Items/FY2017-IPPS-Final-Rule-Data-Files.html?DLPage=1&DLEntries=10&DLSort=0&DLSortDir=ascending for documentation of the GROUPER logic associated with these MS–DRGs.

In addition, in MS–DRG 774, to satisfy the second condition, diagnosis codes that are identified as a complicating diagnosis from among two lists may be reported. The first list is comprised of ICD–10–CM diagnosis codes that may be reported as a principal or secondary diagnosis. The second list is comprised of ICD–10–CM diagnosis codes that may be reported as a secondary diagnosis. Currently, there is only one code listed under the secondary diagnosis list.

Next, our analysis involved reviewing the GROUPER logic for assignment to post-partum MS–DRG 769 (Postpartum and Post Abortion Diagnoses with Major Procedure) and MS–DRG 776 (Postpartum and Post Abortion Diagnoses without O.R. Procedure). The GROUPER logic for these postpartum MS–DRGs requires that a principal diagnosis be reported from a list of several conditions, such as those following pregnancy, those complicating the puerperium, conditions that occurred during or following delivery and conditions associated with lactation disorders. For assignment to MS–DRG 769, the GROUPER logic also requires that a major procedure be reported in addition to a principal diagnosis from the list of conditions.

As a result of our analysis, we agree with the requestor that ICD–10–CM diagnosis code O75.1 should be added to the GROUPER logic for assignment to the postpartum MS–DRGs. This diagnosis code is consistent with other diagnosis codes structured within the GROUPER logic for assignment to MS–DRGs 769 and 776, and clearly represents a post-partum diagnosis with the terminology “during or following labor and delivery” in the title. We believe that adding this diagnosis code to the postpartum MS–DRGs will enable more appropriate MS–DRG assignment for cases where a delivery did not occur. Therefore, we are proposing the following:

- Removing ICD–10–CM diagnosis code O75.1 from the list of principal or secondary diagnosis under the first condition—vaginal delivery GROUPER logic in MS–DRGs 774, 767, and 768;
- Moving ICD–10–CM diagnosis code O75.1 from the list of principal or secondary diagnosis under the second condition—complicating diagnosis for MS–DRG 774 to the secondary diagnosis list only; and
- Adding ICD–10–CM diagnosis code O75.1 to the principal diagnosis list GROUPER logic in MS–DRGs 769 and 776.

We are inviting public comments on our proposals.

7. MDC 15 (Newborns and Other Neonates With Conditions Originating in Perinatal Period); Observation and Evaluation of Newborn

We received a request to add the ICD–10–CM diagnosis codes describing observation and evaluation of newborns for suspected conditions that are ruled out to MS–DRG 795 (Normal Newborn). The 14 diagnosis codes describing observation and evaluation of newborns for suspected conditions ruled out are displayed in the table below.

<table>
<thead>
<tr>
<th>ICD–10–CM code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Z05.0 ..........</td>
<td>Observation and evaluation of newborn for suspected cardiac condition ruled out.</td>
</tr>
<tr>
<td>Z05.1 ..........</td>
<td>Observation and evaluation of newborn for suspected infectious condition ruled out.</td>
</tr>
<tr>
<td>Z05.2 ..........</td>
<td>Observation and evaluation of newborn for suspected respiratory condition ruled out.</td>
</tr>
<tr>
<td>Z05.3 ..........</td>
<td>Observation and evaluation of newborn for suspected metabolic condition ruled out.</td>
</tr>
<tr>
<td>Z05.41 .........</td>
<td>Observation and evaluation of newborn for suspected congenital condition ruled out.</td>
</tr>
<tr>
<td>Z05.42 .........</td>
<td>Observation and evaluation of newborn for suspected metabolic condition ruled out.</td>
</tr>
<tr>
<td>Z05.43 .........</td>
<td>Observation and evaluation of newborn for suspected immunologic condition ruled out.</td>
</tr>
<tr>
<td>Z05.5 ..........</td>
<td>Observation and evaluation of newborn for suspected gastrointestinal condition ruled out.</td>
</tr>
<tr>
<td>Z05.61 .........</td>
<td>Observation and evaluation of newborn for suspected genitourinary condition ruled out.</td>
</tr>
<tr>
<td>Z05.71 .........</td>
<td>Observation and evaluation of newborn for suspected skin and subcutaneous tissue condition ruled out.</td>
</tr>
<tr>
<td>Z05.72 .........</td>
<td>Observation and evaluation of newborn for suspected musculoskeletal condition ruled out.</td>
</tr>
<tr>
<td>Z05.73 .........</td>
<td>Observation and evaluation of newborn for suspected connective tissue condition ruled out.</td>
</tr>
<tr>
<td>Z05.8 ..........</td>
<td>Observation and evaluation of newborn for other specified condition ruled out.</td>
</tr>
<tr>
<td>Z05.9 ..........</td>
<td>Observation and evaluation of newborn for unspecified condition ruled out.</td>
</tr>
</tbody>
</table>

The requestor expressed concern that currently when one of these ruled out codes is added to a newborn encounter with a principal diagnosis described by ICD–10–CM code Z38.00 (Single liveborn infant, delivered vaginally), the case is assigned to MS–DRG 794 (Neonate with Other Significant Problems). The requestor stated that this assignment appears to be in error and that the assignment should instead be to MS–DRG 795 (Normal Newborn).

We reviewed Section 1.C.16.b. of the 2017 ICD–10–CM Official Guidelines for Coding and Reporting which includes the following instructions for the diagnosis codes listed in the table above:

- Assign a code from category Z05 (Observation and evaluation of newborns and infants for suspected conditions ruled out) to identify those instances when a healthy newborn is evaluated for a suspected condition that is determined after study not to be present. Do not use a code from category Z05 when the patient has identified signs or symptoms of a suspected problem; in such cases code the sign or symptom.
- A code from category Z05 may also be assigned as a principal or first-listed code for readmissions or encounters when the code from category Z38 code no longer applies. Codes from category Z05 are for use only for healthy newborns and infants for which no condition after study is found to be present.
- A code from category Z05 is to be used as a secondary code after the code from category Z38. Liveborn infants according to place of birth and type of delivery.

After review of the guidelines and discussion with our clinical advisors, we agree with the requestor that the assignment of these codes to MS–DRG 794 is not accurate because the assignment incorrectly labels the newborns as having a significant problem when the condition does not truly exist. We and our clinical advisors also agree that the above list of diagnosis codes should be added to MS–DRG 795. Therefore, we are proposing to add the 14 diagnosis codes describing observation and evaluation of newborns for suspected conditions that are ruled out listed in the table above to the GROUPER logic for MS–DRGs 794, 767, and 768. We are inviting public comments on our proposals.
8. MDC 21 (Injuries, Poisonings and Toxic Effects of Drugs): Complication Codes

We received a request to examine the ICD–10–CM diagnosis codes in the T85.8-series of codes that describe other specified complications of internal prosthetic devices, implants and grafts, not elsewhere classified and their respective MS–DRG assignments. According to the requestor, the 7th character values in this series of codes impact the MS–DRG assignment under MDC 21 (Injuries, Poisonings and Toxic Effects of Drugs) and MDC 23 (Factors Influencing Health Status & Other Contacts with Health Services) that have resulted in inconsistencies (that is, shifts) between the MS–DRG assignments under Version 33 and Version 34 of the ICD–10 MS–DRGs.

Under ICD–10–CM, diagnosis codes in the range of S00 through T88 require a 7th character value of “A-” initial encounter, “D-” subsequent encounter, or “S-” sequela to identify if the patient is undergoing active treatment for a condition. For complication codes, active treatment refers to treatment for the condition described by the code, even though it may be related to an earlier precipitating problem.

The requestor suggested that the following list of diagnosis codes with the 7th character “A” (initial encounter) may have been inadvertently assigned to the GROUPER logic list of principal diagnoses under MDC 23 because these codes may have been inadvertently assigned to the GROUPER logic list of principal diagnoses for MS–DRGs 949 and 950 (Aftercare with CC/MCC and without CC/MCC, respectively) under MDC 23 because it found claims that grouped to these MS–DRGs (949 and 950) when one of the following diagnosis codes was reported with an O.R. procedure, the requestor found claims grouping to MS–DRG 939, 940, or 941 (O.R. Procedures with Diagnoses of Other Contact with Health Services with MCC, with CC and without CC/MCC, respectively) that had previously grouped to MDC 21 under Version 33 of the ICD–10 MS–DRGs.

### ICD–10–CM diagnosis code | Code description
---|---
T85.818A | Embolism due to other internal prosthetic devices, implants and grafts, initial encounter.
T85.828A | Fibrosis due to other internal prosthetic devices, implants and grafts, initial encounter.
T85.838A | Hemorrhage due to other internal prosthetic devices, implants and grafts, initial encounter.
T85.848A | Pain due to other internal prosthetic devices, implants and grafts, initial encounter.
T85.858A | Stenosis due to other internal prosthetic devices, implants and grafts, initial encounter.
T85.868A | Thrombosis due to other internal prosthetic devices, implants and grafts, initial encounter.
T85.898A | Other specified complication of other internal prosthetic devices, implants and grafts, initial encounter.

The requestor also suggested that the following list of diagnosis codes with the 7th character “D” (subsequent encounter) may have been inadvertently assigned to the GROUPER logic list of principal diagnoses for MS–DRGs 919, 920, or 921 in MDC 21. The requestor noted that these codes were new, effective October 1, 2016 (FY 2017), and the predecessor codes grouped to MS–DRGs 919, 920, and 921 in MDC 21 under Version 33 of the ICD–10 MS–DRGs in FY 2016.

### ICD–10–CM diagnosis code | Code description
---|---
T85.810D | Embolism due to nervous system prosthetic devices, implants and grafts, subsequent encounter.
T85.820D | Fibrosis due to nervous system prosthetic devices, implants and grafts, subsequent encounter.
T85.830D | Hemorrhage due to nervous system prosthetic devices, implants and grafts, subsequent encounter.
T85.840D | Pain due to nervous system prosthetic devices, implants and grafts, subsequent encounter.
T85.850D | Stenosis due to nervous system prosthetic devices, implants and grafts, subsequent encounter.
T85.860D | Thrombosis due to nervous system prosthetic devices, implants and grafts, subsequent encounter.
T85.890D | Other specified complication of nervous system prosthetic devices, implants and grafts, subsequent encounter.

The requestor also suggested that the following list of additional diagnosis codes with the 7th character “D” (subsequent encounter) may have been inadvertently assigned to the GROUPER logic list of principal diagnoses for MS–DRGs 922 and 923 (Other Injury, Poisoning and Toxic Effect with MCC and without MCC, respectively) also under MDC 21. The requestor noted that these codes were also new, effective October 1, 2016 (FY 2017) and that the predecessor codes grouped to MS–DRGs 949 and 950 in MDC 23 under Version 33 of the ICD–10 MS–DRGs in FY 2016.

### ICD–10–CM diagnosis code | Code description
---|---
T85.818D | Embolism due to other internal prosthetic devices, implants and grafts, subsequent encounter.
T85.828D | Fibrosis due to other internal prosthetic devices, implants and grafts, subsequent encounter.
T85.838D | Hemorrhage due to other internal prosthetic devices, implants and grafts, subsequent encounter.
The requestor believed that the lists of diagnosis codes above with 7th character “D” (subsequent encounter) would be more appropriately assigned to MS–DRGs 949 and 950 under MDC 23, according to its review of the 2017 Official Coding Guidelines for use of the 7th character and assignment of other diagnoses of associated complications of care.

We ran test cases to determine if we could duplicate the requestor’s findings with regard to the shifts in MS–DRG assignment between Version 33 and Version 34 of the ICD–10 MS–DRGs.

Results of our review were consistent with the requestor’s findings. We found that the T85.8-series of diagnosis codes with the 7th character of “A” (initial encounter) and 7th character of “D” (subsequent encounter) were inadvertently assigned to the incorrect MDC for Version 34 of the ICD–10 MS–DRGs, which led to inconsistencies (MS–DRG shifts) when compared to Version 33 of the ICD–10 MS–DRGs. Our analysis also included review of all of the diagnosis codes in the T85.8-series and their current MDC and MS–DRG assignments, as well as review of the 2017 Official Coding Guidelines for use of the 7th character and assignment of other diagnoses of associated complications of care. Based on the results of our review, we agree with the requestor’s findings.

In addition, we identified the following list of diagnosis codes with the 7th character “S” (sequela) that appear to have been inadvertently assigned to MS–DRGs 949 and 950 in MDC 23 rather than MDC 21 in MS–DRGs 922 and 923 (Other Injury, Poisoning and Toxic Effect with MCC and without MCC, respectively).

<table>
<thead>
<tr>
<th></th>
<th></th>
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<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>T85.810S</td>
<td>Embolism due to nervous system prosthetic devices, implants and grafts, sequela.</td>
<td>21</td>
<td>919, 920, 921</td>
<td>23</td>
<td>949, 950</td>
</tr>
<tr>
<td>T85.810S</td>
<td>Embolism due to nervous system prosthetic devices, implants and grafts, sequela.</td>
<td>23</td>
<td>949, 950</td>
<td>21</td>
<td>922, 923</td>
</tr>
<tr>
<td>T85.810S</td>
<td>Embolism due to other internal prosthetic devices, implants and grafts, initial encounter.</td>
<td>21</td>
<td>922, 923</td>
<td>23</td>
<td>949, 950</td>
</tr>
<tr>
<td>T85.810S</td>
<td>Embolism due to other internal prosthetic devices, implants and grafts, subsequent encounter.</td>
<td>21</td>
<td>919, 920, 921</td>
<td>23</td>
<td>949, 950</td>
</tr>
<tr>
<td>T85.820S</td>
<td>Fibrosis due to nervous system prosthetic devices, implants and grafts, subsequent encounter.</td>
<td>21</td>
<td>919, 920, 921</td>
<td>23</td>
<td>949, 950</td>
</tr>
<tr>
<td>T85.820S</td>
<td>Fibrosis due to nervous system prosthetic devices, implants and grafts, sequela.</td>
<td>23</td>
<td>949, 950</td>
<td>21</td>
<td>922, 923</td>
</tr>
<tr>
<td>T85.820S</td>
<td>Fibrosis due to other internal prosthetic devices, implants and grafts, initial encounter.</td>
<td>23</td>
<td>949, 950</td>
<td>21</td>
<td>919, 920, 921</td>
</tr>
<tr>
<td>T85.820S</td>
<td>Fibrosis due to other internal prosthetic devices, implants and grafts, subsequent encounter.</td>
<td>21</td>
<td>919, 920, 921</td>
<td>23</td>
<td>949, 950</td>
</tr>
<tr>
<td>T85.830D</td>
<td>Hemorrhage due to nervous system prosthetic devices, implants and grafts, subsequent encounter.</td>
<td>23</td>
<td>949, 950</td>
<td>21</td>
<td>922, 923</td>
</tr>
<tr>
<td>T85.830D</td>
<td>Hemorrhage due to nervous system prosthetic devices, implants and grafts, initial encounter.</td>
<td>23</td>
<td>919, 920, 921</td>
<td>23</td>
<td>949, 950</td>
</tr>
</tbody>
</table>
In FY 2016, we received requests to modify the MS–DRG assignment for MS–DRGs 945 and 946 (Rehabilitation With CC/MCC and Without CC/MCC, Respectively). This issue was addressed in the FY 2017 IPPS/LTCH PPS proposed and final rules (81 FR 24998 through 25000 and 81 FR 56826 through 56831). For FY 2017, we did not change the MS–DRG assignments for MS–DRGs 945 and 946. We did not receive a request to address this issue as part of this FY 2018 IPPS/LTCH PPS proposed rule or suggestions on how to update the MS–DRGs 945 and 946 logic. However, we did refer the FY 2016 requests for a new ICD–10–CM diagnosis code to the Centers for Disease Control and Prevention (CDC) for consideration at a future meeting of the ICD–10 Coordination and Maintenance Committee. CDC has the lead on updating and maintaining ICD–10–CM codes. CDC did not address the issue at the September 13–14, 2016 ICD–10 Coordination and Maintenance Committee meeting. When the topic was not addressed at the September 13–14, 2016 ICD–10 Coordination and Maintenance Committee meeting, we asked CDC to address the code request at the March 7–8, 2017 meeting of the ICD–10 Coordination and Maintenance Committee. The topic was on the agenda for the March 7–8, 2017 ICD–10 Coordination and Maintenance Committee meeting. The deadline for providing comments on proposals considered at this meeting was April 7, 2017. Any new codes approved after this meeting which will be implemented on October 1, 2017 will be posted on the CMS Web site at: http://www.cms.gov/Medicare/Coding/ICD10/index.html and on the CDC Web site at: http://www.cdc.gov/nchs/icd/icd10.html in June 2017. New codes also will be included in the FY 2018 IPPS/LTCH PPS final rule. As addressed in the FY 2017 IPPS/LTCH PPS final rule, the ICD–9–CM MS–DRGs used ICD–9–CM codes reported as the principal diagnosis that clearly identified an encounter for rehabilitation services, such as diagnosis codes V57.89 (Care involving other specified rehabilitation procedure) and V57.9 (Care involving unspecified rehabilitation procedure), and these codes were not included in ICD–10–CM. Given this lack of ICD–10–CM codes to indicate that the reason for the encounter was for rehabilitation, the ICD–10 MS–DRG logic could not reflect the logic of the ICD–9–CM MS–DRGs. Commenters on the final rule recommended that CDC create new diagnosis codes for these concepts in ICD–10–CM so that the MS–DRG logic could be updated to more closely reflect that of the ICD–9–CM MS–DRGs. If new ICD–10–CM codes are created for encounter for rehabilitation services, we would address any updates to MS–DRGs 945 and 946 utilizing these new codes in future rulemaking. In the meantime, we welcome other specific recommendations on how to update MS–DRGs 945 and 946 utilizing these new codes in future rulemaking. In the meantime, we welcome other specific recommendations on how to update MS–DRGs 945 and 946 utilizing these new codes in future rulemaking.
As shown by the tables above, there was a decrease of 3,320 MS–DRG 945 cases (from 3,991 to 671) from FY 2015, when claims were submitted with ICD–9–CM codes, to FY 2016 when ICD–10 codes were submitted. There was a decrease of 1,027 MS–DRG 946 cases (from 1,184 to 157) from FY 2015 to FY 2016. The average length of stay increased 0.5 days (from 10.3 to 10.8 days) for MS–DRG 945 and decreased 0.7 days (from 8.0 to 7.3 days) for MS–DRG 946. The average costs decreased by $428 (from $8,242 to $7,814) for MS–DRG 945 cases and increased by $350 (from $7,322 to $7,672) for MS–DRG 946 cases. The number of cases was significantly lower in FY 2016 compared to FY 2015. However, the difference in average length of stay and average costs did not show large changes.

We also examined possible MS–DRGs where these cases may have been assigned in FY 2016 based on increases in the number of cases. Because there is not a diagnosis code that could be reported as a principal diagnosis, which would indicate if the admissions were for rehabilitation services, we are unable to determine if these were cases admitted for rehabilitation that moved from MS–DRGs 945 and 946 because of the lack of a code for encounter for rehabilitation, or if there was simply a change in the number of cases. The following tables show our findings for MS–DRG 056 (Degenerative Nervous System Disorders with MCC); MS–DRG 057 (Degenerative Nervous System Disorders without MCC); MS–DRG 079 (Hypertensive Encephalopathy without CC/MCC); MS DRG 083 (Traumatic Stupor & Coma, Coma >1 Hour with CC); MS–DRG 084 (Traumatic Stupor & Coma, Coma >1 Hour without CC/MCC); MS–DRG 092 (Other Disorders of Nervous System with MCC); and MS–DRG 093 (Other Disorders of Nervous System without CC/MCC).

<table>
<thead>
<tr>
<th>MS–DRG 056</th>
<th>MS–DRG 057</th>
<th>MS–DRG 079</th>
<th>MS–DRG 083</th>
<th>MS–DRG 084</th>
<th>MS–DRG 092</th>
<th>MS–DRG 093</th>
</tr>
</thead>
<tbody>
<tr>
<td>9,548</td>
<td>25,652</td>
<td>618</td>
<td>2,516</td>
<td>1,955</td>
<td>12,643</td>
<td>7,928</td>
</tr>
<tr>
<td>7.3</td>
<td>5.1</td>
<td>2.7</td>
<td>4.3</td>
<td>2.8</td>
<td>5.7</td>
<td>2.8</td>
</tr>
<tr>
<td>$12,606</td>
<td>7,918</td>
<td>5,212</td>
<td>9,446</td>
<td>6,824</td>
<td>11,158</td>
<td>5,182</td>
</tr>
</tbody>
</table>

As shown by the tables above, the MS–DRGs that show the largest increase in number of cases do not show significant changes in the average length of stay or average costs. For instance, MS–DRG 079 cases doubled from FY 2015 to FY 2016 (from 618 to 1,233). However, the average length of stay did not change from 2.7 days and the average costs increased only $367 (from $5,212 to $5,579). MS–DRG 083 cases increased by 1,542 (from 2,516 to 4,058) with a 1.9 day increase in the average length of stay (from 4.3 to 6.2 days); however, the average costs decreased only $312 (from $9,446 to $9,134).

There were large changes for MS–DRG 092 with cases increasing by 6,749 (from 12,643 to 19,392), the average length of stay decreasing by 1.8 days (from 5.7 to 3.9) and the average costs decreasing by $4,452 (from $11,158 to $6,706). Once again, it is not possible to determine if any changes are a result of the impact of not having a code for the encounter for rehabilitation services to report as a principal diagnosis, or if other factors such as changes in types of patient admissions were involved. Given the lack of a diagnosis code to capture the principal diagnosis of encounter for rehabilitation, we are unable to update MS–DRG 945 or MS–DRG 946 to better identify those cases in which patients are admitted for rehabilitation services. If the CDC creates a new code, we will consider proposing updates to MS–DRGs 945 and 946 in the future.

We are inviting public comments on our proposal not to update MS–DRGs 945 and 946 for FY 2018.

10. Proposed Changes to the Medicare Code Editor (MCE)

The Medicare Code Editor (MCE) is a software program that detects and reports errors in the coding of Medicare.
The age ranges defined within the Age Conflict edits were established with the implementation of the IPPS. The adult age range includes the minimum age of 15 years for those patients who are declared emancipated minors. We note that, historically, we have not provided coding advice in rulemaking with respect to policy. We collaborate with the American Hospital Association (AHA) through the Coding Clinic for ICD–10–CM and ICD–10–PCS to promote proper coding. We recommend that the requestor and other interested parties submit any questions pertaining to correct coding practices for this specific issue to the AHA.

(1) Perinatal/Newborn Diagnosis Category

Under the ICD–10 MCE, the Perinatal/Newborn Diagnosis category under the Age Conflict edit considers the age of 0 years only; a subset of diagnoses which will only occur during the perinatal or newborn period of age 0 to be inclusive. This includes conditions that have their origin in the fetal or perinatal period (before birth through the first 28 days after birth) even if morbidity occurs later. For that reason, the diagnosis codes on this Age Conflict edit list would be expected to apply to conditions or disorders specific to that age group only.

In the ICD–10–CM classification, there are two diagnosis codes that describe conditions as occurring during infancy and the neonatal period that are currently not on the Perinatal/Newborn Diagnosis category edit code list. We consulted with staff at the Centers for Disease Control’s (CDC’s) National Center for Health Statistics (NCHS) because NCHS has the lead responsibility for the ICD–10–CM diagnosis codes. The NCHS’ staff confirmed that, although diagnosis codes D80.7 (Transient hypogammaglobulinemia of infancy) and diagnosis code E71.511 (Neonatal adrenoleukodystrophy) do occur during infancy and the neonatal period, both conditions can last beyond the 28-day timeframe which is used to define the perinatal/newborn period. These diagnosis codes are not intended to be restricted for assignment to newborn patients. Therefore, we are proposing to not add these two diagnosis codes to the Perinatal/Newborn Diagnosis category under the Age Conflict edit. We are inviting public comments on our proposal.

(2) Pediatric Diagnosis Category

Under the ICD–10 MCE, the Pediatric diagnosis category under the Age Conflict edit considers the age range of 0 to 17 years inclusive. For that reason, the diagnosis codes on this Age Conflict edit list would be expected to apply to conditions or disorders specific to that age group only.

The ICD–10–CM diagnosis code list for the Pediatric diagnosis category under the Age Conflict edit currently includes a diagnosis code pertaining to dandruff that is not intended to apply to pediatric patients only. We consulted with staff at the Centers for Disease Control’s (CDC’s) National Center for Health Statistics (NCHS) because NCHS has the lead responsibility for the ICD–10–CM diagnosis codes. The NCHS’ staff confirmed that, although diagnosis code L21.0 (Seborrhea capitis) has an inclusion term of “Cradle cap,” the description of the diagnosis code is not intended to be restricted for assignment of pediatric patients. Therefore, we are proposing to remove diagnosis code L21.0 from the list of diagnosis codes for the Pediatric diagnosis category under the Age Conflict edit. We are inviting public comments on our proposal.
(3) Maternity Diagnoses

Under the ICD–10 MCE, the Maternity diagnosis category under the Age Conflict edit considers the age range of 12 to 55 years inclusive. For that reason, the ICD–10–CM diagnosis codes on this Age Conflict edit list would be expected to apply to conditions or disorders specific to that age group only.

As discussed in section II.F.12. of the preamble of this proposed rule, Table 6A.—New Diagnosis Codes lists the new ICD–10–CM diagnosis codes that have been approved to date, which will become effective with discharges occurring on and after October 1, 2017. Included on this list are a number of diagnosis codes associated with pregnancy and maternal care that we believe are appropriate to add to the list of diagnosis codes for the Maternity diagnoses category under the Age Conflict edit. We refer readers to Table 6P.1a. associated with this proposed rule (which is available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) for a review of the ICD–10–CM diagnosis codes that we are proposing to add to the Age Conflict edit list. We are inviting public comments on our proposal.

b. Sex Conflict Edit

In the MCE, the Sex Conflict edit detects inconsistencies between a patient’s sex and any diagnosis or procedure on the patient’s record; for example, a male patient with cervical cancer (diagnosis) or a female patient with a prostatectomy (procedure). In both instances, the indicated diagnosis or the procedure conflicts with the stated sex of the patient. Therefore, the patient’s diagnosis, procedure, or sex is presumed to be incorrect.

(1) Diagnoses for Males Only Edit

We received a request to review the following ICD–10–CM diagnosis codes pertaining to conditions associated with males for possible inclusion on the list of diagnosis codes for the Diagnoses for Males Only edit.

<table>
<thead>
<tr>
<th>ICD–10–CM code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>B37.42 ..........</td>
<td>Candidal balanitis.</td>
</tr>
<tr>
<td>N35.011 ..........</td>
<td>Post-traumatic bulbous urethral stricture.</td>
</tr>
<tr>
<td>N35.012 ..........</td>
<td>Post-traumatic membranous urethral stricture.</td>
</tr>
<tr>
<td>N35.013 ..........</td>
<td>Post-traumatic anterior urethral stricture.</td>
</tr>
<tr>
<td>N35.112 ..........</td>
<td>Postinfective bulbous urethral stricture, not elsewhere classified.</td>
</tr>
<tr>
<td>N35.113 ..........</td>
<td>Postinfective membranous urethral stricture, not elsewhere classified.</td>
</tr>
<tr>
<td>N35.114 ..........</td>
<td>Postinfective anterior urethral stricture, not elsewhere classified.</td>
</tr>
</tbody>
</table>

We agree with the requestor that diagnosis code B37.42 describes a condition that is applicable only to males. Balanitis is the inflammation of the glans (rounded head) of the penis. We also agree that the diagnosis codes listed above that align under subcategory N35.01 (Post-traumatic urethral stricture, male) and subcategory N35.11 (Postinfection urethral stricture, not elsewhere classified, male) are appropriate to add to the list of diagnosis codes for the Diagnoses for Males Only edit because these diagnosis codes include specific terminology that is applicable only to males. Further, we agree that diagnosis code N99.115 is appropriate to add to the list of diagnosis codes for the Diagnoses for Males Only edit because subcategory N99.11 (Postprocedural urethral stricture, male) includes specific terminology that is applicable to males only as well. Therefore, we are proposing to add the ICD–10–CM diagnosis codes listed in the table above to the list of diagnosis codes for the Diagnoses for Males Only edit.

We also are proposing to remove ICD–10–CM diagnosis code Q64.0 (Epispidias) from the list of diagnosis codes for the Diagnoses for Males Only edit because this rare, congenital condition involving the opening of the urethra can occur in both males and females. In addition, as discussed in section II.F.12. of the preamble of this proposed rule, Table 6A.—New Diagnosis Codes lists the new ICD–10–CM diagnosis codes that have been approved to date, which will become effective with discharges occurring on and after October 1, 2017. Included on this list are a number of diagnosis codes associated with male body parts that we believe are appropriate to add to the list of diagnosis codes for the Diagnoses for Males Only category under the Sex Conflict edit. We refer readers to Table 6P.1b. associated with this proposed rule (which is available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) for a review of the ICD–10–CM diagnosis codes that we are proposing to add to the list of diagnosis codes for the Diagnoses for Males Only category.

We are inviting public comments on our proposals.

(2) Diagnoses for Females Only

We received a request to review the following ICD–10–CM diagnosis codes for possible removal from the list of diagnosis codes for the Diagnoses for Females Only edit.

<table>
<thead>
<tr>
<th>ICD–10–CM code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>F52.6 ..........</td>
<td>Dyspareunia not due to a substance or known physiological condition.</td>
</tr>
<tr>
<td>J84.81 ..........</td>
<td>Lymphangioleiomyomatosis.</td>
</tr>
<tr>
<td>R97.1 ..........</td>
<td>Elevated cancer antigen 125 [CA 125].</td>
</tr>
</tbody>
</table>

The requestor noted that, in the ICD–10–CM classification, the term “Dyspareunia” (painful sexual intercourse) has specified codes for males and females located in the Alphabetic Index to Diseases for Reporting Physiological Dyspareunia. However, the indexing for diagnosis code F52.6 (Dyspareunia not due to a substance or known physiological condition) specifies that it is not due to a physiological condition and the entry is not gender specific. According to the requestor, while the condition is most often associated with female sexual dysfunction, there is a subset of males who also suffer from this condition.
In addition, the requestor stated that diagnosis code J84.81 (Lymphangioleiomyomatosis) describes a rare form of lung disease believed to occur more often in patients with tuberous sclerosis complex (TSC), a disorder due to genetic mutation. Although the condition is described as being exclusive to women, unique cases for men with TSC have also been reported.

Lastly, the requestor indicated that diagnosis code R97.1 (Elevated cancer antigen 125 [CA 125]) describes the tumor marker that commonly identifies ovarian cancer cells in women. However, the requestor stated that high levels have also been demonstrated in men (and women) with lung cancer as well.

We reviewed ICD–10–CM diagnosis codes F52.6, J84.81, and R97.1, and we agree with the requestor that Dyspareunia, not due to a physiological condition, can also occur in males. We also agree that the condition of Lymphangioleiomyomatosis and Elevated CA 125 levels can be found in males. Therefore, we are proposing to remove these three diagnosis codes from the list of diagnosis codes for the Diagnoses for Females Only edit. We are inviting public comments on our proposals.

In addition, we are proposing to add new diagnosis code Z40.03 (Encounter for prophylactic removal of fallopian tube(s)) to the list of diagnosis codes for the Diagnoses for Females Only edit. Currently, diagnosis code Z40.02 (Encounter for prophylactic removal of ovary) is on the edit’s code list; therefore, inclusion of new diagnosis code Z40.03 would be consistent. We refer readers to Table 6A.—New Diagnosis Codes associated with this proposed rule (which is available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html for the list of new ICD–10–CM diagnosis codes finalized to date. We are inviting public comments on our proposal.

c. Non-Covered Procedure Edit: Gender Reassignment Surgery

In the MCE, the Non-Covered Procedure edit identifies procedures for which Medicare does not provide payment. Payment is not provided due to specific criteria that are established in the National Coverage Determination (NCD) process. We refer readers to the Web site at: https://www.cms.gov/Medicare/Coverage/Determination-Process/howtorequestanNCD.html for additional information on this process.

We examined ICD–10–CM diagnosis codes in Chapter 1 (Certain Infectious and Parasitic Diseases) of the Classification Manual that fall within the range of three code categories for “Bacterial and Viral Infectious Agents” (B95 through B97) that, as a result of the instructional note, are not appropriate to report as a principal diagnosis. We are proposing to add the 45 ICD–10–CM diagnosis codes shown in Table 6P.1c. associated with this proposed rule (which is available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) to the list of codes for the Unacceptable Principal Diagnosis edit. We are inviting public comments on our proposal.

(2) Mental Disorders Due to Known Physiological Conditions (F01 Through F09)

We examined ICD–10–CM diagnosis codes in Chapter 5 (Mental and Behavioral Disorders) of the Classification Manual that fall within the range of nine code categories for “Mental Disorders Due to Known

<table>
<thead>
<tr>
<th>ICD–10–CM code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0W4M070</td>
<td>Creation of vagina in male perineum with autologous tissue substitute, open approach.</td>
</tr>
<tr>
<td>0W4M0J0</td>
<td>Creation of vagina in male perineum with synthetic substitute, open approach.</td>
</tr>
<tr>
<td>0W4M0K0</td>
<td>Creation of vagina in male perineum with nonautologous tissue substitute, open approach.</td>
</tr>
<tr>
<td>0W4M0Z0</td>
<td>Creation of vagina in male perineum, open approach.</td>
</tr>
<tr>
<td>0W4N071</td>
<td>Creation of penis in female perineum with autologous tissue substitute, open approach.</td>
</tr>
<tr>
<td>0W4N0J1</td>
<td>Creation of penis in female perineum with synthetic substitute, open approach.</td>
</tr>
<tr>
<td>0W4N0K1</td>
<td>Creation of penis in female perineum with autologous tissue substitute, open approach.</td>
</tr>
<tr>
<td>0W4N0Z1</td>
<td>Creation of penis in female perineum, open approach.</td>
</tr>
</tbody>
</table>

Therefore, we are proposing to remove the ICD–10–PCS procedure codes included in the table above from the list of procedure codes for the Non-Covered Procedure edit to help resolve claims processing issues associated with the reporting of these procedure codes. We are inviting public comments on our proposal.

d. Unacceptable Principal Diagnosis Edit

In the MCE, there are select codes that describe a circumstance that influences an individual’s health status, but does not actually describe a current illness or injury. There are also codes that are not specific manifestations but may be due to an underlying cause. These codes are considered unacceptable as a principal diagnosis. In limited situations, there are a few codes on the MCE Unacceptable Principal Diagnosis edit code list that are considered “acceptable” when a specified secondary diagnosis is also coded and reported on the claim.

(1) Bacterial and Viral Infectious Agents (B95 Through B97)

We examined ICD–10–CM diagnosis codes in Chapter 1 (Certain Infectious and Parasitic Diseases) of the Classification Manual that fall within the range of three code categories for “Bacterial and Viral Infectious Agents” (B95 through B97). The instructional note provided at this section states that these categories are provided for use as supplementary or additional codes to identify the infectious agent(s) in diseases classified elsewhere.

We identified 45 ICD–10–CM diagnosis codes within the range of these code categories for “Bacterial and Viral Infectious Agents” (B95 through B97) that, as a result of the instructional note, are not appropriate to report as a principal diagnosis. We are proposing to add the 45 ICD–10–CM diagnosis codes shown in Table 6P.1c. associated with this proposed rule (which is available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) to the list of codes for the Unacceptable Principal Diagnosis edit. We are inviting public comments on our proposal.

(2) Mental Disorders Due to Known Physiological Conditions (F01 Through F09)

We examined ICD–10–CM diagnosis codes in Chapter 5 (Mental and Behavioral Disorders) of the Classification Manual that fall within the range of nine code categories for “Mental Disorders Due to Known
Physiological Conditions” (F01 through F09). The instructional note provided at this section states that this block comprises a range of mental disorders grouped together on the basis of their having in common a demonstrable etiology in cerebral disease, brain injury, or other insult leading to cerebral dysfunction. The dysfunction may be primary, as in diseases, injuries, and insults that affect the brain directly and selectively; or secondary, as in systemic diseases and disorders that attack the brain only as one of the multiple organs or systems of the body that are involved.

We identified 21 ICD–10–CM diagnosis codes that fall within the range of these code categories for “Mental Disorders Due to Known Physiological Conditions” (F01 through F09). Of these nine code categories, seven have a “Code first the underlying physiological condition” note. For example, at code category F01-Vascular Dementia, the note reads, “Code first the underlying physiological condition or sequelae of cerebrovascular disease.” There are a total of 19 diagnosis codes that fall under these 7 code categories, with a “Code first” note and, therefore, are not appropriate to report as a principal diagnosis. Therefore, we are proposing to add the 19 ICD–10–CM diagnosis codes shown in Table 6P.1d. associated with this proposed rule (which is available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) to the list of codes for the Unacceptable Principal Diagnosis edit. We are inviting public comments on our proposal.

(3) Other Obstetric Conditions, Not Elsewhere Classified (O94 Through O9A)

We examined ICD–10–CM diagnosis codes in Chapter 15 (Pregnancy, Childbirth and the Puerperium) of the Classification Manual that fall within the range of four code categories for “Other Obstetric Conditions, Not Elsewhere Classified” (O94 through O9A). The instructional note provided at this section under category O94 states that “this category is to be used to indicate conditions in O00 through O77, O85 through O94 and O98 through O9A as the cause of late effects. The sequelae include conditions specified as such, or as late effects, which may occur at any time after the puerperium. Code first condition resulting from (sequela) of complication of pregnancy, childbirth, and the puerperium.”

We identified one ICD–10–CM diagnosis code within the range of these code categories for “Other Obstetric Conditions, Not Elsewhere Classified” (O94 through O9A) that, as a result of the instructional note, is not appropriate to report as a principal diagnosis because that code identifies the cause of the late effect. This ICD–10–CM diagnosis code is O94 (Sequelae of complication of pregnancy, childbirth, and the puerperium). We are proposing to add ICD–10–CM diagnosis code O94 to the list of codes for the Unacceptable Principal Diagnosis edit. We are inviting public comments on our proposal.

(5) General Symptoms and Signs (R50 Through R69)

We examined ICD–10–CM diagnosis codes in Chapter 18 (Symptoms, Signs and Abnormal Findings) of the Classification Manual that fall within the range of code categories for “General Symptoms and Signs” (R50 through R69), specifically, at code category R65—Symptoms and signs associated with systemic inflammation and infection. There is an instructional note at subcategory R65.1—Systemic inflammatory response syndrome (SIRS) of non-infectious origin, which states “Code first underlying condition, such as: Heatstroke (T67.0); Injury and trauma (S00–T88).” There is also an instructional note at subcategory R65.2—Severe sepsis, which states “Code first underlying infection, such as:” and provides a list of examples.

We identified four ICD–10–CM diagnosis codes in these subcategories that, as a result of the instructional notes described above, are not appropriate to report as a principal diagnosis. These four ICD–10–CM codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–CM code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>R65.10 ..........</td>
<td>Systemic inflammatory response syndrome (SIRS) of non-infectious origin without acute organ dysfunction.</td>
</tr>
<tr>
<td>R65.11 ..........</td>
<td>Systemic inflammatory response syndrome (SIRS) of non-infectious origin with acute organ dysfunction.</td>
</tr>
<tr>
<td>R65.20 ..........</td>
<td>Severe sepsis without septic shock.</td>
</tr>
<tr>
<td>R65.21 ..........</td>
<td>Severe sepsis with septic shock.</td>
</tr>
</tbody>
</table>

We are proposing to add the four ICD–10–CM diagnosis codes shown in the table above to the list of codes for the Unacceptable Principal Diagnosis edit. We are inviting public comments on our proposal.

(6) Poisoning by, Adverse Effects of, and Underdosing of Drugs, Medicaments and Biological Substances” (T36 through T50). The instructional note provided at this section states “Code first, for adverse effects, the nature of the adverse effect, such as:” and provides a list of examples. In addition, the FY 2017 ICD–10–CM Official Guidelines for Coding and Reporting at Section I.C.19.e.5.c., state that “Codes for underdosing should never be assigned as principal or first-listed codes.”
We identified 996 ICD–10–CM diagnosis codes that, as a result of the instructional note for adverse effects and the guideline for reporting diagnosis codes for underdosing, are not appropriate to report as a principal diagnosis. We are proposing to add the 996 ICD–10–CM diagnosis codes shown in Table 6P.1f. associated with this proposed rule (which is available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) to the list of codes for the Unacceptable Principal Diagnosis edit. We are inviting public comments on our proposal.

(7) Complications of Surgical and Medical Care, Not Elsewhere Classified (T80 Through T88)

We examined ICD–10–CM diagnosis codes in Chapter 19 (Injury and Poisoning) of the Classification Manual that fall within the range of code categories for “Complications of Surgical and Medical Care, Not Elsewhere Classified” (T80 through T88), specifically, at code category T81—Complications of procedures, not elsewhere classified. There is an instructional note at subcategory T81.12x—Postprocedural septic shock, subsequent encounter, which states, “Code first underlying infection.”

We identified two ICD–10–CM diagnosis codes in this subcategory that, as a result of the instructional note, are not appropriate to report as a principal diagnosis. These two ICD–10–CM codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–CM code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>T81.12XD</td>
<td>Postprocedural septic shock, subsequent encounter.</td>
</tr>
<tr>
<td>T81.12XS</td>
<td>Postprocedural septic shock, sequela.</td>
</tr>
</tbody>
</table>

We are proposing to add the two ICD–10–CM diagnosis codes shown in the table above to the list of codes for the Unacceptable Principal Diagnosis edit. We are inviting public comments on our proposal.

(8) Persons Encountering Health Services for Examinations (Z00 Through Z13)

We examined ICD–10–CM diagnosis codes in Chapter 21 (Factors Influencing Health Status) of the Classification Manual that fall within the range of code categories for “Persons Encountering Health Services for Examinations” (Z00 through Z13), specifically, at code category Z00—Encounter for general examination without complaint, suspected or reported diagnosis. The FY 2017 ICD–10–CM Official Guidelines for Coding and Reporting at Section I.C.21.c.16., state that the following ICD–10–CM Z-codes/categories may only be reported as the principal/first-listed diagnosis, except when there are multiple encounters on the same day and the medical records for the encounters are combined:

- Z00 (Encounter for general examination without complaint, suspected or reported diagnosis); except Z00.6 (Encounter for examination for normal comparison and control in clinical research program).

Therefore, diagnosis code Z00.6 should not be reported as a principal/first-listed diagnosis. We are proposing to add ICD–10–CM diagnosis code Z00.6 to the list of codes for the Unacceptable Principal Diagnosis edit. We are inviting public comments on our proposal.

To address a separate issue, we are proposing to remove the diagnosis codes under category Z05 (Encounter for observation and examination of newborn for suspected diseases and conditions ruled out) from the list of codes for the Unacceptable Principal Diagnosis edit. The FY 2017 ICD–10–CM Official Guidelines for Coding and Reporting at Section I.C.16.b. state the following:

- Assign a code from category Z05, Observation and evaluation of newborns and infants for suspected conditions ruled out, to identify those instances when a healthy newborn is evaluated for a suspected condition that is determined after study not to be present. Do not use a code from category Z05 when the patient has identified signs or symptoms of a suspected problem; in such cases code the sign or symptom.

- A code from category Z05 may also be assigned as a principal or first-listed code for readmissions or encounters when the code from category Z38 no longer applies. Codes from category Z05 are for use only for healthy newborns and infants for which no condition after study is found to be present.

- A code from category Z05 is to be used as a secondary code after the code from category Z38, Liveborn infants according to place of birth and type of delivery.

Therefore, the ICD–10–CM diagnosis codes under category Z05 are allowed to be reported as a principal diagnosis. We are proposing to remove the 14 ICD–10–CM diagnosis codes shown in the table below from the list of codes for the Unacceptable Principal Diagnosis edit.

<table>
<thead>
<tr>
<th>ICD–10–CM code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Z05.0</td>
<td>Observation and evaluation of newborn for suspected cardiac condition ruled out.</td>
</tr>
<tr>
<td>Z05.1</td>
<td>Observation and evaluation of newborn for suspected infectious condition ruled out.</td>
</tr>
<tr>
<td>Z05.2</td>
<td>Observation and evaluation of newborn for suspected neurological condition ruled out.</td>
</tr>
<tr>
<td>Z05.3</td>
<td>Observation and evaluation of newborn for suspected respiratory condition ruled out.</td>
</tr>
<tr>
<td>Z05.41</td>
<td>Observation and evaluation of newborn for suspected genetic condition ruled out.</td>
</tr>
<tr>
<td>Z05.42</td>
<td>Observation and evaluation of newborn for suspected metabolic condition ruled out.</td>
</tr>
<tr>
<td>Z05.43</td>
<td>Observation and evaluation of newborn for suspected immunologic condition ruled out.</td>
</tr>
<tr>
<td>Z05.5</td>
<td>Observation and evaluation of newborn for suspected gastrointestinal condition ruled out.</td>
</tr>
<tr>
<td>Z05.6</td>
<td>Observation and evaluation of newborn for suspected genitourinary condition ruled out.</td>
</tr>
<tr>
<td>Z05.71</td>
<td>Observation and evaluation of newborn for suspected skin and subcutaneous tissue condition ruled out.</td>
</tr>
<tr>
<td>Z05.72</td>
<td>Observation and evaluation of newborn for suspected musculoskeletal condition ruled out.</td>
</tr>
<tr>
<td>Z05.73</td>
<td>Observation and evaluation of newborn for suspected connective tissue condition ruled out.</td>
</tr>
<tr>
<td>Z05.8</td>
<td>Observation and evaluation of newborn for other specified suspected condition ruled out.</td>
</tr>
<tr>
<td>Z05.9</td>
<td>Observation and evaluation of newborn for unspecified suspected condition ruled out.</td>
</tr>
</tbody>
</table>
We are inviting public comments on our proposal. (9) Encounters for Other Specific Health Care (Z40 Through Z53)

We examined ICD–10–CM diagnosis codes in Chapter 21 (Factors Influencing Health Status) of the Classification Manual that fall within the range of code categories for “Encounters for Other Specific Health Care” (Z40 through Z53), specifically, at code category Z52—Donors of organs and tissues. The FY 2017 ICD–10–CM Official Guidelines for Coding and Reporting at Section I.C.21.c.16. state that the following Z-codes/categories may only be reported as the principal/ first-listed diagnosis, except when there are multiple encounters on the same day and the medical records for the encounters are combined:

• Z52 (Donors of organs and tissues); except Z52.9 (Donor of unspecified organ or tissue).

Therefore, ICD–10–CM diagnosis code Z52.9 should not be reported as a principal/first-listed diagnosis. We are proposing to add ICD–10–CM diagnosis code Z52.9 to the list of codes for the Unacceptable Principal Diagnosis edit. We are inviting public comments on our proposal.

(10) Persons Encountering Health Services in Other Circumstances (Z69 Through Z76)

We examined ICD–10–CM diagnosis codes in Chapter 21 (Factors Influencing Health Status) of the Classification Manual that fall within the range of code categories for “Persons Encountering Health Services in Other Circumstances” (Z69 through Z76), specifically, at subcategory Z71.8—Other specified counseling. Consistent with ICD–10–CM diagnosis codes Z71.81 (Spiritual or religious counseling) and Z71.89 (Other specified counseling), we are proposing to add new diagnosis code Z71.82 (Exercise counseling) to the list of codes for the Unacceptable Principal Diagnosis edit. We refer readers to Table 6A.—New Diagnosis Codes associated with this proposed rule (which is available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcutelinePatientPPS/index.html) for the list of new ICD–10–CM diagnosis codes finalized to date. We are inviting public comments on our proposal.

(11) Persons With Potential Health Hazards Related to Family and Personal History and Certain Conditions Influencing Health Status (Z77 Through Z99)

We examined ICD–10–CM diagnosis codes in Chapter 21 (Factors Influencing Health Status) of the Classification Manual that fall within the range of code categories for “Persons With Potential Health Hazards Related to Family and Personal History and Certain Conditions Influencing Health Status” (Z77 through Z99), specifically, at code category Z91.8—Other specified personal risk factors, not elsewhere classified. Consistent with ICD–10–CM diagnosis codes Z91.81 (History of falling), Z91.82 (Personal history of military deployment), and Z91.89 (Other specified personal risk factors, not elsewhere classified), we are proposing to add new ICD–10–CM diagnosis codes Z91.841 (Risk for dental caries, low), Z91.842 (Risk for dental caries, moderate), Z91.843 (Risk for dental caries, high), and Z91.849 (Unspecified risk for dental caries) to the list of codes for the Unacceptable Principal Diagnosis edit. We refer readers to Table 6A.—New Diagnosis Codes associated with this proposed rule (which is available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcutelinePatientPPS/index.html) for the list of new ICD–10–CM diagnosis codes finalized to date. We are inviting public comments on our proposal.

e. Future Enhancement

Similar to our discussion in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56843 through 56844), with the implementation of ICD–10, it is clear that there are several new concepts in the classification. Looking ahead to the needs and uses of coded data as the data continue to evolve from the reporting, collection, processing, coverage, payment and analysis aspects, we believe the need to ensure the accuracy of the coded data becomes increasingly significant.

The purpose of the MCE is to ensure that errors and inconsistencies in the coded data are recognized during Medicare claims processing. As we continue to evaluate the purpose and function of the MCE with respect to ICD–10, we encourage public input for future discussion. As we discussed in the FY 2017 IPPS/LTCH PPS final rule, we recognize a need to further examine the current (MS–DRG) to the definitions of those edits. We encourage public comments on whether there are additional concerns with the current edits, including specific edits or language that should be removed or revised, edits that should be combined, or new edits that should be added to assist in detecting errors or inaccuracies in the coded data.

11. Proposed Changes to Surgical Hierarchies

Some inpatient stays entail multiple surgical procedures, each one of which, occurring by itself, could result in assignment of the case to a different MS–DRG within the MDC to which the principal diagnosis is assigned. Therefore, it is necessary to have a decision rule within the GROUPE by which these cases are assigned to a single MS–DRG. The surgical hierarchy, an ordering of surgical classes from most resource-intensive to least resource-intensive, performs that function. Application of this hierarchy ensures that cases involving multiple surgical procedures are assigned to the MS–DRG associated with the most resource-intensive surgical class.

Because the relative resource intensity of surgical classes can shift as a function of MS–DRG recalcification and recalibrations, for FY 2018, we reviewed the surgical hierarchy of each MDC, as we have for previous reclassifications and recalibrations, to determine if the ordering of classes coincides with the intensity of resource utilization. A surgical class can be composed of one or more MS–DRGs. For example, in MDC 11, the surgical class “kidney transplant” consists of a single MS–DRG (MS–DRG 652) and the class “major bladder procedures” consists of three MS–DRGs (MS–DRGs 653, 654, and 655). Consequently, in many cases, the surgical hierarchy has an impact on more than one MS–DRG. The methodology for determining the most resource-intensive surgical class involves weighting the average resources for each MS–DRG by frequency to determine the weighted average resources for each surgical class. For example, assume surgical class A includes MS–DRGs 001 and 002 and surgical class B includes MS–DRGs 003, 004, and 005. Assume also that the average costs of MS–DRG 001 are higher than that of MS–DRG 003, but the average costs of MS–DRGs 004 and 005 are higher than the average costs of MS–DRG 002. To determine whether surgical class A should be higher or lower than surgical class B in the surgical hierarchy, we would weigh the average costs of each MS–DRG in the class by frequency (that is, by the number of cases in the MS–DRG) to determine average resource...
consumption for the surgical class. The surgical classes would then be ordered from the class with the highest average resource utilization to that with the lowest, with the exception of “other O.R. procedures” as discussed in this rule.

This methodology may occasionally result in assignment of a case involving multiple procedures to the lower-weighted MS–DRG (in the highest, most resource-intensive surgical class) of the available alternatives. However, given that the logic underlying the surgical hierarchy provides that the GROUPER search for the procedure in the most resource-intensive surgical class, in cases involving multiple procedures, this result is sometimes unavoidable.

We note that, notwithstanding the foregoing discussion, there are a few instances when a surgical class with a lower average cost is ordered above a surgical class with a higher average cost. For example, the “other O.R. procedures” surgical class is uniformly ordered last in the surgical hierarchy of each MDC in which it occurs, regardless of the fact that the average costs for the MS–DRG or MS–DRGs in that surgical class may be higher than those for other surgical classes in the MDC. The “other O.R. procedures” class is a group of procedures that are only infrequently related to the diagnoses in the MDC, but are still occasionally performed on patients with cases assigned to the MDC with these diagnoses. Therefore, assignment to these surgical classes should only occur if no other surgical class more closely related to the diagnoses in the MDC is appropriate.

A second example occurs when the difference between the average costs for two surgical classes is very small. We have found that small differences generally do not warrant reordering of the hierarchy because, as a result of reassigning cases on the basis of the hierarchy change, the average costs are likely to shift such that the higher-ordered surgical class has lower average costs than the class ordered below it.

We received a request to examine a case involving the principal procedure for excision of pituitary gland (ICD–10–PCS code 0GB0ZZ Excision of pituitary gland, open approach) with a secondary procedure for harvesting of a fat graft (ICD–10–PCS code 0JB80ZZ Excision of abdomen subcutaneous tissue and fascia, open approach) to treat a condition of pituitary adenoma (ICD–10–CM diagnosis code D35.2 (Benign neoplasm of pituitary gland)) and the resulting sela turcica defect. The requestor expressed concern regarding the procedure code for harvesting of the fat graft in the secondary position driving the MS–DRG assignment versus the principal procedure of the excision of pituitary gland.

We analyzed the codes provided by the requestor in the GROUPER to determine if we could duplicate the requestor’s findings. The findings from our analysis were consistent with the requestor’s findings. Our clinical advisors reviewed this issue and agreed that it should be the procedure code for excision of the pituitary gland that is used to determine the MS–DRG assignment in this scenario and not the harvesting of the fat graft procedure code.

Therefore, in this FY 2018 IPPS/LTC PPS proposed rule, we are proposing to move MS–DRGs 614 and 615 above MS–DRGs 622, 623, and 624 in the surgical hierarchy to enable more appropriate MS–DRG assignment for these types of cases.

We are inviting public comments on our proposal.

12. Proposed Changes to the MS–DRG Diagnosis Codes for FY 2018

a. Background of the CC List and the CC Exclusions List

Under the IPPS MS–DRG classification system, we have developed a standard list of diagnoses that are considered CCs. Historically, we developed this list using physician panels that classified each diagnosis code based on whether the diagnosis, when present as a secondary condition, would be considered a substantial complication or comorbidity. A substantial complication or comorbidity was defined as a condition that, because of its presence with a specific principal diagnosis, would cause an increase in the length-of-stay by at least 1 day in at least 75 percent of the patients. However, depending on the principal diagnosis of the patient, some diagnoses on the basic list of complications and comorbidities may be excluded if they are closely related to the principal diagnosis. In FY 2008, we evaluated each diagnosis code to determine its impact on resource use and to determine the most appropriate CC, subclassification (non-CC, CC, or MCC) assignment. We refer readers to sections II.D.2. and 3. of the preamble of the FY 2008 IPPS final rule with comment period for a discussion of the refinement of CCs in relation to the MS–DRGs we adopted for FY 2008 (72 FR 47152 through 47171).

b. Proposed Additions and Deletions to the Diagnosis Code Severity Levels for FY 2018

The following tables identifying the proposed additions and deletions to the MCC severity levels list and the proposed additions and deletions to the CC severity levels list for FY 2018 are available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html.

Table 6I.1—Proposed Additions to the MCC List—FY 2018; Table 6I.2—Proposed Deletions to the MCC List—FY 2018; Table 6J.1—Proposed Additions to the CC List—FY 2018; and Table 6J.2—Proposed Deletions to the CC List—FY 2018.

We are inviting public comments on our proposed severity level designations for the diagnosis codes listed in Table 6I.1 and Table 6J.1. We note that, for Table 6I.2 and Table 6J.2., the proposed deletions are a result of code expansions. Therefore, the diagnosis codes on these lists are no longer valid codes, effective FY 2018. For example, diagnosis code O00.10 (Tubal pregnancy without intrauterine pregnancy) is a current CC for FY 2017 under Version 34 of the ICD–10 MS–DRGs. Effective FY 2018, under Version 35 of the ICD–10 MS–DRGs, this single code has been expanded into three diagnosis codes to include laterality (left/right) and an unspecified option with the addition of a sixth character. Therefore, diagnosis code O00.10 is included in Table 6I.2, for deletion from the CC list because it is no longer a valid code in FY 2018.

c. Principal Diagnosis Is Its Own CC or MCC

CMS’ initial goal in developing the ICD–10 MS–DRGs was to ensure that a patient case was assigned to the same MS–DRG, regardless of whether the patient record was to be coded in ICD–9–CM or ICD–10. When certain ICD–10–CM combination codes are reported as a principal diagnosis, it implies that a CC or MCC is present. This occurs as a result of evaluating the cluster of ICD–
9-CM codes that would have been coded on an ICD-9-CM record. If one of the ICD-9-CM codes in the cluster was a CC or an MCC, the single ICD-10-CM combination code used as a principal diagnosis also must imply that the CC or MCC is present.

The ICD-10-CM diagnosis codes to which this logic applies are included in Appendix J of the ICD-10 MS-DRG Version 34 Definitions Manual (which is available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/IPPS-Final-Rule-Home-Page-Items/IPPS-Final-Rule-Data-Files.html?DLPage=1&DLSort=0&DLSortDir=ascending).

Appendix J includes two lists: Part 1 is the list of principal diagnosis codes where the ICD-10-CM code is its own MCC. Part 2 is the list of principal diagnosis codes where the ICD-10-CM code is its own CC. Part 1 of Appendix J corresponds to Table 6L.—Principal Diagnosis Is Its Own MCC List, and Part 2 of Appendix J corresponds to Table 6M.—Principal Diagnosis Is Its Own CC List.

We received a request to add the ICD-10-CM diagnosis codes for acute myocardial infarction, decompensated heart failure and specified forms of shock, which are currently designated as a CC or an MCC when reported as a secondary diagnosis, to Table 6L.—Principal Diagnosis Is Its Own MCC List. According to the requestor, the addition of these codes to the list is necessary to pay bundled payment initiatives and so that facilities that accept these patients in transfer have resources to care for them.

The purpose of the Principal Diagnosis Is Its Own CC or MCC Lists was to ensure consistent MS-DRG assignment between the ICD-9-CM and ICD-10 MS-DRGs due to the clusters and combination codes. There are a number of other ICD-10-CM combination codes that, due to their prior designation as a CC or an MCC when reported as a secondary diagnosis, are not on either of these lists. Having multiple lists for CC and MCC diagnoses when reported as a principal and/or secondary diagnosis may not provide an accurate representation of resource utilization for the MS-DRGs. As discussed in further detail below, we have plans to conduct a comprehensive review of the CC and MCC lists for FY 2019. We believe the results of that review will help to inform the future of these lists.

Therefore, we are not proposing to add the ICD-10-CM diagnosis codes for acute myocardial infarction, decompensated heart failure and specified forms of shock to Table 6L.—Principal Diagnosis Is Its Own MCC List. In addition, we are not proposing any changes to Table 6L.—Principal Diagnosis Is Its Own MCC List and Table 6M.—Principal Diagnosis Is Its Own CC List. We are inviting public comments on our proposal to maintain the existing lists of principal diagnosis codes in Tables 6L. and 6M for FY 2018.

d. Proposed CC Exclusions List for FY 2018

In the September 1, 1987 final notice (52 FR 33143), concerning changes to the DRG classification system, we modified the GROUPER logic so that certain diagnoses included on the standard list of CCs would not be considered valid CCs in combination with a particular principal diagnosis. We created the CC Exclusions List for the following reasons: (1) To preclude coding of CCs for closely related conditions; (2) to preclude duplicative or inconsistent coding from being treated as CCs; and (3) to ensure that cases are appropriately classified between the complicated and uncomplicated DRGs in a pair. As previously indicated, we developed a list of diagnoses, using physician panels, to include those diagnoses that, when present as a secondary condition, would be considered a substantial complication or comorbidity.

In previous years, we made changes to the list of CCs, either by adding new CCs or deleting CCs already on the list. In the May 19, 1987 proposed notice (52 FR 18877) and the September 1, 1987 final notice (52 FR 33154), we explained that the excluded secondary diagnoses were established using the following five principles:

- Chronic and acute manifestations of the same condition should not be considered CCs for one another;
- Specific and nonspecific (that is, not otherwise specified (NOS)) diagnosis codes for the same condition should not be considered CCs for one another;
- Codes for the same condition that cannot coexist, such as partial/total, unilaterial/bilateral, obstructed/unobstructed, and benign/malignant, should not be considered CCs for one another;
- Codes for the same condition in anatomically proximal sites should not be considered CCs for one another; and
- Closely related conditions should not be considered CCs for one another.

The creation of the CC Exclusions List was a major project involving hundreds of codes. We have continued to review the remaining CCs to identify additional exclusions and to remove diagnoses from the master list that have been shown not to meet the definition of a CC. We refer readers to the FY 2014 IPPS/LTCH PPS final rule (78 FR 50541 through 50544) for detailed information regarding revisions that were made to the CC and CC Exclusion Lists under the ICD-9-CM MS-DRGs.

For FY 2018, we are proposing changes to the ICD-10 MS-DRGs Version 35 CC Exclusion List. Therefore, we have developed Table 6G.1.—Proposed Secondary Diagnosis Order Additions to the CC Exclusions List—FY 2018; Table 6G.2.—Proposed Principal Diagnosis Order Additions to the CC Exclusions List—FY 2018; Table 6H.1.—Proposed Secondary Diagnosis Order Deletions to the CC Exclusions List—FY 2018; and Table 6H.2.—Proposed Principal Diagnosis Order Deletions to the CC Exclusions List—FY 2018. Each of these principal diagnosis codes for which there is a CC exclusion is shown in Table 6G.2. with an asterisk and the conditions that will not count as a CC are provided in an indented column immediately following the affected principal diagnosis. Beginning with discharges on or after October 1 of each year, the indented diagnoses are not recognized by the GROUPER as valid CCs for the asterisked principal diagnoses. Tables 6G. and 6H. associated with this proposed rule are available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html.

To identify new, revised and deleted diagnosis and procedure codes, for FY 2018, we have developed Table 6A.—New Diagnosis Codes, Table 6B.—New Procedure Codes, Table 6C.—Invalid Diagnosis Codes, Table 6D.—Invalid Procedure Codes, Table 6E.—Revised Diagnosis Code Titles, and Table 6F.—Revised Procedure Code Titles for this proposed rule.

These tables are not published in the Addendum to this proposed rule but are available via the Internet on the CMS Web site at: (https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) as described in section VI. of the Addendum to this proposed rule. As discussed in section II.F.15. of the preamble of this proposed rule, the code titles are adopted as part of the ICD-10 (previously ICD-9-CM) Coordination and Maintenance Committee process. Therefore, although we publish the code titles in the IPPS proposed and final rules, they are not subject to comment in this proposed rule. We are inviting public comments on the MDC and MS-DRG assignments for the new
diagnosis and procedure codes as set forth in Table 6A.—New Diagnosis Codes and Table 6B.—New Procedure Codes. In addition, we are inviting public comments on the proposed severity level designations for the new diagnosis codes as set forth in Table 6A and the proposed O.R. status for the new procedure codes as set forth in Table 6B.

13. Comprehensive Review of CC List for FY 2019

In the FY 2008 IPPS final rule (72 FR 47153 through 47175), we discussed our efforts to better recognize severity of illness which began with a comprehensive review of the CC list and, ultimately, the implementation of the MS–DRGs. Similar to the analysis that was performed at that time, we are providing the public with notice of our plans to conduct a comprehensive review of the CC and MCC lists for FY 2019.

As a result of the time that has elapsed since that review and changes to how inpatient care is currently delivered, we plan to analyze if further refinements to these lists are warranted. For example, over the past several years, there has been a steady increase in the proportion of cases grouping to the MS–DRGs with an MCC severity level than had previously occurred. Our evaluation will assist in determining if the conditions designated as an MCC continue to represent significant increases in resource utilization that support the MCC designation.

We currently utilize a statistical algorithm to determine the impact on resource use of each secondary diagnosis. Each diagnosis for which Medicare data are available is evaluated to determine its impact on resource use and to determine the most appropriate CC subclass (non-CC, CC, or MCC) assignment. In order to make this determination, the average costs for each subset of cases is compared to the expected costs for cases in that subset. The following format is used to evaluate each diagnosis:

<table>
<thead>
<tr>
<th>Code</th>
<th>Diagnosis</th>
<th>Cnt1</th>
<th>C1</th>
<th>Cnt2</th>
<th>C2</th>
<th>Cnt3</th>
<th>C3</th>
</tr>
</thead>
</table>

Count (Cnt) is the number of patients in each subset and C1, C2, and C3 are a measure of the impact on resource use of patients in each of the subsets. The C1, C2, and C3 values are a measure of the ratio of average costs for patients with these conditions to the expected average costs across all cases. The C1 value reflects a patient with no other secondary diagnosis or with all other secondary diagnoses that are non-CCs. The C2 value reflects a patient with at least one other secondary diagnosis that is a CC but none that is an MCC. The C3 value reflects a patient with at least one other secondary diagnosis that is an MCC. A value close to 1.0 in the C1 field would suggest that the code produces the same expected value as a non-CC diagnosis. That is, average costs for the case are similar to the expected average costs for that subset and the diagnosis is not expected to increase resource usage. A higher value in the C1 (or C2 and C3) field suggests more resource usage is associated with the diagnosis and an increased likelihood that it is more like a CC or major CC than a non-CC. Thus, a value close to 2.0 suggests the condition is more like a CC than a non-CC but not as significant in resource usage as an MCC. A value close to 3.0 suggests the condition is expected to consume resources more similar to an MCC than a CC or non-CC. For example, a C1 value of 1.8 for a secondary diagnosis means that for the subset of patients who have the secondary diagnosis and have either no other secondary diagnosis present, or all the other secondary diagnoses present are non-CCs, the impact on resource use of the secondary diagnoses is greater than the expected value for a non-CC by an amount equal to 80 percent of the difference between the expected value of a CC and a non-CC (that is, the impact on resource use of the secondary diagnosis is closer to a CC than a non-CC).

We are inviting public comments regarding other possible ways we can incorporate meaningful indicators of clinical severity.

14. Review of Procedure Codes in MS DRGs 981 Through 986; and 987 Through 989

Each year, we review cases assigned to MS–DRGs 981, 982, and 983 (Extensive O.R. Procedure Unrelated to Principal Diagnosis with MCC, with CC, and without CC/MCC, respectively); MS–DRGs 984, 985, and 986 (Prostatic O.R. Procedure Unrelated to Principal Diagnosis with MCC, with CC, and without CC/MCC, respectively); and MS–DRGs 987, 988, and 989 (Nonextensive O.R. Procedure Unrelated to Principal Diagnosis with MCC, with CC, and without CC/MCC, respectively) or MS–DRGs 981 through 983 and 987 through 989 assigned to those discharges in which the only procedures performed are unrelated to the principal diagnosis.

We refer the reader to the FY 2017 IPPS/LTCH PPS final rule (81 FR 56847 through 56848) for a discussion of the movement and redesignation of procedure codes from MS–DRGs 984 through 986 related to the transition of the ICD–10 MS–DRGs.

Our review of MedPAR claims data showed that there are no cases that merited movement or should logically be reassigned from ICD–10 MS–DRGs 984 through 986 to any of the other MDCs for FY 2018. Therefore, for FY 2018, we are not proposing to change the procedures assigned among these MS–DRGs. We are inviting public comments on our proposal to maintain the current structure of these MS–DRGs.

a. Moving Procedure Codes From MS–DRGs 981 Through 983 or MS–DRGs 987 Through 989 Into MDCs

We annually conduct a review of procedures producing assignment to MS–DRGs 981 through 983 (Extensive O.R. Procedure Unrelated to Principal Diagnosis with MCC, with CC, and without CC/MCC, respectively) or MS–DRGs 987 through 989 (Nonextensive O.R. Procedure Unrelated to Principal Diagnosis with MCC, with CC, and without CC/MCC, respectively) on the basis of volume, by procedure, to see if it would be appropriate to move procedure codes out of these MS–DRGs.
into one of the surgical MS–DRGs for the MDC into which the principal diagnosis falls. The data are arrayed in two ways for comparison purposes. We look at a frequency count of each major operative procedure code. We also compare procedures across MDCs by volume of procedure codes within each MDC.

We identify those procedures occurring in conjunction with certain principal diagnoses with sufficient frequency to justify adding them to one of the surgical MS–DRGs for the MDC in which the diagnosis falls. Upon review of the claims data from the December 2016 update of the FY 2016 MedPAR file, we did not find any cases that merited movement or that should logically be assigned to any of the other MDCs. Therefore, for FY 2018, we are not proposing to remove any procedures from MS–DRGs 981 through 983 or MS–DRGs 987 through 989 into one of the surgical MS–DRGs for the MDC into which the principal diagnosis is assigned. We are inviting public comments on our proposal to maintain the current structure of these MS–DRGs.

b. Reassignment of Procedures Among MS–DRGs 981 Through 983, 984 Through 986, and 987 Through 989

We also review the list of ICD–10–PCS procedures that, when in combination with their principal diagnosis code, result in assignment to MS–DRGs 981 through 983, 984 through 986, or 987 through 989, to ascertain whether any of those procedures should be reassigned from one of those groups of MS–DRGs to another of the three groups of MS–DRGs based on average costs and the length of stay. We look at the data for trends such as shifts in treatment practice or reporting practice that would make the resulting MS–DRG assignment illogical. If we find these shifts, we would propose to move cases to keep the MS–DRGs clinically similar or to provide payment for the cases in a similar manner. Generally, we move only those procedures for which we have an adequate number of discharges to analyze the data.

Based on the results of our review of the December 2016 update of the FY 2016 MedPAR file, we are proposing to reassign the procedure codes currently assigned to MS–DRGs 984 through 986 (Prostatic O.R. Procedure Unrelated to Principal Diagnosis with MCC, with CC and without CC/MCC, respectively) to MS–DRGs 987 through 989 (Non–extensive O.R. Procedure Unrelated to Principal Diagnosis with MCC, with CC and without CC/MCC, respectively). As shown in the table below, we found a total of 1,001 cases in MS–DRGs 984 through 986 with an average length-of-stay of 7.5 days and average costs of $16,539. In MS–DRGs 987 through 989, we found a total of 17,772 cases, with an average length of stay of 7.5 days and average costs of $16,193.

### O.R. Procedures Unrelated to Principal Diagnosis

<table>
<thead>
<tr>
<th>MS–DRG</th>
<th>Number of cases</th>
<th>Average length of stay</th>
<th>Average costs</th>
</tr>
</thead>
<tbody>
<tr>
<td>MS–DRGs 984, 985 and 986</td>
<td>1,001</td>
<td>7.5</td>
<td>$16,539</td>
</tr>
<tr>
<td>MS–DRGs 987, 988 and 989</td>
<td>17,772</td>
<td>7.5</td>
<td>$16,193</td>
</tr>
</tbody>
</table>

The claims data demonstrate that it is no longer necessary to maintain a separate set of MS–DRGs specifically for the prostatic O.R. procedures. The average length of stay of 7.5 days is identical in both sets of MS–DRGs and the average costs are very similar with a difference of only $346. Our clinical advisors reviewed the data and support movement of these 1,001 cases into the nonextensive O.R. procedures MS–DRGs. They noted that treatment practices have shifted since the inception of the prostatic O.R. procedures grouping and the average costs are in alignment.

Therefore, for FY 2018, we are proposing to reassign the prostatic O.R. procedure codes from MS–DRGs 984 through 986 to MS–DRGs 987 through 989 and to delete MS–DRGs 984, 985 and 986 because they would no longer be needed as a result of this proposed movement. We are inviting public comments on our proposals.

15. Proposed Changes to the ICD–10–CM and ICD–10–PCS Coding Systems

In September 1985, the ICD–9–CM Coordination and Maintenance Committee was formed. This is a Federal interdepartmental committee, co-chaired by the National Center for Health Statistics (NCHS), the Centers for Disease Control and Prevention, and CMS, charged with maintaining and updating the ICD–9–CM system. The final update to ICD–9–CM codes was made on October 1, 2013. Thereafter, the name of the Committee was changed to the ICD–10 Coordination and Maintenance Committee, effective with the March 19–20, 2014 meeting. The ICD–10 Coordination and Maintenance Committee addresses updates to the ICD–10–CM and ICD–10–PCS coding systems. The Committee is jointly responsible for approving coding changes, and developing errata, addenda, and other modifications to the coding systems to reflect newly developed procedures and technologies and newly identified diseases. The Committee is also responsible for promoting the use of Federal and non-Federal educational programs and other communication techniques with a view toward standardizing coding applications and upgrading the quality of the classification system.

The official list of ICD–10–CM and ICD–10–PCS codes can be found on the CMS Web site at: [http://www.cms.gov/Medicare/Coding/ICD10ProviderDiagnosticCodes/codes.html](http://www.cms.gov/Medicare/Coding/ICD10ProviderDiagnosticCodes/codes.html). The NCHS has lead responsibility for the ICD–10–CM and ICD–9–CM diagnosis codes included in the Tabular List and Alphabetic Index for Diseases, while CMS has lead responsibility for the ICD–10–PCS and ICD–9–CM procedure codes included in the Tabular List and Alphabetic Index for Procedures.

The Committee encourages participation in the previously mentioned process by health-related organizations. In this regard, the Committee holds public meetings for discussion of educational issues and proposed coding changes. These meetings provide an opportunity for representatives of recognized organizations in the coding field, such as the American Health Information Management Association (AHIMA), the American Hospital Association (AHA), and various physician specialty groups, as well as individual physicians, health information management professionals, and other members of the public, to
contribute ideas on coding matters. After considering the opinions expressed at the public meetings and in writing, the Committee formulates recommendations, which then must be approved by the agencies.

The Committee presented proposals for coding changes for implementation in FY 2018 at a public meeting held on September 13–14, 2016, and finalized the coding changes after consideration of comments received at the meetings and in writing by November 13, 2016. The Committee held its 2017 meeting on March 7–8, 2017. The deadline for submitting comments on these code proposals was April 7, 2017. It was announced at this meeting that any new ICD–10–CM/PCS codes for which there was consensus of public support and for which complete tabular and indexing changes would be made by May 2017 would be included in the October 1, 2017 update to ICD–10–CM/ICD–10–PCS. As discussed in earlier sections of the preamble of this proposed rule, there are new, revised, and deleted ICD–10–CM diagnosis codes and ICD–10–PCS procedure codes that are captured in Table 6A.—New Diagnosis Codes, Table 6B.—New Procedure Codes, Table 6C.—Invalid Diagnosis Codes, Table 6D.—Invalid Procedure Codes, Table 6E.—Revised Diagnosis Code Titles, and Table 6F.—Revised Procedure Code Titles for this proposed rule, which are available via the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Coding/index.html. Because of the length of these tables, they are not published in the Addendum to this proposed rule. Rather, they are available via the Internet as discussed in section VI. of the Addendum to this proposed rule.

Live Webcast recordings of the discussions of procedure codes at the Committee’s September 13–14, 2016 meeting and March 7–8, 2017 meeting can be obtained from the CMS Web site at: http://cms.hhs.gov/Medicare/Coding/ICD9ProviderDiagnosticCodes/index.html?redirect=icd9ProviderDiagnosticCodes03_meetings.asp. The minutes of the discussions of diagnosis codes at the September 13–14, 2016 meeting and March 7–8, 2017 meeting can be found at: http://www.cdc.gov/nchs/icd/icd10cm_maintenance.html. These Web sites also provide detailed information about the Committee, including information on requesting a new code, attending a Committee meeting, and timeline requirements and meeting dates.

We encourage commenters to address suggestions on coding issues involving diagnosis codes to: Donna Pickett, Co-Chairperson, ICD–10 Coordination and Maintenance Committee, NCHS, Room 2402, 3311 Toledo Road, Hyattsville, MD 20782. Comments may be sent by Email to nchsicd10@cdc.gov.

Questions and comments concerning the procedure codes should be addressed to: Patricia Brooks, Co-Chairperson, ICD–10 Coordination and Maintenance Committee, CMS, Center for Medicare Management, Hospital and Ambulatory Policy Group, Division of Acute Care, C4–08–06, 7500 Security Boulevard, Baltimore, MD 21244–1850. Comments may be sent by Email to ICDProcedureCodeRequest@cms.hhs.gov.

In the September 7, 2001 final rule implementing the IPPS new technology add-on payments (66 FR 46906), we indicated we would attempt to include proposals for procedure codes that would describe new technology discussed and approved at the Spring meeting as part of the code revisions effective the following October. Section 503(a) of Public Law 108–173 included a requirement for updating diagnosis and procedure codes twice a year instead of a single update on October 1 of each year. This requirement was included as part of the amendments to the Act relating to recognition of new technology under the IPPS. Section 503(a) amended section 1886(d)(5)(K) of the Act by adding a clause (vii) which states that the Secretary shall provide for the addition of new diagnosis and procedure codes on April 1 of each year, but the addition of such codes shall not require the Secretary to adjust the payment (or diagnosis-related group classification) until the fiscal year that begins after such date. This requirement improves the recognition of new technologies under the IPPS system by providing information on these new technologies at an earlier date. Data will be available 6 months earlier than would be possible with updates occurring only once a year on October 1.

While section 1886(d)(5)(K)(vii) of the Act states that the addition of new diagnosis and procedure codes on April 1 of each year shall not require the Secretary to adjust the payment, or DRG classification, under section 1886(d) of the Act until the fiscal year that begins after such date, we have to update the DRG software and other systems in order to recognize and accept the new codes. We also publicize the code changes and the need for a mid-year systems update by providers to identify the new codes and to obtain the new code books and encoder updates, and make other system changes in order to identify and report the new codes.

The ICD–10 (previously ICD–9–CM) Coordination and Maintenance Committee holds its meetings in the spring and fall in order to update the codes and the applicable payment and reporting systems by October 1 of each year. Items are placed on the agenda for the Committee meeting if the request is received at least 2 months prior to the meeting. This requirement allows time for staff to review and research the coding issues and prepare material for discussion at the meeting. It also allows time for the topic to be publicized in meeting announcements in the Federal Register as well as on the CMS Web site. Final decisions on code title revisions are currently made by March 1 so that these titles can be included in the IPPS proposed rule. A complete addendum describing details of all diagnosis and procedure coding changes, both tabular and index, is published on the CMS and NCHS Web sites in June of each year. Publishers of coding books and software use this information to modify their products that are used by health care providers. This 5-month time period has proved to be necessary for hospitals and other providers to update their systems.

A discussion of this timeline and the need for changes are included in the December 4–5, 2005 ICD–9–CM Coordination and Maintenance Committee Meeting minutes. The public agreed that there was a need to hold the fall meetings earlier, in September or October, in order to meet the new implementation dates. The public provided comment that additional time would be needed to update hospital systems and obtain new code books and coding software. There was considerable concern expressed about the impact this new April update would have on providers.

In the FY 2005 IPPS final rule, we implemented section 1886(d)(5)(K)(vii) of the Act, as added by section 503(a) of Public Law 108–173, by developing a mechanism for approving, in time for the April update, diagnosis and procedure code revisions needed to describe new technologies and medical services for purposes of the new technology add-on payment process. We also established the following process for making these determinations. Topics considered during the Fall ICD–10 (previously ICD–9–CM) Coordination and Maintenance Committee meeting are considered for an April 1 update if a strong and convincing case is made by the requester at the Committee’s public meeting. The requester must justify the reason why a new code is needed in April for purposes of the new
technology process. The participants at the meeting and those reviewing the Committee meeting summary report are provided the opportunity to comment on this expedited request. All other topics are considered for the October 1 update. Participants at the Committee meeting are encouraged to comment on all such requests. There were no requests approved for an expedited April 1, 2017 implementation of a code at the September 13–14, 2016 Committee meeting. Therefore, there were no new codes implemented on April 1, 2017.


CMS also sends copies of all ICD–10–CM and ICD–10–PCS coding changes to its Medicare contractors for use in updating their systems and providing education to providers.

The code titles are adopted as part of the ICD–10 (previously ICD–9–CM) Coordination and Maintenance Committee process. Therefore, although we publish the code titles in the IPPS proposed and final rules, they are not subject to comment in the proposed or final rules.

The following chart shows the number of ICD–10–CM and ICD–10–PCS codes and code changes since FY 2016 when ICD–10 was implemented.

### TOTAL NUMBER OF CODES AND CHANGES IN TOTAL NUMBER OF CODES PER FISCAL YEAR ICD–10–CM AND ICD–10–PCS CODES

<table>
<thead>
<tr>
<th>Fiscal year</th>
<th>Number</th>
<th>Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>FY 2016:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ICD–10–CM</td>
<td>69,823</td>
<td></td>
</tr>
<tr>
<td>ICD–10–PCS</td>
<td>71,974</td>
<td></td>
</tr>
<tr>
<td>FY 2017:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ICD–10–CM</td>
<td>71,486</td>
<td>+1,663</td>
</tr>
<tr>
<td>ICD–10–PCS</td>
<td>75,799</td>
<td>+3,815</td>
</tr>
<tr>
<td>FY 2018:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ICD–10–CM</td>
<td>71,772</td>
<td>+286</td>
</tr>
<tr>
<td>ICD–10–PCS</td>
<td>78,299</td>
<td>+2,510</td>
</tr>
</tbody>
</table>

As mentioned previously, the public is provided the opportunity to comment on any requests for new diagnosis or procedure codes discussed at the ICD–10 Coordination and Maintenance Committee meeting.

At the September 12–13, 2016 and March 7–8, 2017 Committee meetings, we discussed any requests we had received for new ICD–10–CM diagnosis codes and ICD–10–PCS procedure codes that were to be implemented on October 1, 2017. We invited public comments on any code requests discussed at the September 12–13, 2016 and March 7–8, 2017 Committee meetings for implementation as part of the October 1, 2017 update. The deadline for commenting on code proposals discussed at the September 12–13, 2016 Committee meeting was November 13, 2016. The deadline for commenting on code proposals discussed at the March 7–8, 2017 Committee meeting was April 7, 2017.

16. Proposed Replaced Devices Offered Without Cost or With a Credit

a. Background

In the FY 2008 IPPS final rule with comment period (72 FR 47246 through 47251), we discussed the topic of Medicare payment for devices that are replaced without cost or where credit for a replaced device is furnished to the hospital. We implemented a policy to reduce a hospital’s IPPS payment for certain MS–DRGs where the implantation of a device that has been recalled determined the base MS–DRG assignment. At that time, we specified that we will reduce a hospital’s IPPS payment for those MS–DRGs where the hospital received a credit for a replaced device equal to 50 percent or more of the cost of the device.

In the FY 2012 IPPS/LTCH PPS final rule (76 FR 51556 through 51557), we clarified this policy to state that the policy applies if the hospital received a credit equal to 50 percent or more of the cost of the replacement device and issued instructions to hospitals accordingly.

b. Proposed Changes for FY 2018

For FY 2018, we are not proposing to add any MS–DRGs to the policy for replaced devices offered without cost or with a credit. We are proposing to continue to include the existing MS–DRGs currently subject to the policy as displayed in the table below.

<table>
<thead>
<tr>
<th>MDC</th>
<th>MS–DRG</th>
<th>MS–DRG title</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-MDC</td>
<td>001</td>
<td>Heart Transplant or Implant of Heart Assist System with MCC.</td>
</tr>
<tr>
<td>Pre-MDC</td>
<td>002</td>
<td>Heart Transplant or Implant of Heart Assist System without MCC.</td>
</tr>
<tr>
<td>1</td>
<td>023</td>
<td>Craniotomy with Major Device Implant/Acute Complex CNS Principal Diagnosis with MCC or Chemo Implant.</td>
</tr>
<tr>
<td>1</td>
<td>024</td>
<td>Craniotomy with Major Device Implant/Acute Complex CNS Principal Diagnosis without MCC.</td>
</tr>
<tr>
<td>1</td>
<td>025</td>
<td>Craniotomy &amp; Endovascular Intracranial Procedures with MCC.</td>
</tr>
<tr>
<td>1</td>
<td>026</td>
<td>Craniotomy &amp; Endovascular Intracranial Procedures with CC.</td>
</tr>
<tr>
<td>1</td>
<td>027</td>
<td>Craniotomy &amp; Endovascular Intracranial Procedures without CC/MCC.</td>
</tr>
<tr>
<td>1</td>
<td>040</td>
<td>Peripheral, Cranial Nerve &amp; Other Nervous System Procedures with MCC.</td>
</tr>
<tr>
<td>1</td>
<td>041</td>
<td>Peripheral, Cranial Nerve &amp; Other Nervous System Procedures with CC or Peripheral Neurostimulator.</td>
</tr>
<tr>
<td>1</td>
<td>042</td>
<td>Peripheral, Cranial Nerve &amp; Other Nervous System Procedures without CC/MCC.</td>
</tr>
<tr>
<td>3</td>
<td>129</td>
<td>Major Head &amp; Neck Procedures with CC/MCC or Major Device.</td>
</tr>
<tr>
<td>3</td>
<td>130</td>
<td>Major Head &amp; Neck Procedures without CC/MCC.</td>
</tr>
<tr>
<td>5</td>
<td>215</td>
<td>Other Heart Assist System Implant.</td>
</tr>
<tr>
<td>5</td>
<td>216</td>
<td>Cardiac Valve &amp; Other Major Cardiothoracic Procedure with Cardiac Catheterization with MCC.</td>
</tr>
<tr>
<td>5</td>
<td>217</td>
<td>Cardiac Valve &amp; Other Major Cardiothoracic Procedure with Cardiac Catheterization without MCC.</td>
</tr>
<tr>
<td>5</td>
<td>218</td>
<td>Cardiac Valve &amp; Other Major Cardiothoracic Procedure with Cardiac Catheterization without CC/MCC.</td>
</tr>
<tr>
<td>5</td>
<td>219</td>
<td>Cardiac Valve &amp; Other Major Cardiothoracic Procedure without Cardiac Catheterization with MCC.</td>
</tr>
</tbody>
</table>
We are soliciting public comments on our proposal to continue to include the existing MS–DRGs currently subject to the policy for replaced devices offered without cost or with credit and to not add any additional MS–DRGs to the policy. We note that, as discussed in section II.F.2.b. and in section II.F.5.a. of the preamble of this proposed rule, we are proposing to revise the titles for MS–DRG 023 and MS–DRGs 469 and 470. We refer readers to those discussions of the specific proposed MS–DRG titles. The final list of MS–DRGs subject to the payment policy for devices provided at no cost or with credit for FY 2018 will be listed in the FY 2018 IPPS/LTCH PPS final rule, as well as issued to providers through guidance and instructions in the form of a Change Request (CR).

17. Other Policy Changes: Other Operating Room (O.R.) and Non-O.R. Issues


For this FY 2018 IPPS/LTCH PPS proposed rule, we continued our efforts to address the recommendations for consideration that we received in response to some of the proposals set forth in the FY 2017 IPPS/LTCH PPS proposed rule pertaining to changing the designation of ICD–10–PCS procedure codes from O.R. procedures to non-O.R. procedures. As we stated in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56871), we received requests and recommendations for over 800 procedure codes that we were not able to fully evaluate and finalize for FY 2017. We discuss these requests and recommendations below.

We also are addressing separate requests that we received regarding changing the designation of specific ICD–10–PCS procedure codes. For each group summarized below, the detailed lists of procedure codes are shown in Tables 6P.4a through 6P.4p. (Proposed ICD–10–CM and ICD–10–PCS Code Designations, MCE and MS–DRG Changes—FY 2018) associated with this proposed rule (which are available via the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(2) Percutaneous Insertion of Intraluminal or Monitoring Device

One commenter identified 28 ICD–10–PCS procedure codes describing procedures involving the percutaneous insertion of intraluminal and monitoring devices into central nervous system and other cardiovascular body parts that generally would not require the resources of an operating room and can be performed at the bedside. We agree with the commenter. Therefore, we are proposing that the 135 ICD–10–PCS procedure codes listed in Table 6P.4a. be designated as non-O.R. procedures. We are inviting public comments on our proposal.

### Table: Proposed MS–DRG Titles

<table>
<thead>
<tr>
<th>MDC</th>
<th>MS–DRG</th>
<th>MS–DRG title</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>220</td>
<td>Cardiac Valve &amp; Other Major Cardiothoracic Procedure without Cardiac Catheterization with CC.</td>
</tr>
<tr>
<td>5</td>
<td>221</td>
<td>Cardiac Valve &amp; Other Major Cardiothoracic Procedure without Cardiac Catheterization without CC/ MCC.</td>
</tr>
<tr>
<td>5</td>
<td>222</td>
<td>Cardiac Defibrillator Implant with Cardiac Catheterization with AMI/Heart Failure/Shock with MCC.</td>
</tr>
<tr>
<td>5</td>
<td>223</td>
<td>Cardiac Defibrillator Implant with Cardiac Catheterization with AMI/Heart Failure/Shock without MCC.</td>
</tr>
<tr>
<td>5</td>
<td>224</td>
<td>Cardiac Defibrillator Implant with Cardiac Catheterization without AMI/Heart Failure/Shock with MCC.</td>
</tr>
<tr>
<td>5</td>
<td>225</td>
<td>Cardiac Defibrillator Implant with Cardiac Catheterization without AMI/Heart Failure/Shock without MCC.</td>
</tr>
<tr>
<td>5</td>
<td>226</td>
<td>Cardiac Defibrillator Implant without Cardiac Catheterization with MCC.</td>
</tr>
<tr>
<td>5</td>
<td>227</td>
<td>Cardiac Defibrillator Implant without Cardiac Catheterization without MCC.</td>
</tr>
<tr>
<td>5</td>
<td>242</td>
<td>Permanent Cardiac Pacemaker Implant with MCC.</td>
</tr>
<tr>
<td>5</td>
<td>243</td>
<td>Permanent Cardiac Pacemaker Implant with CC.</td>
</tr>
<tr>
<td>5</td>
<td>244</td>
<td>Permanent Cardiac Pacemaker Implant without CC/MCC.</td>
</tr>
<tr>
<td>5</td>
<td>245</td>
<td>AICD Generator Procedures.</td>
</tr>
<tr>
<td>5</td>
<td>258</td>
<td>Cardiac Pacemaker Device Replacement with MCC.</td>
</tr>
<tr>
<td>5</td>
<td>259</td>
<td>Cardiac Pacemaker Device Replacement without MCC.</td>
</tr>
<tr>
<td>5</td>
<td>260</td>
<td>Cardiac Pacemaker Revision Except Device Replacement with MCC.</td>
</tr>
<tr>
<td>5</td>
<td>261</td>
<td>Cardiac Pacemaker Revision Except Device Replacement with CC.</td>
</tr>
<tr>
<td>5</td>
<td>262</td>
<td>Cardiac Pacemaker Revision Except Device Replacement without CC/MCC.</td>
</tr>
<tr>
<td>5</td>
<td>265</td>
<td>AICD Lead Procedures.</td>
</tr>
<tr>
<td>5</td>
<td>266</td>
<td>Endovascular Cardiac Valve Replacement with MCC.</td>
</tr>
<tr>
<td>5</td>
<td>267</td>
<td>Endovascular Cardiac Valve Replacement without MCC.</td>
</tr>
<tr>
<td>5</td>
<td>268</td>
<td>Aortic and Heart Assist Procedures Except Pulsion Balloon with MCC.</td>
</tr>
<tr>
<td>5</td>
<td>269</td>
<td>Aortic and Heart Assist Procedures Except Pulsion Balloon without MCC.</td>
</tr>
<tr>
<td>5</td>
<td>270</td>
<td>Other Major Cardiovascular Procedures with MCC.</td>
</tr>
<tr>
<td>5</td>
<td>271</td>
<td>Other Major Cardiovascular Procedures with CC.</td>
</tr>
<tr>
<td>5</td>
<td>272</td>
<td>Other Major Cardiovascular Procedures without CC/MCC.</td>
</tr>
<tr>
<td>8</td>
<td>461</td>
<td>Bilateral or Multiple Major Joint Procedures Of Lower Extremity with MCC.</td>
</tr>
<tr>
<td>8</td>
<td>462</td>
<td>Bilateral or Multiple Major Joint Procedures Of Lower Extremity without MCC.</td>
</tr>
<tr>
<td>8</td>
<td>466</td>
<td>Revision of Hip or Knee Replacement with MCC.</td>
</tr>
<tr>
<td>8</td>
<td>467</td>
<td>Revision of Hip or Knee Replacement with CC.</td>
</tr>
<tr>
<td>8</td>
<td>468</td>
<td>Revision of Hip or Knee Replacement without CC/MCC.</td>
</tr>
<tr>
<td>8</td>
<td>469</td>
<td>Major Joint Replacement or Reattachment of Lower Extremity with MCC.</td>
</tr>
<tr>
<td>8</td>
<td>470</td>
<td>Major Joint Replacement or Reattachment of Lower Extremity without MCC.</td>
</tr>
</tbody>
</table>

We refer readers to those lists of procedure codes. For each group summarized below, the detailed lists of procedure codes are shown in Tables 6P.4a through 6P.4p.
PCS procedure code lists in Table 6P.4b. associated with this proposed rule (which is available via the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(7) External/Diagnostic Drainage

One commenter identified 28 ICD–10–PCS procedure codes that describe procedures involving the percutaneous revision of drainage, infusion, intraluminal and monitoring devices from central nervous system and other vascular body parts that generally would not require the resources of an operating room and can be performed at the bedside. We agree with the commenter. Therefore, we are proposing that the 28 ICD–10–PCS procedure codes listed in Table 6P.4c. associated with this proposed rule (which is available via the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) be designated as non-O.R. procedures. We are inviting public comments on our proposal.

ICD–10–PCS code | Code description
--- | ---
02PAXMZ | Removal of cardiac lead from heart, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
01PYXMZ | Removal of neurostimulator lead from peripheral nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00PEXMZ | Removal of neurostimulator lead from cranial nerve, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
00P6XMZ | Removal of neurostimulator lead from cerebral ventricle, external approach.
We agree with the commenter. Therefore, we are proposing that the four ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(9) External Removal of Radioactive Element or Synthetic Substitute

One commenter identified three ICD–10–PCS procedure codes that describe procedures involving the external removal of radioactive or synthetic substitutes from the eye that generally would not require the resources of an operating room and can be performed at the bedside. These three ICD–10–PCS codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>08P0X1Z</td>
<td>Removal of radioactive element from right eye, external approach.</td>
</tr>
<tr>
<td>08P0XJZ</td>
<td>Removal of synthetic substitute from right eye, external approach.</td>
</tr>
<tr>
<td>08P1XJZ</td>
<td>Removal of synthetic substitute from left eye, external approach.</td>
</tr>
</tbody>
</table>

We agree with the commenter. Therefore, we are proposing that the three ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(10) Endoscopic/Transorifice Diagnostic Drainage

One commenter identified eight ICD–10–PCS procedure codes that describe procedures involving endoscopic/transorifice (via natural or artificial opening) drainage of ear structures that generally would not require the resources of an operating room and can be performed at the bedside. These eight ICD–10–PCS codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>09977ZX</td>
<td>Drainage of right tympanic membrane, via natural or artificial opening, diagnostic.</td>
</tr>
<tr>
<td>09976ZX</td>
<td>Drainage of right tympanic membrane, via natural or artificial opening endoscopic, diagnostic.</td>
</tr>
<tr>
<td>09987ZX</td>
<td>Drainage of left tympanic membrane, via natural or artificial opening, diagnostic.</td>
</tr>
<tr>
<td>09988ZX</td>
<td>Drainage of left tympanic membrane, via natural or artificial opening endoscopic, diagnostic.</td>
</tr>
<tr>
<td>099F7ZX</td>
<td>Drainage of right eustachian tube, via natural or artificial opening, diagnostic.</td>
</tr>
<tr>
<td>099F8ZX</td>
<td>Drainage of right eustachian tube, via natural or artificial opening endoscopic, diagnostic.</td>
</tr>
<tr>
<td>099G7ZX</td>
<td>Drainage of left eustachian tube, via natural or artificial opening, diagnostic.</td>
</tr>
<tr>
<td>099G8ZX</td>
<td>Drainage of left eustachian tube, via natural or artificial opening endoscopic, diagnostic.</td>
</tr>
</tbody>
</table>

We agree with the commenter. Therefore, we are proposing that the eight ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(11) External Release

One commenter identified four ICD–10–PCS procedure codes that describe procedures involving the external release of ear structures that generally would not require the resources of an operating room and can be performed at the bedside. These four ICD–10–PCS codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>09N0XZZ</td>
<td>Release right external ear, external approach.</td>
</tr>
<tr>
<td>09N1XZZ</td>
<td>Release left external ear, external approach.</td>
</tr>
<tr>
<td>09N3XZZ</td>
<td>Release right external auditory canal, external approach.</td>
</tr>
<tr>
<td>09N4XZZ</td>
<td>Release left external auditory canal, external approach.</td>
</tr>
</tbody>
</table>

We agree with the commenter. Therefore, we are proposing that the four ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(12) External Repair

One commenter identified three ICD–10–PCS procedure codes that describe procedures involving the external repair of body parts that generally would not require the resources of an operating room and can be performed at the bedside. These three ICD–10–PCS codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>09QKXZZ</td>
<td>Repair nose, external approach.</td>
</tr>
<tr>
<td>0CQ4XZZ</td>
<td>Repair buccal mucosa, external approach.</td>
</tr>
<tr>
<td>0CQ7XZZ</td>
<td>Repair tongue, external approach.</td>
</tr>
</tbody>
</table>

We agree with the commenter. Therefore, we are proposing that the three ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(13) Endoscopic/Transorifice Destruction

One commenter identified eight ICD–10–PCS procedure codes that describe procedures involving endoscopic/transorifice destruction of respiratory system body parts that generally would not require the resources of an operating room and can be performed at the bedside. These eight ICD–10–PCS codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>09QKXZZ</td>
<td>Repair nose, external approach.</td>
</tr>
<tr>
<td>0CQ4XZZ</td>
<td>Repair buccal mucosa, external approach.</td>
</tr>
<tr>
<td>0CQ7XZZ</td>
<td>Repair tongue, external approach.</td>
</tr>
</tbody>
</table>
We agree with the commenter. Therefore, we are proposing that the eight ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(14) Endoscopic/Transorifice Drainage

One commenter identified 40 ICD–10–PCS procedure codes that describe procedures involving endoscopic/ transorifice (via natural or artificial opening) drainage of respiratory system body parts that generally would not require the resources of an operating room and can be performed at the bedside. We agree with the commenter. Therefore, we are proposing that the 40 ICD–10–PCS procedure codes listed in Table 6P.4f. associated with this proposed rule (which is available via the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) be designated as non-O.R. procedures. We are inviting public comments on our proposal.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0B538ZZ</td>
<td>Destruction of right main bronchus, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0B548ZZ</td>
<td>Destruction of right upper lobe bronchus, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0B558ZZ</td>
<td>Destruction of right middle lobe bronchus, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0B568ZZ</td>
<td>Destruction of right lower lobe bronchus, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0B578ZZ</td>
<td>Destruction of left main bronchus, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0B588ZZ</td>
<td>Destruction of left upper lobe bronchus, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0B598ZZ</td>
<td>Destruction of lingula bronchus, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0B5B8ZZ</td>
<td>Destruction of left lower lobe bronchus, via natural or artificial opening endoscopic.</td>
</tr>
</tbody>
</table>

We agree with the commenter. Therefore, we are proposing that the nine ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(15) Endoscopic/Transorifice Extirpation

One commenter identified nine ICD–10–PCS procedure codes that describe procedures involving endoscopic/ transorifice extirpation of matter from respiratory system body parts that generally would not require the resources of an operating room and can be performed at the bedside. These nine ICD–10–PCS codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0BCC8ZZ</td>
<td>Extirpation of matter from right upper lung lobe, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0BCD8ZZ</td>
<td>Extirpation of matter from right middle lung lobe, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0BCF8ZZ</td>
<td>Extirpation of matter from right lower lung lobe, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0BCG8ZZ</td>
<td>Extirpation of matter from left upper lung lobe, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0BCH8ZZ</td>
<td>Extirpation of matter from lung lingula, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0BCJ8ZZ</td>
<td>Extirpation of matter from left lower lung lobe, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0BCK8ZZ</td>
<td>Extirpation of matter from right lung, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0BCL8ZZ</td>
<td>Extirpation of matter from left lung, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0BCM8ZZ</td>
<td>Extirpation of matter from bilateral lungs, via natural or artificial opening endoscopic.</td>
</tr>
</tbody>
</table>

We agree with the commenter. Therefore, we are proposing that the nine ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(16) Endoscopic/Transorifice Fragmentation

One commenter identified 16 ICD–10–PCS procedure codes that describe procedures involving endoscopic/ transorifice fragmentation of respiratory system body parts that generally would not require the resources of an operating room and can be performed at the bedside. These 16 ICD–10–PCS codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0BF37ZZ</td>
<td>Fragmentation in right main bronchus, via natural or artificial opening.</td>
</tr>
<tr>
<td>0BF38ZZ</td>
<td>Fragmentation in right main bronchus, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0BF47ZZ</td>
<td>Fragmentation in right upper lobe bronchus, via natural or artificial opening.</td>
</tr>
<tr>
<td>0BF48ZZ</td>
<td>Fragmentation in right upper lobe bronchus, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0BF57ZZ</td>
<td>Fragmentation in right middle lobe bronchus, via natural or artificial opening.</td>
</tr>
<tr>
<td>0BF58ZZ</td>
<td>Fragmentation in right middle lobe bronchus, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0BF67ZZ</td>
<td>Fragmentation in right lower lobe bronchus, via natural or artificial opening.</td>
</tr>
<tr>
<td>0BF68ZZ</td>
<td>Fragmentation in right lower lobe bronchus, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0BF77ZZ</td>
<td>Fragmentation in left main bronchus, via natural or artificial opening.</td>
</tr>
<tr>
<td>0BF78ZZ</td>
<td>Fragmentation in left main bronchus, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0BF87ZZ</td>
<td>Fragmentation in left upper lobe bronchus, via natural or artificial opening.</td>
</tr>
<tr>
<td>0BF88ZZ</td>
<td>Fragmentation in left upper lobe bronchus, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0BF97ZZ</td>
<td>Fragmentation in lingula bronchus, via natural or artificial opening.</td>
</tr>
<tr>
<td>0BF98ZZ</td>
<td>Fragmentation in lingula bronchus, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0BF99ZZ</td>
<td>Fragmentation in left lower lobe bronchus, via natural or artificial opening.</td>
</tr>
<tr>
<td>0BFB8ZZ</td>
<td>Fragmentation in left lower lobe bronchus, via natural or artificial opening endoscopic.</td>
</tr>
</tbody>
</table>
We agree with the commenter. Therefore, we are proposing that the 16 ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(17) Endoscopic/Transorifice Insertion of Intraluminal Device

One commenter identified two ICD–10–PCS procedure codes that describe procedures involving an endoscopic/transorifice (via natural or artificial opening) insertion of intraluminal devices into respiratory system body parts that generally would not require the resources of an operating room and can be performed at the bedside. These two ICD–10–PCS codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0BH17DZ</td>
<td>Insertion of intraluminal device into trachea, via natural or artificial opening.</td>
</tr>
<tr>
<td>0BH18DZ</td>
<td>Insertion of intraluminal device into trachea, via natural or artificial opening endoscopic.</td>
</tr>
</tbody>
</table>

We agree with the commenter. Therefore, we are proposing that the two ICD–10–PCS procedure codes shown in the table above be designated non-O.R. procedures. We are inviting public comments on our proposal.

(18) Endoscopic/Transorifice Removal of Radioactive Element

One commenter identified two ICD–10–PCS procedure codes that describe procedures involving the endoscopic/transorifice removal of radioactive elements from respiratory system body parts that generally would not require the resources of an operating room and can be performed at the bedside. These two ICD–10–PCS codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0BPK71Z</td>
<td>Removal of radioactive element from right lung, via natural or artificial opening.</td>
</tr>
<tr>
<td>0BPK81Z</td>
<td>Removal of radioactive element from right lung, via natural or artificial opening endoscopic.</td>
</tr>
</tbody>
</table>

We agree with the commenter. Therefore, we are proposing that the two ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(19) Endoscopic/Transorifice Revision of Drainage, Infusion, Intraluminal or Monitoring Device

One commenter identified 18 ICD–10–PCS procedure codes that describe procedures involving the revision of drainage, infusion, intraluminal, or monitoring devices from respiratory system body parts that generally would not require the resources of an operating room and can be performed at the bedside. We agree with the commenter. Therefore, we are proposing that the 18 ICD–10–PCS procedure codes listed in Table 6P.4g., associated with this proposed rule (which is available via the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(20) Endoscopic/Transorifice Excision

One commenter identified one ICD–10–PCS procedure code that describes the procedure involving endoscopic/transorifice (via natural or artificial opening) excision of the digestive system body parts that generally would not require the resources of an operating room and can be performed at the bedside. This code is 0DBQ8ZZ (Excision of anus, via natural or artificial opening endoscopic. We agree with the commenter. Therefore, we are proposing that ICD–10–PCS procedure code 0DBQ8ZZ be designated as a non-O.R. procedure. We are inviting public comments on our proposal.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0DH67DZ</td>
<td>Insertion of intraluminal device into stomach, via natural or artificial opening.</td>
</tr>
<tr>
<td>0DH88DZ</td>
<td>Insertion of intraluminal device into stomach, via natural or artificial opening endoscopic.</td>
</tr>
</tbody>
</table>

We agree with the commenter. Therefore, we are proposing that the two ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(22) Endoscopic/Transorifice Removal of Feeding Device

One commenter identified six ICD–10–PCS procedure codes that describe procedures involving endoscopic/transorifice (via natural or artificial opening) removal of feeding devices that generally would not require the resources of an operating room and can be performed at the bedside. These six ICD–10–PCS codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0DP07UZ</td>
<td>Removal of feeding device from upper intestinal tract, via natural or artificial opening.</td>
</tr>
<tr>
<td>0DP08UZ</td>
<td>Removal of feeding device from upper intestinal tract, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0DP67UZ</td>
<td>Removal of feeding device from stomach, via natural or artificial opening.</td>
</tr>
<tr>
<td>0DP68UZ</td>
<td>Removal of feeding device from stomach, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0DP77UZ</td>
<td>Removal of feeding device from lower intestinal tract, via natural or artificial opening.</td>
</tr>
</tbody>
</table>
We agree with the commenter. Therefore, we are proposing that the six ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0DPD8UZ</td>
<td>Removal of feeding device from lower intestinal tract, via natural or artificial opening endoscopic.</td>
</tr>
</tbody>
</table>

(23) External Reposition

One commenter identified two ICD–10–PCS procedure codes that describe procedures involving external reposition of gastrointestinal body parts that generally would not require the resources of an operating room and can be performed at the bedside. These two ICD–10–PCS codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0DS5XZZ</td>
<td>Reposition esophagus, external approach.</td>
</tr>
<tr>
<td>0DSQXZZ</td>
<td>Reposition anus, external approach.</td>
</tr>
</tbody>
</table>

We agree with the commenter. Therefore, we are proposing that the two ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0EPD8UZ</td>
<td>Draining of right hepatic duct with drainage device, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0EPQXZZ</td>
<td>Draining of right hepatic duct, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0EP96UZ</td>
<td>Draining of left hepatic duct with drainage device, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0EP98UZ</td>
<td>Draining of cystic duct with drainage device, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0EP99UZ</td>
<td>Draining of cystic duct, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0EPD8ZZ</td>
<td>Draining of pancreatic duct, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0EPF8ZZ</td>
<td>Draining of accessory pancreatic duct, via natural or artificial opening endoscopic.</td>
</tr>
</tbody>
</table>

We agree with the commenter. Therefore, we are proposing that the eight ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0EPF8ZZ</td>
<td>Fragmentation in accessory pancreatic duct, via natural or artificial opening endoscopic.</td>
</tr>
</tbody>
</table>

(25) Endoscopic/Transorifice Fragmentation

One commenter identified two ICD–10–PCS procedure codes that describe procedures involving endoscopic/transorifice (via natural or artificial opening) fragmentation of hepatobiliary system and pancreatic body parts that generally would not require the resources of an operating room and can be performed at the bedside. These two ICD–10–PCS codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0FFD8ZZ</td>
<td>Fragmentation in pancreatic duct, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0FFF8ZZ</td>
<td>Fragmentation in accessory pancreatic duct, via natural or artificial opening endoscopic.</td>
</tr>
</tbody>
</table>

We agree with the commenter. Therefore, we are proposing that the two ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0H0T3JZ</td>
<td>Alteration of right breast with synthetic substitute, percutaneous approach.</td>
</tr>
<tr>
<td>0H0U3JZ</td>
<td>Alteration of left breast with synthetic substitute, percutaneous approach.</td>
</tr>
<tr>
<td>0H0V3JZ</td>
<td>Alteration of bilateral breast with synthetic substitute, percutaneous approach.</td>
</tr>
</tbody>
</table>

We agree with the commenter. Therefore, we are proposing that the three ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0H0T3JZ</td>
<td>Alteration of right breast with synthetic substitute, percutaneous approach.</td>
</tr>
<tr>
<td>0H0U3JZ</td>
<td>Alteration of left breast with synthetic substitute, percutaneous approach.</td>
</tr>
<tr>
<td>0H0V3JZ</td>
<td>Alteration of bilateral breast with synthetic substitute, percutaneous approach.</td>
</tr>
</tbody>
</table>
We agree with the commenter. Therefore, we are proposing that the three ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(27) External Division and Excision of Skin

One commenter identified 41 ICD–10–PCS procedure codes that describe procedures involving external division and excision of the skin for body parts that generally would not require the resources of an operating room and can be performed at the bedside. We agree with the commenter. Therefore, we are proposing that the 41 ICD–10–PCS procedure codes listed in Table 6P.4h. associated with this proposed rule (which is available via the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(28) Percutaneous Supplement

One commenter identified three ICD–10–PCS procedure codes that describe procedures involving percutaneous supplement of the breast with synthetic substitute that generally would not require the resources of an operating room and can be performed at the bedside. These three ICD–10–PCS codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0HUT3JZ ..........</td>
<td>Supplement right breast with synthetic substitute, percutaneous approach.</td>
</tr>
<tr>
<td>0HUU3JZ ..........</td>
<td>Supplement left breast with synthetic substitute, percutaneous approach.</td>
</tr>
<tr>
<td>0HUV3JZ ..........</td>
<td>Supplement bilateral breast with synthetic substitute, percutaneous approach.</td>
</tr>
</tbody>
</table>

We disagree with the commenter because these procedure codes describe various types of surgery performed on the breast or nipple (for example, partial mastectomy) that would typically involve the use of general anesthesia. Therefore, we are proposing that the six ICD–10–PCS procedure codes shown in the table above remain designated as O.R. procedures. We are inviting public comments on our proposal.

(29) Percutaneous Drainage

One commenter identified 25 ICD–10–PCS procedure codes that describe procedures involving percutaneous drainage of subcutaneous tissue and fascia body parts that generally would not require the resources of an operating room and can be performed at the bedside. The list includes procedure codes for drainage with or without placement of a drainage device. We agree with the commenter. Therefore, we are proposing that the 25 ICD–10–PCS procedure codes listed in Table 6P.4i. associated with this proposed rule (which is available via the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(30) Open Drainage

One commenter identified 25 ICD–10–PCS procedure codes that describe procedures involving open drainage of subcutaneous tissue and fascia body parts that generally would not require the resources of an operating room and can be performed at the bedside. We agree with the commenter. Therefore, we are proposing that the 25 ICD–10–PCS procedure codes listed in Table 6P.4i. associated with this proposed rule (which is available via the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(31) Percutaneous Extraction

One commenter identified 22 ICD–10–PCS procedure codes that describe procedures involving percutaneous extraction of subcutaneous tissue and fascia body parts that generally would not require the resources of an operating room and can be performed at the bedside. We agree with the commenter. Therefore, we are proposing that the 22 ICD–10–PCS procedure codes listed in Table 6P.4j. associated with this proposed rule (which is available via the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) be...
designated as non-O.R. procedures. We are inviting public comments on our proposal.

(33) Open Extraction

One commenter identified 22 ICD–10–PCS procedure codes that describe procedures involving open extraction of subcutaneous tissue and fascia body parts that the commenter believed would generally not require the resources of an operating room and can be performed at the bedside. We disagree with the commenter because these codes describe procedures that utilize an open approach and are being performed on the skin and subcutaneous tissue. Depending on the medical reason for the open extraction, the procedures may require an O.R. setting. Therefore, we are proposing that the 22 ICD–10–PCS procedure codes listed in Table 6P.4k associated with this proposed rule (which is available via the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) remain designated as O.R. procedures. We are inviting public comments on our proposal.

(34) Percutaneous and Open Repair

One commenter identified 44 ICD–10–PCS procedure codes that describe procedures involving percutaneous and open repair of subcutaneous tissue and fascia body parts that generally would not require the resources of an operating room and can be performed at the bedside. We agree with the commenter. Therefore, we are proposing that the 44 ICD–10–PCS procedure codes listed in Table 6P.4l associated with this proposed rule (which is available via the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(35) External Release

One commenter identified 28 ICD–10–PCS procedure codes that describe procedures involving external release of bursa and ligament body parts that generally would not require the resources of an operating room and can be performed at the bedside. We agree with the commenter. Therefore, we are proposing that the 28 ICD–10–PCS procedure codes listed in Table 6P.4m associated with this proposed rule (which is available via the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(36) External Repair

One commenter identified 135 ICD–10–PCS procedure codes that describe procedures involving external repair of various bones and joints. We believe that these procedures generally would not be performed in the operating room. We are proposing that the 135 ICD–10–PCS procedure codes listed in Table 6P.4n associated with this proposed rule (which is available via the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html) be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(37) External Reposition

One commenter identified 14 ICD–10–PCS procedure codes that describe procedures involving external reposition of various bones. These 14 ICD–10–PCS codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ONS0XZZ</td>
<td>Reposition skull, external approach.</td>
</tr>
<tr>
<td>ONS1XZZ</td>
<td>Reposition right frontal bone, external approach.</td>
</tr>
<tr>
<td>ONS2XZZ</td>
<td>Reposition left frontal bone, external approach.</td>
</tr>
<tr>
<td>ONS3XZZ</td>
<td>Reposition right parietal bone, external approach.</td>
</tr>
<tr>
<td>ONS4XZZ</td>
<td>Reposition left parietal bone, external approach.</td>
</tr>
<tr>
<td>ONS5XZZ</td>
<td>Reposition right temporal bone, external approach.</td>
</tr>
<tr>
<td>ONS6XZZ</td>
<td>Reposition left temporal bone, external approach.</td>
</tr>
<tr>
<td>ONS7XZZ</td>
<td>Reposition right occipital bone, external approach.</td>
</tr>
<tr>
<td>ONS8XZZ</td>
<td>Reposition left occipital bone, external approach.</td>
</tr>
<tr>
<td>OPS3XZZ</td>
<td>Reposition cervical vertebra, external approach.</td>
</tr>
<tr>
<td>OPS4XZZ</td>
<td>Reposition thoracic vertebra, external approach.</td>
</tr>
<tr>
<td>OQS0XZZ</td>
<td>Reposition lumbar vertebra, external approach.</td>
</tr>
<tr>
<td>OQS1XZZ</td>
<td>Reposition sacrum, external approach.</td>
</tr>
<tr>
<td>OQS2XZZ</td>
<td>Reposition coccyx, external approach.</td>
</tr>
</tbody>
</table>

We believe that these procedures generally would not be performed in the operating room. Therefore, we are proposing that the 14 ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

(38) Endoscopic/Transorifice Dilation

One commenter identified eight ICD–10–PCS procedure codes that describe procedures involving endoscopic/transorifice (via natural or artificial opening) dilation of urinary system body parts that generally would not require the resources of an operating room and can be performed at the bedside. These eight ICD–10–PCS codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0T767ZZ</td>
<td>Dilation of right ureter, via natural or artificial opening.</td>
</tr>
<tr>
<td>0T768ZZ</td>
<td>Dilation of right ureter, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0T777ZZ</td>
<td>Dilation of left ureter, via natural or artificial opening.</td>
</tr>
<tr>
<td>0T778ZZ</td>
<td>Dilation of left ureter, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0T779ZZ</td>
<td>Dilation of bladder with intraluminal device, via natural or artificial opening.</td>
</tr>
<tr>
<td>0T787ZZ</td>
<td>Dilation of bladder with intraluminal device, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0T776ZZ</td>
<td>Dilation of bladder, via natural or artificial opening.</td>
</tr>
<tr>
<td>0T77B0DZ</td>
<td>Dilation of bladder, via natural or artificial opening endoscopic.</td>
</tr>
</tbody>
</table>
We agree with the commenter. Therefore, we are proposing that the eight ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>4A0C35Z</td>
<td>Measurement of biliary flow, percutaneous approach.</td>
</tr>
<tr>
<td>4A0635Z</td>
<td>Measurement of lymphatic flow, percutaneous approach.</td>
</tr>
<tr>
<td>3E1N88Z</td>
<td>Irrigation of male reproductive using irrigating substance, via natural or artificial opening.</td>
</tr>
<tr>
<td>3E1N88X</td>
<td>Irrigation of male reproductive using irrigating substance, via natural or artificial opening, diagnostic.</td>
</tr>
<tr>
<td>4A069BZ</td>
<td>Measurement of lymphatic pressure, percutaneous approach.</td>
</tr>
<tr>
<td>4AOC35Z</td>
<td>Measurement of biliary flow, percutaneous approach.</td>
</tr>
</tbody>
</table>

One commenter identified 20 ICD–10–PCS procedure codes that describe procedures involving percutaneous/transorifice (via natural or artificial opening) excision of urinary system body parts that the commenter believed would generally not require the resources of an operating room and can be performed at the bedside. These three ICD–10–PCS codes are shown in the table below.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>4A0C35Z</td>
<td>Measurement of biliary flow, percutaneous approach.</td>
</tr>
<tr>
<td>4A0635Z</td>
<td>Measurement of lymphatic flow, percutaneous approach.</td>
</tr>
<tr>
<td>3E1N88Z</td>
<td>Irrigation of male reproductive using irrigating substance, via natural or artificial opening.</td>
</tr>
<tr>
<td>3E1N88X</td>
<td>Irrigation of male reproductive using irrigating substance, via natural or artificial opening, diagnostic.</td>
</tr>
<tr>
<td>4A069BZ</td>
<td>Measurement of lymphatic pressure, percutaneous approach.</td>
</tr>
<tr>
<td>4AOC35Z</td>
<td>Measurement of biliary flow, percutaneous approach.</td>
</tr>
</tbody>
</table>

We disagree with the commenter because, depending on the medical reason for the excision, the procedures may require an O.R. setting. Therefore, we are proposing that the three ICD–10–PCS procedure codes shown in the table above remain designated as O.R. procedures. We are inviting public comments on our proposal.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0TBD7ZZ</td>
<td>Excision of urethra, via natural or artificial opening.</td>
</tr>
<tr>
<td>0TBD8ZZ</td>
<td>Excision of urethra, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0TBDXZZ</td>
<td>Excision of urethra, external approach.</td>
</tr>
</tbody>
</table>

We agree with the commenter. Therefore, we are proposing that these three ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0UQG7ZZ</td>
<td>Repair vagina, via natural or artificial opening.</td>
</tr>
<tr>
<td>0UQGXZZ</td>
<td>Repair vagina, external approach.</td>
</tr>
<tr>
<td>0UQMXZZ</td>
<td>Repair vulva, external approach.</td>
</tr>
</tbody>
</table>

We disagree with the commenter because, depending on the medical reason for the excision, the procedures may require an O.R. setting. Therefore, we are proposing that the three ICD–10–PCS procedure codes shown in the table above remain designated as O.R. procedures. We are inviting public comments on our proposal.

<table>
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We agree with the commenter. Therefore, we are proposing that these three ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

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</tr>
<tr>
<td>0TBD8ZZ</td>
<td>Excision of urethra, via natural or artificial opening endoscopic.</td>
</tr>
<tr>
<td>0TBDXZZ</td>
<td>Excision of urethra, external approach.</td>
</tr>
</tbody>
</table>

One commenter identified 51 ICD–10–PCS procedure codes that describe procedures involving external/transorifice repair of the vagina body part that generally would not require the resources of an operating room and can be performed at the bedside. These three ICD–10–PCS codes are shown in the table below.

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We agree with the commenter. Therefore, we are proposing that these three ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

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We disagree with the commenter because, depending on the medical reason for the excision, the procedures may require an O.R. setting. Therefore, we are proposing that the three ICD–10–PCS procedure codes shown in the table above remain designated as O.R. procedures. We are inviting public comments on our proposal.

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<tr>
<td>0TBDXZZ</td>
<td>Excision of urethra, external approach.</td>
</tr>
</tbody>
</table>

One commenter identified 15 ICD–10–PCS procedure codes that describe procedures involving percutaneous/diagnostic and endoscopic/transorifice (via natural or artificial opening) irrigation, measurement and monitoring of structures, pressures and flow that generally would not require the resources of an operating room and can be performed at the bedside. These 15 ICD–10–PCS codes are shown in the table below.

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<th>Code description</th>
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</thead>
<tbody>
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<td>Repair vagina, via natural or artificial opening.</td>
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<td>Repair vagina, external approach.</td>
</tr>
<tr>
<td>0UQMXZZ</td>
<td>Repair vulva, external approach.</td>
</tr>
</tbody>
</table>
We agree with the commenter. Therefore, we are proposing that the 15 ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

We agree with the commenter. Therefore, we are proposing that the six ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

We agree with the commenter. Therefore, we are proposing that the five ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

We received a request to review three ICD–10–PCS procedure codes that describe procedures for revision of a neurostimulator generator that are currently designated as O.R. procedures and assigned to MS–DRGs 252, 253 and 254 (Other Vascular Procedures with contrast of hepatobiliary system body parts that generally would not require the resources of an operating room and can be performed at the bedside. These six ICD–10–PCS codes are shown in the table below.

We agree with the commenter. Therefore, we are proposing that the five ICD–10–PCS procedure codes shown in the table above be designated as non-O.R. procedures. We are inviting public comments on our proposal.

The requester expressed concern with the MS–DRG assignments and noted that although these codes are used to report revision of a carotid sinus stimulator pulse generator and appropriately assigned to MS–DRGs 252, 253 and 254 in MDC 5 (Diseases and Disorders of the Nervous System), when either code is reported in combination with one of the three procedure codes that describe revision of neurostimulator generator codes (currently assigned to MDC 5), the resulting MS–DRG assignment is to MS–DRGs 981, 982 and 983 (Extensive O.R. Procedure Unrelated to Principal Diagnosis with MCC, with CC and without CC/MCC, respectively). The three codes are 4A0C3BZ (Revision of stimulator generator in trunk subcutaneous tissue and fascia, percutaneous approach), 4A0C7BZ (Revision of stimulator generator in trunk subcutaneous tissue and fascia, percutaneous approach), and 4A0C85Z (Revision of stimulator generator, initial encounter) or T85.890A (Other specified complication of nervous system prosthetic devices, implants and grafts, initial encounter). Because both of these diagnosis codes are assigned to MDC 1 (Diseases and Disorders of the Nervous System), when either code is reported in combination with one of the three procedure codes that describe revision of neurostimulator generator codes (currently assigned to MDC 5), the resulting MS–DRG assignment is to MS–DRGs 981, 982 and 983 (Extensive O.R. Procedure Unrelated to Principal Diagnosis with MCC, with CC and without CC/MCC, respectively).

The requester also stated that the indication for revision of a neurostimulator generator is typically due to a complication, which would be reflected in a complication code such as ICD–10–CM diagnosis code T85.734A (Infection and inflammatory reaction due to implanted electronic neurostimulator, generator, initial encounter) or T85.890A (Other specified complication of nervous system prosthetic devices, implants and grafts, initial encounter).
sinus stimulators and the other types of neurostimulator cases would appropriately group to medical MS–
DRGs.
• Add the ICD–10–PCS procedure codes to MDC 1, such as to MS–DRGs 040, 041 and 042 (Peripheral, Cranial Nerve and Other Nervous System Procedures with MCC, with CC, or Peripheral Neurostimulator and without CC/MCC, respectively) under MDC 1.

The requestor stated that this option would resolve the inconsistency between a revision of a carotid sinus stimulator generator being classified as an O.R. procedure, while the other comparable procedures involving a revision of a regular neurostimulator generator are not. The requestor also stated that this option would preclude cases being assigned to MS–DRGs 981 through 983.
• Stop classifying the ICD–10–PCS procedure codes as O.R. procedures entirely. The requestor stated that, under this option, all cases would then group to medical MS–DRGs, regardless of the type of neurostimulator generator. We analyzed claims data for the three revision of neurostimulator generator procedure codes from the December 2016 update of the FY 2016 MedPAR file and identified cases under MDC 1 (Diseases and Disorders of the Nervous System) in MS–DRGs 025, 026, and 027 (Granulotomy and Endovascular Intracranial Procedures with MCC, with CC and without CC/MCC, respectively); MS–DRGs 029 and 030 (Spinal Procedures with CC or Neurostimulators and Spinal Procedures without CC/ MCC, respectively); and MS–DRGs 041 and 042 (Peripheral, Cranial Nerve and Other Nervous System Procedures with CC or Peripheral Neurostimulator and without CC/MCC, respectively). We also identified cases in MS–DRGs 982 and 983 (Extensive O.R. Procedure Unrelated to Principal Diagnosis with CC and without CC/MCC, respectively). Lastly, we identified cases under MDC 5 (Diseases and Disorders of the Circulatory System) in MS–DRGs 252, 253 and 254 (Other Vascular Procedures with MCC, with CC and without CC/MCC, respectively). Our findings are shown in the table below.

**MS–DRGs for Revision of Neurostimulator Generator**

<table>
<thead>
<tr>
<th>MS–DRG</th>
<th>Number of cases</th>
<th>Average length of stay</th>
<th>Average costs</th>
</tr>
</thead>
<tbody>
<tr>
<td>MS–DRG 025—All cases</td>
<td>18,442</td>
<td>9.1</td>
<td>$29,984</td>
</tr>
<tr>
<td>MS–DRG 025—Cases with revision of neurostimulator generator</td>
<td>1</td>
<td>12.0</td>
<td>73,716</td>
</tr>
<tr>
<td>MS–DRG 026—All cases</td>
<td>8,415</td>
<td>5.6</td>
<td>21,557</td>
</tr>
<tr>
<td>MS–DRG 026—Cases with revision of neurostimulator generator</td>
<td>1</td>
<td>6.0</td>
<td>4,537</td>
</tr>
<tr>
<td>MS–DRG 027—All cases</td>
<td>10,087</td>
<td>2.9</td>
<td>17,320</td>
</tr>
<tr>
<td>MS–DRG 027—Cases with revision of neurostimulator generator</td>
<td>4</td>
<td>1.8</td>
<td>13,906</td>
</tr>
<tr>
<td>MS–DRG 029—All cases</td>
<td>3,192</td>
<td>5.9</td>
<td>23,145</td>
</tr>
<tr>
<td>MS–DRG 029—Cases with revision of neurostimulator generator</td>
<td>6</td>
<td>3.5</td>
<td>32,799</td>
</tr>
<tr>
<td>MS–DRG 030—All cases</td>
<td>1,933</td>
<td>2.9</td>
<td>14,901</td>
</tr>
<tr>
<td>MS–DRG 030—Cases with revision of neurostimulator generator</td>
<td>11</td>
<td>2.2</td>
<td>18,294</td>
</tr>
<tr>
<td>MS–DRG 041—All cases</td>
<td>5,154</td>
<td>5.5</td>
<td>16,633</td>
</tr>
<tr>
<td>MS–DRG 041—Cases with revision of neurostimulator generator</td>
<td>1</td>
<td>1.0</td>
<td>14,145</td>
</tr>
<tr>
<td>MS–DRG 042—All cases</td>
<td>2,099</td>
<td>3.2</td>
<td>13,725</td>
</tr>
<tr>
<td>MS–DRG 042—Cases with revision of neurostimulator generator</td>
<td>2</td>
<td>2.0</td>
<td>28,587</td>
</tr>
<tr>
<td>MS–DRG 982—All cases</td>
<td>15,216</td>
<td>6.6</td>
<td>17,341</td>
</tr>
<tr>
<td>MS–DRG 982—Cases with revision of neurostimulator generator</td>
<td>11</td>
<td>3.0</td>
<td>15,336</td>
</tr>
<tr>
<td>MS–DRG 983—All cases</td>
<td>3,508</td>
<td>3.2</td>
<td>11,627</td>
</tr>
<tr>
<td>MS–DRG 983—Cases with revision of neurostimulator generator</td>
<td>9</td>
<td>4.2</td>
<td>19,951</td>
</tr>
<tr>
<td>MS–DRG 252—All cases</td>
<td>33,817</td>
<td>7.6</td>
<td>23,384</td>
</tr>
<tr>
<td>MS–DRG 252—Cases with revision of neurostimulator generator</td>
<td>1</td>
<td>7.0</td>
<td>18,740</td>
</tr>
<tr>
<td>MS–DRG 253—All cases</td>
<td>27,456</td>
<td>5.5</td>
<td>18,519</td>
</tr>
<tr>
<td>MS–DRG 253—Cases with revision of neurostimulator generator</td>
<td>7</td>
<td>2.4</td>
<td>19,078</td>
</tr>
<tr>
<td>MS–DRG 254—All cases</td>
<td>10,089</td>
<td>2.9</td>
<td>14,901</td>
</tr>
<tr>
<td>MS–DRG 254—Cases with revision of neurostimulator generator</td>
<td>3</td>
<td>3.0</td>
<td>11,981</td>
</tr>
</tbody>
</table>

As shown in the table above, the overall volume of cases reporting revision of neurostimulator generator is low, with a total of only 57 cases found across all of the MS–DRGs reviewed. The average length of stay for these cases reporting revision of neurostimulator generators is, in most cases, consistent with the average length of stay for all cases in the respective MS–DRG, with the majority having an average length of stay below the average length of stay for all cases in the respective MS–DRG. Finally, the average costs for cases reporting revision of neurostimulator generator reflect a wide range, with a low of $4,537 in MS–DRG 026 to a high of $73,716 in MS–DRG 025. It is clear that, for MS–DRG 025 where the average costs of all cases were $29,984 and the average costs of the one case reporting revision of a neurostimulator generator was $73,716, this is an atypical case. It is also clear from the data that there were other procedures reported on the claims where a procedure code for a revision of a neurostimulator generator was assigned due to the various MS–DRG assignments.

After review of the claims data and discussion with our clinical advisors, we agree with and support the requestor’s first option—to reclassify the three ICD–10–PCS procedure codes for revision of neurostimulator generators from O.R. procedures to non-O.R. procedures that affect the assignment for MS–DRGs 252, 253 and 254 to account for the subset of patients undergoing revision of a carotid sinus neurostimulator generator specifically. In cases where one of the more common (for example, gastric, intracranial, sacral and spinal) neurostimulator generators are undergoing revision, in the absence of another O.R. procedure, these cases would group to a medical MS–DRG. We are inviting public comments on our proposal.

c. External Repair of Hymen
We received a request to examine ICD–10–PCS procedure code 0UQKXXZ (Repair Hymen, External Approach).

This procedure code is currently designated as an O.R. procedure in MS–
DRGs 746 and 747 (Vagina, Cervix and Vulva Procedures with CC/MCC and without CC/MCC, respectively) under
MDC 13. The requestor provided examples and expressed concern that procedure code 0UQKXZZ was assigned to MS-DRG 987 (Non-Extensive O.R. Procedures Unrelated to Principal Diagnosis with MCC) when reported on a maternal delivery claim. The requestor noted that when a similar code was reported with an external approach (for example, procedure code 0UQMXZZ (Repair hymen, external approach)), the case was appropriately assigned to MS-DRG 774 (Vaginal Delivery with Complicating Diagnosis). The requestor stated that the physician documentation was simply more specific to the location of the repair and this should not affect assignment to one of the MS-DRGs for vaginal delivery.

We reviewed claims data involving the examples provided by the requestor involving ICD-10-PCS procedure code 0UQKXZZ (Repair hymen, external approach). Our clinical advisors agree with the requestor that reporting of this procedure code should not affect assignment to one of the MS-DRGs for vaginal delivery. As discussed earlier in section II.F.15.a. of the preamble of this proposed rule, we are proposing to change the designation for a number of procedure codes from O.R. procedures to non-O.R. procedures. Included in that proposal are ICD-10-PCS procedure codes 0UQGXZZ (Repair vagina, external approach) and 0UQMXZZ (Repair vulva, external approach). Consistent with the change in designation for these procedure codes, we also are proposing to designate ICD-10-PCS procedure code 0UQKXZZ (Repair hymen, external approach) as a non-O.R. procedure. The procedure by itself would generally not require the resources of an operating room. If the procedure is performed following a vaginal delivery, it is the vaginal delivery procedure code 10E0XXZ (Delivery of products of conception) that determines the MS-DRG assignment because this code is designated as a non-O.R. procedure affecting the MS-DRG.

Therefore, we are proposing to change the designation of ICD-10-PCS procedure code 0UQKXZZ (Repair hymen, external approach) to a non-O.R. procedure. This redesignation will enable more appropriate MS-DRG assignment for these cases by eliminating erroneous assignment to MS-DRGs 987 through 989. We are inviting public comments on our proposal.

d. Non-O.R. Procedures in MDC 17 (Myeloproliferative Diseases and Disorders, Poorly Differentiated Neoplasms)

Under MDC 17 (Myeloproliferative Diseases and Disorders, Poorly Differentiated Neoplasms), there are 11 surgical MS-DRGs. Of these 11 surgical MS-DRGs, there are 5 MS-DRGs containing GROUPER logic that includes ICD-10-PCS procedure codes designated as O.R. procedures as well as non-O.R. procedures that affect the MS-DRG. These five MS-DRGs are MS-DRGs 823, 824, and 825 (Lymphoma and Non-Acute Leukemia with Other O.R. Procedure with MCC, with CC and without CC/MCC, respectively) and MS-DRGs 829 and 830 (Myeloproliferative Disorders or Poorly Differentiated Neoplasms without O.R. Procedure with CC/MCC and without CC/MCC, respectively). We refer the reader to the ICD-10 Version 34 MS-DRG Definitions Manual which is available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/FY2017-IPPS-Final-Rule-Home-Page-Items/FY2017-IPPS-Final-Rule-Data-Files.html?DLPage=1&DLEntries=10&DLSort=0&DLSortDir=ascending for the complete list of ICD-10-PCS procedure codes assigned to these five MS-DRGs under MDC 17.

We reviewed the list of 244 ICD-10-PCS non-O.R. procedure codes currently assigned to these 5 MS-DRGs. Of these 244 procedure codes, we determined that 55 of the procedure codes do not warrant being designated as non-O.R. procedures that affect these MS-DRGs because they describe procedures that would generally not require a greater intensity of resources for facilities to manage the cases included in the definition (logic) of these MS-DRGs. Therefore, we are proposing that the 55 codes be removed from the logic for MS-DRGs 823, 824, 825, 829 and 830 as non-O.R. procedures affecting the MS-DRG.

In developing the proposed FY 2018 system of weights, we used two data sources: Claims data and cost report data. As in previous years, the claims data source is the MedPAR file. This file is based on fully coded diagnostic and procedure data for all Medicare inpatient hospital stays. The FY 2016 MedPAR data used in this proposed rule includes discharges occurring on October 1, 2015, through September 30, 2016, based on bills received by CMS through December 31, 2016, from all hospitals subject to the IPPS and short-term, acute care hospitals in Maryland (which at that time were under a waiver from the IPPS).

The FY 2016 MedPAR file used in calculating the proposed relative weights includes data for approximately 9,607,103 Medicare discharges from IPPS providers. Discharges for Medicare beneficiaries enrolled in a Medicare Advantage managed care plan are excluded from this analysis. These discharges are excluded when the MedPAR “GH0 Paid” indicator field on the claim record is equal to “1” or when the MedPAR DRG payment field, which represents the total payment for the claim, is equal to the MedPAR “Indirect Medical Education (IME)” payment field, indicating that the claim was an “IME only” claim submitted by a teaching hospital on behalf of a beneficiary enrolled in a Medicare Advantage managed care plan. In addition, the December 31, 2016 update of the FY 2016 MedPAR file complies with version 5010 of the X12 HIPAA Transaction and Code Set Standards, and includes a variable called “claim type.” Claim type “60” indicates that the claim was an inpatient claim paid as fee-for-service. Claim types “61,” “62,” “63,” and “64” relate to encounter claims, Medicare Advantage IME claims, and HMO no-pay claims.

The calculation of the proposed relative weights for FY 2018 also excludes claims with claim type values not equal to “60.” The data exclude CAHs, including hospitals that subsequently became CAHs after the period from which the data were taken. We note that the proposed FY 2018 relative weights are based on the ICD-


10–CM diagnoses and ICD–10–PCS procedure codes from the FY 2016 MedPAR claims data, grouped through the ICD–10 version of the proposed FY 2018 GROUPER (Version 35).

The second data source used in the cost-based relative weighting methodology is the Medicare cost report data files from the HCRIS. Normally, we use the HCRIS dataset that is 3 years prior to the IPPS fiscal year. Specifically, we used cost report data from the December 31, 2016 update of the FY 2015 HCRIS for calculating the proposed FY 2018 cost-based relative weights.

2. Methodology for Calculation of the Proposed Relative Weights

As we explained in section II.E.2. of the preamble of this proposed rule, we calculated the proposed FY 2018 relative weights based on 19 CCRs, as we did for FY 2017. The methodology we are proposing to use to calculate the FY 2018 cost-based relative weights based on claims data in the FY 2016 MedPAR file and data from the FY 2015 Medicare cost reports is as follows. We note that we have provided additional precision in our description of the methodology for FY 2018.

• To the extent possible, all the claims were regrouped using the proposed FY 2018 MS–DRG classifications discussed in sections II.B. and II.F. of the preamble of this proposed rule.

• The transplant cases that were used to establish the proposed relative weights for heart and heart-lung, liver and/or intestinal, and lung transplants (MS–DRGs 001, 002, 005, 006, and 007, respectively) were limited to those Medicare-approved transplant centers that have cases in the FY 2016 MedPAR file. (Medicare coverage for heart, heart-lung, liver and/or intestinal, and lung transplants is limited to those facilities that have received approval from CMS as transplant centers.)

• Organ acquisition costs for kidney, heart, heart-lung, liver, lung, pancreas, and intestinal (or multivisceral organs) transplants continue to be paid on a reasonable cost basis. Because these acquisition costs are paid separately from the prospective payment rate, it is necessary to subtract the acquisition charges from the total charges on each transplant bill that showed acquisition charges before computing the average cost for each MS–DRG and before eliminating statistical outliers.

• Claims with total charges or total lengths of stay less than or equal to zero were deleted. Claims that had an amount in the total charge field that differed by more than $30.00 from the sum of the routine day charges, intensive care charges, pharmacy charges, implantable devices charges, supplies and equipment charges, therapy services charges, operating room charges, cardiology charges, laboratory charges, radiology charges, other service charges, labor and delivery charges, inhalation therapy charges, emergency room charges, blood and blood products charges, anesthesia charges, cardiac catheterization charges, CT scan charges, and MRI charges were also deleted.

• At least 92.2 percent of the providers in the MedPAR file had charges for 14 of the 19 cost centers. All claims of providers that did not have charges greater than zero for at least 14 of the 19 cost centers were deleted. In other words, a provider must have no more than five blank cost centers. If a provider did not have charges greater than zero in more than five cost centers, the claims for the provider were deleted.

• Statistical outliers were eliminated by removing all cases that were beyond 3.0 standard deviations from the geometric mean of the log distribution of both the total charges per case and the total charges per day for each MS–DRG.

• Effective October 1, 2008, because hospital inpatient claims include a POA indicator for each diagnosis present on the claim, only for purposes of relative weight-setting, the POA indicator field was reset to “Y” for “Yes” for all claims that otherwise have an ‘‘N’’ (No) or a ‘‘U’’ (documentation insufficient to determine if the condition was present at the time of inpatient admission) in the POA field.

Under current payment policy, the presence of specific HAC codes, as indicated by the POA field values, can generate a lower payment for the claim. Specifically, if the particular condition is present on admission (that is, a “Y” indicator is associated with the diagnosis on the claim), it is not a HAC, and the hospital is paid for the higher severity (and, therefore, the higher weighted MS–DRG). If the particular condition is not present on admission (that is, an “N” indicator is associated with the diagnosis on the claim) and there are no other complicating conditions, the DRG GROUPER assigns the claim to a lower severity (and, therefore, the lower weighted MS–DRG) as a penalty for allowing a Medicare inpatient to contract a HAC. While the POA reporting meets policy goals of encouraging quality care and generates program savings, it presents an issue for the relative weight-setting process. Because cases identified as HACs are likely to be more complex than similar cases that are not identified as HACs, the charges associated with HAC cases are likely to be higher as well. Therefore, if the higher charges of these HAC claims are grouped into lower severity MS–DRGs prior to the relative weight-setting process, the relative weights of these particular MS–DRGs would become artificially inflated, potentially skewing the relative weights. In addition, we want to protect the integrity of the budget neutrality process by ensuring that, in estimating payments, no increase to the standardized amount occurs as a result of lower overall payments in a previous year that stem from using weights and case-mix that are based on lower severity MS–DRG assignments. If this would occur, the anticipated cost savings from the HAC policy would be lost.

To avoid these problems, we reset the POA indicator field to “Y” only for relative weight-setting purposes for all claims that otherwise have an “N” or a “U” in the POA field. That is, resetting “forced” the more costly HAC claims into the higher severity MS–DRGs as appropriate, and the relative weights calculated for each MS–DRG more closely reflect the true costs of those cases.

In addition, in the FY 2013 IPPS/LTCH PPS final rule, for FY 2013 and subsequent fiscal years, we finalized a policy to treat hospitals that participate in the Bundled Payments for Care Improvement (BPCI) initiative the same as prior fiscal years for the IPPS payment modeling and ratesetting process without regard to hospitals’ participation within these bundled payment models (that is, if hospitals were not participating in those models under the BPCI initiative). The BPCI initiative, developed under the authority of section 3021 of the Affordable Care Act (codified at section 1115A of the Act), is comprised of four broadly defined models of care, which link payments for multiple services beneficiaries receive during an episode of care. Under the BPCI initiative, organizations enter into payment arrangements that include financial and performance accountability for episodes of care. For FY 2018, we are proposing to continue to include all applicable data from subsection (d) hospitals participating in BPCI Models 1, 2, and 4 in our IPPS payment modeling and ratesetting calculations. We refer readers to the FY 2013 IPPS/LTCH PPS final rule for a complete discussion on our final policy for the treatment of hospitals participating in the BPCI initiative in our ratesetting process. For additional information on
The charges for each of the 19 cost groups for each claim were standardized to remove the effects of differences in proposed area wage levels, IME and DSH payments, and for hospitals located in Alaska and Hawaii, the applicable proposed cost-of-living adjustment. Because hospital charges include charges for both operating and capital costs, we standardized total charges to remove the effects of differences in proposed geographic adjustment factors, cost-of-living adjustments, and DSH payments under the capital IPPS as well. Charges were then summed by MS–DRG for each of the 19 cost groups so that each MS–DRG had 19 standardized charge totals. Statistical outliers were then removed. These charges were then adjusted to cost by applying the proposed national average CCRs developed from the FY 2015 cost report data.

The 19 cost centers that we used in the proposed relative weight calculation are shown in the following table. The table shows the lines on the cost report and the corresponding revenue codes that we used to create the proposed 19 national cost center CCRs. If stakeholders have comments about the groupings in this table, we may consider those comments as we finalize our policy.

<table>
<thead>
<tr>
<th>Cost center group name (19 total)</th>
<th>MedPAR charge field</th>
<th>Revenue codes contained in MedPAR charge field</th>
<th>Cost report line description</th>
<th>Cost from HCRIS (Worksheet C, Part 1, Column 5 and line number) Form CMS–2552–10</th>
<th>Charges from HCRIS (Worksheet C, Part 1, Column 6 &amp; 7 and line number) Form CMS–2552–10</th>
<th>Medicare charges from HCRIS (Worksheet D–3, Column &amp; line number) Form CMS–2552–10</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drugs .......................</td>
<td>Pharmacy Charges ...</td>
<td>025X, 026X and 063X.</td>
<td>DME–Sold ..................</td>
<td>C_1.C5.34</td>
<td>C_1.C6.34</td>
<td>D3.HOS.C2.34</td>
</tr>
<tr>
<td>Operating Room ........</td>
<td>Operating Room Charges.</td>
<td>036X ................</td>
<td>Operating Room ..........</td>
<td>C_1.C5.43</td>
<td>C_1.C6.43</td>
<td>D3.HOS.C2.43</td>
</tr>
<tr>
<td>Cardiology ..............</td>
<td>Cardiology Charges</td>
<td>048X and 073X ....</td>
<td>Electrocardiology ..........</td>
<td>C_1.C5.46</td>
<td>C_1.C6.46</td>
<td>D3.HOS.C2.46</td>
</tr>
</tbody>
</table>

For more information on the BPCI initiative, we refer readers to the CMS’ Center for Medicare and Medicaid Innovation’s Web site at: [http://innovation.cms.gov/initiatives/Bundled-Payments/index.html](http://innovation.cms.gov/initiatives/Bundled-Payments/index.html) and to section IV.H.4. of the preamble of the FY 2013 IPPS/LTC PPS final rule (77 FR 53341 through 53343).
<table>
<thead>
<tr>
<th>Cost center group name (19 total)</th>
<th>MedPAR charge field</th>
<th>Revenue codes contained in MedPAR charge field</th>
<th>Cost report line description</th>
<th>Cost from HCRIS (Worksheet C, Part 1, Column 5 and line number) Form CMS–2552–10</th>
<th>Charges from HCRIS (Worksheet C, Part 1, Column 6 &amp; 7 and line number) Form CMS–2552–10</th>
<th>Medicare charges from HCRIS (Worksheet D–3, Column &amp; line number) Form CMS–2552–10</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laboratory</td>
<td>030X, 031X, and 075X</td>
<td>Laboratory</td>
<td>C.1.C5.60</td>
<td>C.1.C6.60</td>
<td>D.3.HOS.C2.60</td>
<td></td>
</tr>
<tr>
<td>CT Scan Charges</td>
<td>035X</td>
<td>Computed Tomography (CT) Scan</td>
<td>C.1.C5.56</td>
<td>C.1.C6.56</td>
<td>D.3.HOS.C2.56</td>
<td></td>
</tr>
<tr>
<td>Outpatient Service Charges</td>
<td>049X</td>
<td>ASC (Non Distinct Part)</td>
<td>C.1.C5.75</td>
<td>C.1.C6.75</td>
<td>D.3.HOS.C2.75</td>
<td></td>
</tr>
<tr>
<td>Lithotripsy Charge</td>
<td>079X</td>
<td>Other Ancillary</td>
<td>C.1.C5.76</td>
<td>C.1.C6.76</td>
<td>D.3.HOS.C2.76</td>
<td></td>
</tr>
<tr>
<td>Clinic Visit Charges</td>
<td>051X</td>
<td>Clinic</td>
<td>C.1.C5.76</td>
<td>C.1.C6.76</td>
<td>D.3.HOS.C2.76</td>
<td></td>
</tr>
<tr>
<td>Professional Fees Charges</td>
<td>096X, 097X, and 098X</td>
<td>Observation beds</td>
<td>C.1.C5.92.01</td>
<td>C.1.C6.92.01</td>
<td>D.3.HOS.C2.92.01</td>
<td></td>
</tr>
<tr>
<td>Rural Health Clinic</td>
<td>055X</td>
<td>Ambulance</td>
<td>C.1.C5.95</td>
<td>C.1.C6.95</td>
<td>D.3.HOS.C2.95</td>
<td></td>
</tr>
</tbody>
</table>

3. Development of Proposed National Average CCRs

We developed the proposed national average CCRs as follows:

Using the FY 2015 cost report data, we removed CAHs, Indian Health Service hospitals, all-inclusive rate hospitals, and cost reports that represented time periods of less than 1 year (365 days). We included hospitals located in Maryland because we included their charges in our claims database. We then created CCRs for each provider for each cost center (see prior table for line items used in the calculations) and removed any CCRs that were greater than 10 or less than 0.01. We normalized the departmental CCRs by dividing the CCR for each department by the total CCR for the hospital for the
The purpose of trimming the data is to identify cost center CCRs and remove any cost center CCRs where the log of the cost center CCR was greater than or less than the mean log plus/minus 3 times the standard deviation for the log of that cost center CCR. Once the cost report data were trimmed, we calculated a Medicare-specific CCR. The Medicare-specific CCR was determined by taking the Medicare charges for each line item from Worksheet D–3 and deriving the Medicare-specific costs by applying the hospital-specific departmental CCRs to the Medicare-specific charges for each line item from Worksheet D–3. Once each hospital’s Medicare-specific costs were established, we summed the total Medicare-specific costs and divided by the sum of the total Medicare-specific charges to produce national average, charge-weighted CCRs.

After we multiplied the total charges for each MS–DRG in each of the 19 cost centers by the corresponding national average CCR, we summed the 19 “costs” across each MS–DRG to produce a total standardized cost for the MS–DRG. The average standardized cost for each MS–DRG was then computed as the total standardized cost for the MS–DRG divided by the transfer-adjusted case count for the MS–DRG. The average cost for each MS–DRG was then divided by the national average standardized cost per case to determine the proposed relative weight.

The proposed FY 2018 cost-based relative weights were then normalized by a proposed adjustment factor of 1.736047 so that the average case weight after recalibration was equal to the average case weight before recalibration. The proposed normalization adjustment is intended to ensure that recalibration by itself neither increases nor decreases total payments under the IPPS, as required by section 1886(d)(4)(C)(iii) of the Act.

The proposed 19 national average CCRs for FY 2018 are as follows:

<table>
<thead>
<tr>
<th>Group</th>
<th>CCR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Routine Days</td>
<td>0.449</td>
</tr>
<tr>
<td>Intensive Days</td>
<td>0.375</td>
</tr>
<tr>
<td>Drugs</td>
<td>0.197</td>
</tr>
<tr>
<td>Supplies &amp; Equipment</td>
<td>0.300</td>
</tr>
<tr>
<td>Implantable Devices</td>
<td>0.327</td>
</tr>
<tr>
<td>Therapy Services</td>
<td>0.314</td>
</tr>
<tr>
<td>Laboratory</td>
<td>0.116</td>
</tr>
<tr>
<td>Operating Room</td>
<td>0.186</td>
</tr>
<tr>
<td>Cardiology</td>
<td>0.108</td>
</tr>
<tr>
<td>Cardiac Catheterization</td>
<td>0.115</td>
</tr>
<tr>
<td>Radiology</td>
<td>0.149</td>
</tr>
<tr>
<td>MRIs</td>
<td>0.077</td>
</tr>
<tr>
<td>CT Scans</td>
<td>0.037</td>
</tr>
</tbody>
</table>

Since FY 2009, the relative weights have been based on 100 percent cost weights based on our MS–DRG grouping system.

When we recalibrated the DRG weights for previous years, we set a threshold of 10 cases as the minimum number of cases required to compute a reasonable weight. We are proposing to use that same case threshold in recalibrating the MS–DRG relative weights for FY 2018. Using data from the FY 2016 MedPAR file, there were 10 MS–DRGs that contain fewer than 10 cases. For FY 2018, because we do not have sufficient MedPAR data to set accurate and stable cost relative weights for these low-volume MS–DRGs, we are proposing to compute proposed relative weights for the low-volume MS–DRGs by adjusting their final FY 2017 relative weights by the percentage change in the average weight of the cases in other MS–DRGs. The crosswalk table is shown:

We are inviting public comments on our proposals.

1. Background

Sections 1886(d)(5)(K) and (L) of the Act establish a process of identifying and ensuring adequate payment for new medical services and technologies: (sometimes collectively referred to in this section as “new technologies”) under the IPPS. Section 1886(d)(5)(K)(vi) of the Act specifies that a medical service or technology will be considered new if it meets criteria established by the Secretary after notice and opportunity for public comment. The regulations at 42 CFR 412.87 implement these provisions and specify three criteria for a new medical service or technology to receive the additional payment: (1) The medical service or technology must be new; (2) the medical service or technology must be costly such that the DRG rate otherwise applicable to such discharges under this subsection is inadequate. We note that, beginning with discharges occurring in FY 2008, CMS transitioned from CMS–DRGs to MS–DRGs.
applicable to discharges involving the medical service or technology is determined to be inadequate; and (3) the service or technology must demonstrate a substantial clinical improvement over existing services or technologies. Below we highlight some of the major statutory and regulatory provisions relevant to the new technology add-on payment criteria, as well as other information.

For a complete discussion on the new technology add-on payment criteria, we refer readers to the FY 2012 IPPS/LTCH PPS final rule (76 FR 51572 through 51574).

Under the first criterion, as reflected in § 412.87(b)(2), a specific medical service or technology will be considered “new” for purposes of new medical service or technology add-on payments until such time as Medicare data are available to fully reflect the cost of the technology in the MS–DRG weights through recalibration. We note that we do not consider a service or technology to be new if it is substantially similar to one or more existing technologies. That is, even if a technology receives a new FDA approval or clearance, it may not necessarily be considered “new” for purposes of new technology add-on payments if it is “substantially similar” to a technology that was approved or cleared by FDA and has been on the market for more than 2 to 3 years. In the FY 2010 IPPS/RY 2010 LTCH PPS final rule (74 FR 43813 through 43814), we established criteria for evaluating whether a new technology is substantially similar to an existing technology:

1. Whether a product uses the same or a similar mechanism of action to achieve a therapeutic outcome; (2) whether a product is assigned to the same or a different MS–DRG; and (3) whether the new use of the technology involves the treatment of the same or similar type of disease and the same or similar patient population. If a technology meets all three of these criteria, it would be considered substantially similar to an existing technology and would not be considered “new” for purposes of new technology add-on payments. For a detailed discussion of the criteria for substantial similarity, we refer readers to the FY 2006 IPPS final rule (70 FR 47351 through 47352), and the FY 2010 IPPS/LTCH PPS final rule (74 FR 43813 through 43814).

Under the second criterion, § 412.87(b)(3) further provides that, to be eligible for the add-on payment for new medical services or technologies, the MS–DRG prospective payment rate otherwise applicable to discharges involving the new medical service or technology must be assessed for adequacy. Under the cost criterion, consistent with the formula specified in section 1886(d)(5)(K)(ii)(I) of the Act, to assess the adequacy of payment for a new technology paid under the applicable MS–DRG prospective payment rate, we evaluate whether the charges for cases involving the new technology exceed certain threshold amounts. Table 10 that was released with the FY 2017 IPPS/LTCH PPS final rule contains the final thresholds that we used to evaluate applications for new medical service and new technology add-on payments for FY 2018. We refer readers to the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/ AcuteInpatientPPS/FY2017-IPPS-Final- Rule-Home-Page-Items/FY2017-IPPS- Final-Rule-Tables.html to download and view Table 10.

In the September 7, 2001 final rule that established the new technology add-on payment regulations (66 FR 46917), we discussed the issue of whether the Health Insurance Portability and Accountability Act (HIPAA) Privacy Rule at 45 CFR parts 160 and 164 applies to claims information that providers submit with applications for new medical service and new technology add-on payments. We refer readers to the FY 2012 IPPS/LTCH PPS final rule (76 FR 51573) for complete information on this issue.

Under the third criterion, § 412.87(b)(1) of our existing regulations provides that a new technology is an appropriate candidate for an additional payment when it represents an advance that substantially improves, relative to technologies previously available, the diagnosis or treatment of Medicare beneficiaries. For example, a new technology represents a substantial clinical improvement when it reduces mortality, decreases the number of hospitalizations or physician visits, or reduces recovery time compared to the technologies previously available. (We refer readers to the September 7, 2001 final rule for a more detailed discussion of this criterion (66 FR 46902)).

The new medical service or technology add-on payment policy under the IPPS provides additional payments for cases with relatively high costs involving eligible new medical services or technologies, while preserving some of the incentives inherent under an average-based prospective payment system. The payment mechanism is based on the cost to hospitals for the new medical service or technology. Under § 412.88, if the new service or technology is determined by applying cost-to-charge ratios (CCRs) as described in § 412.84(h) to exceed the full DRG payment (including payments for IME and DSH, but excluding outlier payments), Medicare will make an additional Medicare payment equal to the lesser of: (1) 50 percent of the estimated costs of the new technology or medical service (if the estimated costs for the case involving the new technology or medical service exceed Medicare’s payment); or (2) 50 percent of the difference between the full DRG payment and the hospital’s estimated cost for the case. Unless the discharge qualifies for an outlier payment, the additional Medicare payment is limited to the full MS–DRG payment plus 50 percent of the estimated costs of the new technology or new medical service.

Section 503(d)(2) of Public Law 108–173 provides that there shall be no reduction or adjustment in aggregate payments under the IPPS due to add-on payments for new medical services and technologies. Therefore, in accordance with section 503(d)(2) of Public Law 108–173, add-on payments for new medical services or technologies for FY 2006 and later years have not been subjected to budget neutrality.

In the FY 2009 IPPS final rule (73 FR 48561 through 48563), we modified our regulations at § 412.87 to codify our longstanding practice of how CMS evaluates the eligibility criteria for new medical service or technology add-on payment applications. That is, we first determine whether a medical service or technology meets the newness criterion, and only if so, do we then make a determination as to whether the technology meets the cost threshold and represents a substantial clinical improvement over existing medical services or technologies. We amended § 412.87(c) to specify that all applicants for new technology add-on payments must have FDA approval or clearance for their new medical service or technology by July 1 of each year prior to the beginning of the fiscal year that the application is being considered.

The Council on Technology and Innovation (CTI) at CMS oversees the agency’s cross-cutting priority on coordinating coverage, coding and payment processes for Medicare with respect to new technologies and procedures, including new drug therapies, as well as promoting the exchange of information on new technologies and medical services between CMS and other entities. The CTI, composed of senior CMS staff and clinicians, was established under section 942(a) of Public Law 108–173. The Council is co-chaired by the Director of the Center for Clinical Standards and Quality (CCSQ) and the Director of the Center for Medicare
The specific processes for coverage, coding, and payment are implemented by CM, CCSQ, and the local Medicare Administrative Contractors (MACs) (in the case of local coverage and payment decisions). The CTI supplements, rather than replaces, these processes by working to assure that all of these activities reflect the agency-wide priority to promote high-quality, innovative care. At the same time, the CTI also works to streamline, accelerate, and improve coordination of these processes to ensure that they remain up to date as new issues arise. To achieve its goals, the CTI works to streamline and create a more transparent coding and payment process, improve the quality of medical decisions, and speed patient access to effective new treatments. It is also dedicated to supporting better decisions by patients and doctors in using Medicare-covered services through the promotion of better evidence development, which is critical for improving the quality of care for Medicare beneficiaries.

To improve the understanding of CMS’ processes for coverage, coding, and payment and how to access them, the CTI has developed an “Innovator’s Guide” to these processes. The intent is to consolidate this information, much of which is already available in a variety of CMS documents and in various places on the CMS Web site, in a user-friendly format. This guide was published in 2010 and is available on the CMS Web site at: [http://www.cms.gov/Innovations/InnovatorsGuide5_10_10.pdf](http://www.cms.gov/Innovations/InnovatorsGuide5_10_10.pdf).

As we indicated in the FY 2009 IPPS final rule (73 FR 48554), we invite any product developers or manufacturers of new medical services or technologies to contact the agency early in the process of product development if they have questions or concerns about the evidence that would be needed later in the development process for the agency’s coverage decisions for Medicare.

The CTI aims to provide useful information on its activities and initiatives to stakeholders, including Medicare beneficiaries, advocates, medical product manufacturers, providers, and health policy experts. Stakeholders with further questions about Medicare’s coverage, coding, and payment processes, or who want further guidance about how they can navigate these processes, can contact the CTI at [CTI@cms.hhs.gov](mailto:CTI@cms.hhs.gov).

In order to receive applications for add-on payments for new medical services or technologies for FY 2019 must submit a formal request, including a full description of the clinical applications of the medical service or technology and the results of any clinical evaluations demonstrating that the new medical service or technology represents a substantial clinical improvement, along with a significant sample of data to demonstrate that the medical service or technology meets the high-cost threshold. Complete application information, along with final deadlines for submitting a full application, will be posted as it becomes available on the CMS Web site at: [http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/newtech.html](http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/newtech.html). To allow interested parties to identify the new medical services or technologies under review before the publication of the proposed rule for FY 2019, the CMS Web site also will post the tracking forms completed by each applicant.

2. Public Input Before Publication of a Notice of Proposed Rulemaking on Add-On Payments

Section 1886(d)(5)(K)(vi) of the Act, as amended by section 503(b)(2) of Public Law 108–173, provides for a mechanism for public input before publication of a notice of proposed rulemaking regarding whether a medical service or technology represents a substantial clinical improvement or advancement. The process for evaluating new medical service and technology applications requires the Secretary to:

- Provide, before publication of a proposed rule, for public input regarding whether a new service or technology represents an advance in medical technology that substantially improves the diagnosis or treatment of Medicare beneficiaries;
- Make public and periodically update a list of the services and technologies for which applications for add-on payments are pending;
- Accept comments, recommendations, and data from the public regarding whether a service or technology represents a substantial clinical improvement; and
- Provide, before publication of a proposed rule, for a meeting at which organizations representing hospitals, physicians, manufacturers, and any other interested party may present comments, recommendations, and data regarding whether a new medical service or technology represents a substantial clinical improvement to the clinical staff of CMS.

In order to provide an opportunity for public input regarding add-on payments for new medical services and technologies for FY 2018 prior to publication of the FY 2018 IPPS/LTCH PPS proposed rule, we published a notice in the Federal Register on November 9, 2016 (81 FR 78814), and held a town hall meeting at the CMS Headquarters Office in Baltimore, MD, on February 14, 2017. In the announcement notice for the meeting, we stated that the opinions and presentations provided during the meeting would assist us in our evaluations of applications by allowing public discussion of the substantial clinical improvement criterion for each of the FY 2018 new medical service and technology add-on payment applications before the publication of the FY 2018 IPPS/LTCH PPS proposed rule.

Approximately 66 individuals registered to attend the town hall meeting in person, while additional individuals listened over an open telephone line. We also live-streamed the town hall meeting and posted the town hall on the CMS YouTube Web page at: [https://www.youtube.com/watch?v=9niqfxE4oA&t=217s](https://www.youtube.com/watch?v=9niqfxE4oA&t=217s). We considered each applicant’s presentation made at the town hall meeting, as well as written comments submitted on the applications that were received by the due date of February 24, 2017, in our evaluation of the new technology add-on payment applications for FY 2018 in this proposed rule.

In response to the published notice and the February 14, 2017 New Technology Town Hall meeting, we received written comments regarding the applications for FY 2018 new technology add-on payments. We note that we do not summarize comments that are unrelated to the “substantial clinical improvement” criterion. As explained above and in the Federal Register notice announcing the New Technology Town Hall meeting (81 FR78814 through 78816), the purpose of the meeting was specifically to discuss the substantial clinical improvement criterion in regard to pending new technology add-on payment applications for FY 2018. Therefore, we are not summarizing these comments in this proposed rule. We summarize below a general comment that does not relate to a specific application for FY 2018 new technology add-on payments. We also summarize comments regarding individual applications, or, if applicable, indicate that there were no comments received in section II.H.5. of the preamble of this proposed rule at the end of each discussion of the individual applications.
Comment: One commenter recommended that CMS: (1) Prohibit local MACs from denying coverage and add-on payments for new medical services or technologies approved by the Secretary; and (2) broaden the criteria applied in making substantial clinical improvement determinations to require, in addition to existing criteria, that the Secretary consider whether the new technology or medical service meets one or more of the following criteria: (a) Results in a reduction of the length of a hospital stay; (b) improves patient quality of life; (c) creates long-term clinical efficiencies in treatment; (d) addresses patient-centered objectives as defined by the Secretary; or (e) meets such other criteria as the Secretary may specify.

Response: We appreciate the commenter’s comments and will consider them in future rulemaking.

3. ICD–10–PCS Section “X” Codes for Certain New Medical Services and Technologies

As discussed in the FY 2016 IPPS/LTCH final rule (80 FR 49434), the ICD–10–PCS includes a new section containing the new Section “X” codes, which began being used with discharges occurring on or after October 1, 2015. Decisions regarding changes to ICD–10–PCS Section “X” codes will be handled in the same manner as the decisions for all of the other ICD–10–PCS code changes. That is, proposals to create, delete, or revise Section “X” codes under the ICD–10–PCS structure will be referred to the ICD–10 Coordination and Maintenance Committee. In addition, several of the new medical services and technologies that have been, or may be, approved for new technology add-on payments may now, and in the future, be assigned a Section “X” code within the structure of the ICD–10–PCS. We posted ICD–10–PCS Guidelines on the CMS Web site at: http://www.cms.gov/Medicare/Coding/ICD10/2016-ICD-10-PCS-and-GEMS.html, including guidelines for ICD–10–PCS Section “X” codes. We encourage providers to view the material provided on ICD–10–PCS Section “X” codes.

4. Proposal To Revise the Reference to an ICD–9–CM Code in § 412.87(b)(2) of the Regulations

The existing regulations under § 412.87(b)(2) state that a medical service or technology may be considered new within 2 or 3 years after the point at which data begin to become available reflecting the ICD–9–CM code assigned to the new service or technology (depending on when a new code is assigned and data on the new service or technology become available for DRG recalculation). After CMS has recalibrated the DRGs, based on available data, to reflect the costs of an otherwise new medical service or technology, the medical service or technology will no longer be considered “new” under the criterion of this section.

As discussed in the FY 2016 IPPS/LTCH final rule (80 FR 49454), HIPAA covered entities are required, as of October 1, 2015, to use the ICD–10 coding system (ICD–10–PCS codes for procedures and ICD–10–CM codes for diagnoses). Instead of the ICD–9–CM coding system, to report diagnoses and procedures for Medicare hospital inpatient services provided to Medicare beneficiaries as classified under the MS–DRG system and paid for under the IPPS. The language in § 412.87(b)(2) only references an “ICD–9–CM code.” Therefore, we are proposing to revise the regulations at § 412.87(b)(2) to replace the term “ICD–9–CM code” with the term “inpatient hospital code,” as defined in section 1886(d)(i)(K)(ii) of the Act. Section 1886(d)(i)(K)(iii) of the Act defines an “inpatient hospital code” as any code that is used with respect to inpatient hospital services for which payment may be made under this subsection of the Act and includes an alphanumeric code issued under the International Classification of Diseases, 9th Revision, Clinical Modification (“ICD–9–CM”) and its subsequent revisions. We are inviting public comments on our proposal.

5. Proposed FY 2018 Status of Technologies Approved for FY 2017 Add-On Payments

a. CardioMEMSTM HF (Heart Failure) Monitoring System

CardioMEMSTM, Inc. submitted an application for new technology add-on payments for FY 2015 for the CardioMEMSTM HF (Heart Failure) Monitoring System, which is an implantable hemodynamic monitoring system comprised of an implantable sensor/monitor placed in the distal pulmonary artery. Pulmonary artery hemodynamic monitoring is used in the management of heart failure. The CardioMEMSTM HF Monitoring System measures multiple pulmonary artery pressure parameters for an ambulatory patient to measure and transmit data via a wireless sensor to a secure Web site. The CardioMEMSTM HF Monitoring System utilizes radiofrequency (RF) energy to power the sensor and to measure pulmonary artery (PA) pressure and consists of three components: An Implantable Sensor with Delivery Catheter, an External Electronics Unit, and a Pulmonary Artery Pressure Database. The system provides the physician with the patient’s PA pressure waveform (including systolic, diastolic, and mean pressures) as well as heart rate. The sensor is permanently implanted in the distal pulmonary artery using transcatheter techniques in the catheterization laboratory where it is calibrated using a Swan-Ganz catheter. PA pressures are transmitted by the patient at home in a supine position on a padded antenna, pushing one button which records an 18-second continuous waveform. The data also can be recorded from the hospital, physician’s office, or clinic.

The hemodynamic data, including a detailed waveform, are transmitted to a secure Web site that serves as the Pulmonary Artery Pressure Database, so that information regarding PA pressure is available to the physician or nurse at any time via the Internet. Interpretation of trend data allows the clinician to make adjustments to therapy and can be used along with heart failure signs and symptoms to adjust medications.

The applicant received FDA approval on May 28, 2014. After evaluation of the newness, costs, and substantial clinical improvement criteria for new technology add-on payments for the CardioMEMSTM HF Monitoring System and consideration of the public comments we received in response to the FY 2015 IPPS/LTCH PPS proposed rule, we approved the CardioMEMSTM HF Monitoring System for new technology add-on payments for FY 2015 (79 FR 49940). Cases involving the CardioMEMSTM HF Monitoring System that are eligible for new technology add-on payments are identified by either ICD–10–PCS procedure code 02HQ30Z (Insertion of pressure sensor monitoring device into right pulmonary artery, percutaneous approach) or ICD–10–PCS procedure code 02HR30Z (Insertion of pressure sensor monitoring device into left pulmonary artery, percutaneous approach). With the new technology add-on payment application, the applicant stated that the total operating cost of the CardioMEMSTM HF Monitoring System is $17,750. Under § 412.88(a)(2), we limit new technology add-on payments to the lesser of 50 percent of the average cost of the device or 50 percent of the costs in excess of the MS–DRG payment for the case. As a result, the maximum new technology add-on payment for a case involving the CardioMEMSTM HF Monitoring System is $8,875. We refer the reader to the FY 2015 IPPS/LTCH PPS Final Rule (79 FR 49937) for complete details on the CardioMEMSTM HF Monitoring System.
Our policy is that a medical service or technology may be considered new within 2 or 3 years after the point at which data begin to become available reflecting the inpatient hospital code assigned to the new service or technology. Our practice has been to begin and end new technology add-on payments on the basis of a fiscal year, and we have generally followed a guideline that uses a 6-month window before and after the start of the fiscal year to determine whether to extend the new technology add-on payment for an additional fiscal year. In general, we extend add-on payments for an additional year only if the 3-year anniversary date of the product’s entry onto the U.S. market occurs in the latter half of the fiscal year (70 FR 47362).

With regard to the newness criterion for the CardioMEMSTM HF Monitoring System, we considered the beginning of the newness period to commence when the CardioMEMSTM HF Monitoring System was approved by the FDA on May 28, 2014. The 3-year anniversary date of the entry of the CardioMEMSTM HF Monitoring System onto the U.S. market (May 28, 2017) will occur prior to the beginning of FY 2018. Therefore, we are proposing to discontinue new technology add-on payments for this technology for FY 2018. We are inviting public comments on this proposal.

b. Defitelio® (Defibrotide)

Jazz Pharmaceuticals submitted an application for new technology add-on payments for FY 2017 for defibrotide (Defitelio®), a treatment for patients diagnosed with hepatic veno-occlusive disease (VOD) with evidence of multiorgan dysfunction. VOD, also known as sinusoidal obstruction syndrome (SOS), is a potentially life-threatening complication of hematopoietic stem cell transplantation (HSCT), with an incidence rate of 8 percent to 15 percent. Diagnoses of VOD range in severity from what has been classically defined as a disease limited to the liver (mild) and reversible, to a severe syndrome associated with multiorgan dysfunction or failure and death. Patients treated with HSCT who develop VOD with multi-organ failure face an immediate risk of death, with a mortality rate of more than 80 percent when only supportive care is used. The applicant asserted that Defitelio® improves the survival rate of patients diagnosed with VOD with multi-organ failure by 23 percent.

Defitelio® was granted Orphan Drug Designation for the treatment of VOD in 2003 and for prevention of VOD in 2007. It has been available to patients as an investigational drug through an expanded access program since 2007. The applicant’s New Drug Application (NDA) for Defitelio® received FDA approval on March 30, 2016. The applicant confirmed that Defitelio® was not available on the U.S. market as of the FDA NDA approval date of March 30, 2016. According to the applicant, commercial packaging could not be completed until the label for Defitelio® was finalized with FDA approval, and that commercial shipments of Defitelio® to hospitals and treatment centers began on April 4, 2016. Therefore, we agreed that, based on this information, the newness period for Defitelio® begins on April 4, 2016, the date of its first commercial availability.

The applicant received unique ICD-10-PCS procedure codes to describe the use of Defitelio® that became effective October 1, 2016. The approved procedure codes are XW03392 (Introduction of defibrotide sodium anticoagulant into peripheral vein, percutaneous approach) and XW04392 (Introduction of defibrotide sodium anticoagulant into central vein, percutaneous approach).

After evaluation of the newness, costs, and substantial clinical improvement criteria for new technology add-on payments for Defitelio® and consideration of the public comments we received in response to the FY 2017 IPPS/LTCF PPS proposed rule, we approved Defitelio® for new technology add-on payments for FY 2017 (81 FR 56906). With the new technology add-on payment application, the applicant estimated that any Medicare beneficiary would require a dosage of 25 mg/kg/day for a minimum of 21 days of treatment. The recommended dose is 6.25 mg/kg given as a 2-hour intravenous infusion every 6 hours. Dosing should be based on a patient’s baseline body weight, which is assumed to be 70 kg for an average adult patient. All vials contain 200 mg at a cost of $825 per vial. Therefore, we determined that cases involving the use of the Defitelio® technology would incur an average cost per case of $151,800 (70 kg adult × 25 mg/kg/day × 21 days = 36,750 mg per patient/200 mg vial = 184 vials per patient × $825 per vial = $151,800). Under § 412.88(a)(2), we limit new technology add-on payments to the lesser of 50 percent of the average cost of the technology or 50 percent of the costs in excess of the MS–DRG payment for the case. As a result, the annual new technology add-on payment amount for a case involving the use of Defitelio® is $75,900. Because the 3-year anniversary date of the entry of Defitelio® onto the U.S. market will occur after FY 2018 (April 4, 2019), we are proposing to continue new technology add-on payments for this technology for FY 2018. The maximum payment for a case involving Defitelio® would remain at $75,900 for FY 2018. We are inviting public comments on our proposal to continue new technology add-on payments for Defitelio®.

c. GORE® EXCLUDER® Iliac Branch Endoprosthesi(Gore IBE Device)

W. L. Gore and Associates, Inc. submitted an application for new technology add-on payments for the GORE® EXCLUDER® Iliac Branch Endoprosthesis (GORE IBE device) for FY 2017. The device consists of two components: The Iliac Branch Component (IBC) and the Internal Iliac Component (IIC). The applicant indicated that each endoprosthesi is pre-mounted on a customized delivery and deployment system allowing for controlled endovascular delivery via bilateral femoral access. According to the applicant, the device is designed to be used in conjunction with the GORE® EXCLUDER® AAA Endoprosthesis for the treatment of patients requiring repair of common iliac or aortoiliac aneurysms. When deployed, the GORE IBE device excludes the common iliac aneurysm from systemic blood flow, while preserving blood flow in the external and internal iliac arteries.

With regard to the newness criterion, the applicant received pre-market FDA approval of the GORE IBE device on February 29, 2016. The applicant submitted a request for an unique ICD-10-PCS procedure code and was granted approval for the following procedure codes to describe to use of this technology: 04VC0EZ (Restriction of right common iliac artery with branched or fenestrated intraluminal device, one or two arteries, open approach); 04VC0DFZ (Restriction of right common iliac artery with branched or fenestrated intraluminal device, three or more arteries, open approach); 04VC3EZ (Restriction of right common iliac artery with branched or fenestrated intraluminal device, one or two arteries, percutaneous approach); 04VC3FZ (Restriction of right common iliac artery with branched or fenestrated intraluminal device, three or more arteries, percutaneous approach); 04VC4EZ (Restriction of right common iliac artery with branched or fenestrated intraluminal device, one or two arteries, percutaneous approach); 04VC4FZ (Restriction of right common iliac artery with branched or fenestrated intraluminal device, three or more arteries, percutaneous approach); 04VDOEZ (Restriction of left
common iliac artery with branched or fenestrated intraluminal device, one or two arteries, open approach); 04VD0FZ (Restriction of left common iliac artery with branched or fenestrated, intraluminal device, three or more arteries, open approach); 04VD3EZ (Restriction of left common iliac artery with branched or fenestrated intraluminal device, one or two arteries, percutaneous approach); 04VD3FZ (Restriction of left common iliac artery with branched or fenestrated intraluminal device, three or more arteries, percutaneous approach); 04VD4EZ (Restriction of left common iliac artery with branched or fenestrated intraluminal device, one or two arteries, percutaneous endoscopic approach); and 04VD4FZ (Restriction of left common iliac artery with branched or fenestrated intraluminal device, three or more arteries, percutaneous endoscopic approach). These new ICD–10–PCS procedure codes became effective on October 1, 2016.

After evaluation of the newness, costs, and substantial clinical improvement criteria for new technology add-on payments for the GORE IBE device and consideration of the public comments we received in response to the FY 2017 IPPS/LTCH PPS proposed rule, we approved the GORE IBE device for new technology add-on payments for FY 2017 (81 FR 56909). With the new technology add-on payment application, the applicant indicated that the total operating cost of the GORE IBE device is $10,500. Under § 412.88(a)(2), we limit new technology add-on payments to the lesser of 50 percent of the average cost of the device or 50 percent of the costs in excess of the MS–DRG payment for the case. As a result, the maximum new technology add-on payment for a case involving the GORE IBE device is $5,250.

With regard to the newness criterion for the GORE IBE device, we considered the beginning of the newness period to commence when Idarucizumab was approved by the FDA on October 16, 2015. Because the 3-year anniversary date of the entry of Idarucizumab onto the U.S. market will occur after FY 2018 (October 15, 2018), we are proposing to continue new technology add-on payments for this technology for FY 2018. The maximum payment for a case involving Idarucizumab would remain at $1,750. We are inviting public comments on our proposal to continue new technology add-on payments for Idarucizumab.

e. Lutonix® Drug Coated Balloon PTA Catheter and In.PACT™ Admiral™ Paclitaxel Coated Percutaneous Transluminal Angioplasty (PTA) Balloon Catheter

Two manufacturers, CR Bard Inc. and Medtronic, submitted applications for new technology add-on payments for FY 2016 for LUTONIX® Drug-Coated Balloon (DCB) Percutaneous Transluminal Angioplasty (PTA) Catheter (LUTONIX®) and In.PACT™ Admiral™ Paclitaxel Coated Percutaneous Transluminal Angioplasty (PTA) Balloon Catheter (In.PACT™ Admiral™), respectively. Both of these technologies are drug-coated balloon angioplasty treatments for patients diagnosed with peripheral artery disease (PAD). Typical treatments for patients with PAD include angioplasty, stenting, atherectomy and vascular bypass surgery. PAD most commonly occurs in the femoropopliteal segment of the peripheral arteries, is associated with significant levels of morbidity and impairment in quality of life, and requires treatment to reduce symptoms and prevent or treat ischemic events.1

Treatment options for symptomatic PAD include noninvasive treatment such as medication and life-style modification (for example, exercise programs, diet, and smoking cessation) and invasive options, which include endovascular treatment and surgical bypass. The 2013 American College of Cardiology and American Heart Association (ACC/AHA) guidelines for the management of PAD recommend endovascular therapy as the first-line treatment for femoropopliteal artery lesions in patients suffering from claudication (Class I, Level A recommendation).2

According to both applicants, LUTONIX® and IN.PACT™ Admiral™ are the first drug coated balloons that can be used for treatment of patients who are diagnosed with PAD. In the FY 2016 IPPS/LTC Ha final rule, we stated that cases eligible for the two devices would group to the same MS–DRGs and we believe that these devices are substantially similar to each other (that is, they are intended to treat the same or similar disease in the same or similar patient population and are purposed to achieve the same therapeutic outcome using the same or similar mechanism of action), we evaluated both technologies as one application for new technology add-on payments under the IPPS. The applicants submitted separate cost and clinical data, and we reviewed and discussed each set of data separately. However, we made one determination regarding new technology add-on payments that applied to both devices. We believe that this is consistent with our policy statements in the past regarding substantial similarity. Specifically, we have noted that approval of new technology add-on payments would extend to all technologies that are substantially similar (66 FR 46915), and we believe that continuing our current practice of extending a new technology add-on payment without a further application from the manufacturer of the competing product or a specific finding on cost and clinical improvement if we make a finding of substantial similarity among two products is the better policy because we avoid—

- Creating manufacturer-specific codes for substantially similar products;
- Requiring different manufacturers of substantially similar products from having to submit separate new technology add-on payment applications;
- Having to compare the merits of competing technologies on the basis of substantial clinical improvement; and
- Bestowing an advantage to the first applicant representing a particular new technology to receive approval (70 FR 47351).

<table>
<thead>
<tr>
<th>ICD–10–PCS code</th>
<th>Code description</th>
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<tbody>
<tr>
<td>047K041</td>
<td>Dilation of right femoral artery with drug-eluting intraluminal device using drug-coated balloon, open approach.</td>
</tr>
<tr>
<td>047K0D1</td>
<td>Dilation of right femoral artery with intraluminal device using drug-coated balloon, open approach.</td>
</tr>
<tr>
<td>047K0Z2</td>
<td>Dilation of right femoral artery using drug-coated balloon, open approach.</td>
</tr>
<tr>
<td>047K041</td>
<td>Dilation of right femoral artery with drug-eluting intraluminal device using drug-coated balloon, percutaneous approach.</td>
</tr>
<tr>
<td>047K0D1</td>
<td>Dilation of right femoral artery with intraluminal device using drug-coated balloon, percutaneous approach.</td>
</tr>
<tr>
<td>047K0Z1</td>
<td>Dilation of right femoral artery using drug-coated balloon, percutaneous approach.</td>
</tr>
<tr>
<td>047K441</td>
<td>Dilation of right femoral artery with drug-eluting intraluminal device using drug-coated balloon, percutaneous endoscopic approach.</td>
</tr>
<tr>
<td>047K4D1</td>
<td>Dilation of right femoral artery with intraluminal device using drug-coated balloon, percutaneous endoscopic approach.</td>
</tr>
<tr>
<td>047K4Z1</td>
<td>Dilation of left femoral artery with drug-eluting intraluminal device using drug-coated balloon, open approach.</td>
</tr>
<tr>
<td>047L041</td>
<td>Dilation of left femoral artery with intraluminal device using drug-coated balloon, open approach.</td>
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<tr>
<td>047L0Z1</td>
<td>Dilation of left femoral artery using drug-coated balloon, open approach.</td>
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<tr>
<td>047L341</td>
<td>Dilation of left femoral artery with drug-eluting intraluminal device using drug-coated balloon, percutaneous approach.</td>
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<tr>
<td>047L3D1</td>
<td>Dilation of left femoral artery with intraluminal device using drug-coated balloon, percutaneous approach.</td>
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<tr>
<td>047M041</td>
<td>Dilation of right popliteal artery with drug-eluting intraluminal device using drug-coated balloon, open approach.</td>
</tr>
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LUTONIX®

which resulted in the following case-

equalled 34,875 cases (26,000 plus

cases for each of the applicants, which

compute the weighted cost average, we

the maximum new technology add-on

based on the projected number of cases

captured by using the same ICD–10–PCS

codes are not manufacturer specific, we

manufacturer specific). Because ICD–10

codes listed above (which are not

described by the ICD–10–PCS procedure

tcology add-on payment for cases

As discussed in the FY 2016 IPPS/

LTCH final rule (80 FR 49469), each of the applicants submitted operating costs

g for its DCB. The manufacturer of the

LUTONIX® stated that a mean of 1.37

drug-coated balloons was used during

during the LEVANT 2 clinical trial. The

acquisition price for the hospital will be

$1,900 per drug-coated balloon, or

$2,603 per case (1.37 × $1,900). The

applicant projected that approximately

8,875 cases will involve use of the

LUTONIX® for FY 2016. The

manufacturer for the IN.PACT™

Admiral™ stated that a mean of 1.4

drug-coated balloons was used during

during the IN.PACT™ Admiral™ DCB arm. The

acquisition price for the hospital will be

$1,350 per drug-coated balloon, or

$1,890 per case (1.4 × $1,350). The

applicant projected that approximately

26,000 cases will involve use of the

IN.PACT™ Admiral™ for FY 2016.

For FY 2016, we based the new

technology add-on payment for cases

involving these technologies on the

weighted average cost of the two DCBs

described by the ICD–10–PCS procedure
codes listed above (which are not

manufacturer specific). Because ICD–10
codes are not manufacturer specific, we

cannot set one new technology add-on

payment amount for IN.PACT™

Admiral™ and a different new

technology add-on payment amount for

LUTONIX®; both technologies will be

captured by using the same ICD–10–PCS

procedure code. As such, we stated that

we believe that the use of a weighted

average of the cost of the standard DCBs

based on the projected number of cases

involving each technology to determine

the maximum new technology add-on

payment would be most appropriate. To

compute the weighted cost average, we

summed the total number of projected
cases for each of the applicants, which

equalled 34,875 cases (26,000 plus

8,875). We then divided the number of

projected cases for each of the

applicants by the total number of cases,

which resulted in the following case-

weighted percentages: 25 percent for the

LUTONIX® and 75 percent for the

IN.PACT™ Admiral™. We then

multiplied the cost per case for the

manufacturer specific DCB by the case-

weighted percentage (0.25 × $2,603 =

$662.41 for LUTONIX® and 0.75 ×

$1,890 = $1,409.03 for the IN.PACT™

Admiral™). This resulted in a case-

weighted average cost of $2,071.45 for

DCBs. Under § 412.88(a)(2), we limit

new technology add-on payments to the

lesser of 50 percent of the average cost

of the device or 50 percent of the costs

in excess of the MS–DRG payment for

the case. As a result, the maximum

payment for a case involving the

LUTONIX® or IN.PACT™ Admiral™

DCBs is $1,035.72.

With regard to the newness criterion

for the LUTONIX® and IN.PACT™

Admiral™ technologies, we considered

the beginning of the newness period to

commence when LUTONIX® gained

entry onto the U.S. market on October

10, 2014. As discussed previously in

this section, in general, we extend new

technology add-on payments for an

additional year only if the 3-year

anniversary date of the product’s entry

onto the U.S. market occurs in the latter

half of the upcoming fiscal year.

Because the 3-year anniversary date of

the entry of LUTONIX® onto the U.S.

market (October 10, 2017) will occur in

the first half of FY 2018, we are

proposing to discontinue new

technology add-on payments for both

the LUTONIX® and IN.PACT™

Admiral™ technologies for FY 2018.

We are inviting public comments on

this proposal.

f. MAGEC® Spinal Bracing and

Distraction System (MAGEC® Spine)

Ellipse Technologies, Inc. submitted

an application for new technology add-
on payments for FY 2017 for the

MAGEC® Spine. According to the

applicant, the MAGEC® Spine has been

developed for use in the treatment of

children diagnosed with severe spinal

deformities, such as scoliosis. The

system can be used in the treatment of

skeletally immature patients less than

10 years of age who have been

diagnosed with severe progressive

spinal deformities associated with or at

risk of Thoracic Insufficiency Syndrome

(TIS).

The MAGEC® Spine consists of a

(spine growth) rod that can be

lengthened through the use of magnets

that are controlled by an external remote

controller (ERC). The rod(s) can be

implanted into children as young as 2

years of age. According to the applicant,

use of the MAGEC® Spine has proven to

be successfully used in the treatment of

patients diagnosed with scoliosis who

have not been responsive to other

treatments.

The MAGEC® Spine initially received

FDA clearance for use of the predicate

device, which used a Harrington Rod on

February 27, 2014. The applicant

verified that, due to manufacturing

delays, the MAGEC® Spine was not

available for implant until April 1, 2014.

Specifically, the complete MAGEC®

Spine system was produced and

available for shipment for the first

implant on April 1, 2014. Therefore, the

newness period for the MAGEC® Spine

began on April 1, 2014. Subsequent

FDA clearance was granted for use of the

modified device, which uses a shorter

70 mm rod on September 18, 2014. After

minor modification of the product, the

MAGEC® Spine received

FDA clearances on March 24, 2015, and

May 29, 2015, respectively.

After evaluation of the newness, costs,

and substantial clinical improvement

criteria for new technology add-on

payments for the MAGEC® Spine and

consideration of the public comments

we received in response to the FY 2017

IPPS/LTCH PPS proposed rule, we

approved the MAGEC® Spine for new

technology add-on payments for FY

2017 (81 FR 56891). Cases involving the

MAGEC® Spine that are eligible for new

technology add-on payments are

identified by ICD–10–PCS procedure

codes XNS0032 (Reposition of lumbar

vertebra using magnetically controlled
growth rod(s), open approach);

XNS0432 (Reposition of lumbar vertebra

using magnetically controlled growth

growth

procedure code.

ICD–10–PCS code Code description

047M4Z1 .............. Dilation of right popliteal artery using drug-coated balloon, percutaneous endoscopic approach.

047N0D1 .............. Dilation of left popliteal artery with intraluminal device using drug-coated balloon, open approach.

047N0Z1 .............. Dilation of left popliteal artery using drug-coated balloon, open approach.

047N341 .............. Dilation of left popliteal artery with drug-eluting intraluminal device using drug-coated balloon, percutaneous approach.

047N3D1 .............. Dilation of left popliteal artery with intraluminal device using drug-coated balloon, percutaneous approach.

047N3Z1 .............. Dilation of left popliteal artery with drug-eluting intraluminal device using drug-coated balloon, percutaneous endoscopic approach.

047N4D1 .............. Dilation of left popliteal artery with intraluminal device using drug-coated balloon, percutaneous endoscopic approach.

047N4Z1 .............. Dilation of left popliteal artery using drug-coated balloon, percutaneous endoscopic approach.
rod(s), percutaneous endoscopic approach); XNS3032 (Reposition of cervical vertebra using magnetically controlled growth rod(s), open approach); XNS3342 (Reposition of cervical vertebra using magnetically controlled growth rod(s), percutaneous endoscopic approach); XNS4032 (Reposition of thoracic vertebra using magnetically controlled growth rod(s), open approach); and XNS4432 (Reposition of thoracic vertebra using magnetically controlled growth rod(s)).

With the new technology add-on payment application, the applicant stated that the total operating cost of the MAGEC Spine was $17,500 for a single rod and $35,000 for a dual rod. It is historical practice for CMS to make the new technology add-on payment based on the average cost of the technology and not the maximum. For example, in the FY 2013 IPPS/LTCH PPS final rule (77 FR 53358), we approved new technology add-on payments for DIFICID based on the average dosage of 6.2 days, rather than the maximum 10-day dosage. The applicant noted that 20 percent of cases use a single rod, while 80 percent of cases use a dual rod. As a result, the weighted average cost for a single and dual MAGEC Spine is $31,500 (((0.2 * $17,500) + (0.8 * $35,000))). Under § 412.88(a)(2), we limit new technology add-on payments to the lesser of 50 percent of the average cost of the device or 50 percent of the costs in excess of the MS–DRG payment for the case. As a result, the maximum new technology add-on payment for a case involving Vistogard is $15,750. We refer the reader to the FY 2017 IPPS/LTCH PPS final rule (81 FR 56888) for complete details on the MAGEC Spine.

With regard to the newness criterion for the MAGEC Spine, we considered the beginning of the newness period to commence when the MAGEC Spine was produced and available for shipment for the first implant on April 1, 2014. As discussed previously in this section, in general, we extend new technology add-on payments for an additional year only if the 3-year anniversary date of the product’s entry onto the U.S. market occurs in the latter half of the upcoming fiscal year. Because the 3-year anniversary date of the entry of the MAGEC Spine onto the U.S. market (April 1, 2017) will occur prior to the beginning of FY 2018, we are proposing to discontinue new technology add-on payments for this technology for FY 2018. We are inviting public comments on this proposal.

g. Vistogard™ (Uridine Triacetate)

BTG International Inc., submitted an application for new technology add-on payments for the Vistogard™ for FY 2017. Vistogard™ was developed as an antidote to Fluorouracil toxicity. Chemotherapeutic agent 5-fluorouracil (5–FU) is used to treat specific solid tumors. It acts upon deoxyribonucleic acid (DNA) and ribonucleic acid (RNA) in the body, as uracil is a naturally occurring building block for genetic material. Fluorouracil is a fluorinated pyrimidine. As a chemotherapy agent, Fluorouracil is absorbed by cells and causes the cell to metabolize into byproducts that are toxic and used to destroy cancerous cells. According to the applicant, the byproducts fluorouridine monophosphate (F–UMP) and fluorouridine triphosphate (FUTP) are believed to do the following: (1) Reduce DNA synthesis; (2) lead to DNA fragmentation; and (3) disrupt RNA synthesis. Fluorouracil is used to treat a variety of solid tumors such as colorectal, head and neck, breast, and ovarian cancer. Different tumor treatments, different dosages, and different dosing schedules, there is a risk for toxicity in these patients. Patients may suffer from fluorouracil toxicity/death if 5–FU is delivered in slight excess or at faster infusion rates than prescribed. The cause of overdose can happen for a variety of reasons including: Pump malfunction, incorrect pump programming or miscalculated doses, and accidental or intentional ingestion.

Vistogard™ is an antidote to Fluorouracil toxicity and is a prodrug of uridine. Once the drug is metabolized into uridine, it competes with the toxic byproduct FUTP in binding to RNA, thereby reducing the impact FUTP has on cell death.

The Vistogard™ received FDA approval on December 11, 2015. In the FY 2017 IPPS/LTCH PPS final rule (81 FR 56910), we stated that we agreed with the manufacturer that, due to the delay in availability, the date the newness period begins for Vistogard™ is March 2, 2016, instead of December 11, 2015.

The applicant noted that the Vistogard™ is the first FDA-approved antidote used to reverse fluorouracil toxicity. The applicant received a unique ICD–10–PCS procedure code that became effective October 1, 2016, to describe the use of this technology. The approved procedure code is WX0DX82 (Ingestion of Uracil Triacetate into Mouth and Pharynx, External Approach, New Technology Group 2).

After evaluation of the newness, costs, and substantial clinical improvement criteria for new technology add-on payments for Vistogard™ and consideration of the public comments we received in response to the FY 2017 IPPS/LTCH PPS proposed rule, we approved Vistogard™ for new technology add-on payments for FY 2017 (81 FR 56912). With the new technology add-on payment application, the applicant stated that the total operating cost of Vistogard™ is $75,000. Under § 412.88(a)(2), we limit new technology add-on payments to the lesser of 50 percent of the average cost of the technology or 50 percent of the costs in excess of the MS–DRG payment for the case. As a result, the maximum new technology add-on payment for a case involving Vistogard™ is $37,500.

As noted previously, with regard to the newness criterion for the Vistogard™, we considered the beginning of the newness period to commence on March 2, 2016. Because the 3-year anniversary date of the entry of the Vistogard™ onto the U.S. market (March 2, 2019) will occur after FY 2018, we are proposing to continue new technology add-on payments for this technology for FY 2018. The maximum payment for a case involving the Vistogard™ would remain at $37,500 for FY 2018. We are inviting public comments on our proposal to continue new technology add-on payments for the Vistogard™.

h. Blinatumomab (BLINCYTO®)

Amgen, Inc. submitted an application for new technology add-on payments for FY 2016 for Blinatumomab (BLINCYTO®), a bi-specific T-cell engager (BiTE) used for the treatment of Philadelphia chromosome-negative (Ph−) relapsed or refractory (R/R) B-cell precursor acute lymphoblastic leukemia (ALL), which is a rare aggressive cancer of the blood and bone marrow. Approximately 6,050 individuals are diagnosed with Ph− R/R B-cell precursor ALL in the United States each year, and approximately 2,400 individuals, representing 30 percent of all new cases, are adults. Ph− R/R B-cell precursor ALL occurs when there are malignant transformations of B-cell or T-cell progenitor cells, causing an accumulation of lymphoblasts in the blood, bone marrow, and occasionally throughout the body. As a bi-specific T-cell engager, the BLINCYTO® technology attaches to a molecule on the surface of the tumorous cell, as well as to a molecule on the surface of normal T-cells, bringing the two cell types into closer proximity and allowing the normal T-cell to destroy the tumorous cell.
Specifically, the BLINCYTO® technology attaches to a cell identified as CD19, which is present on all of the cells of the malignant transformations that cause Ph-R/R B-cell precursor ALL and helps attract the cell into close proximity of the T-cell CD3 with the intent of getting close enough to allow the T-cell to inject toxins that destroy the cancerous cell. According to the applicant, the BLINCYTO® technology is the first, and the only, bi-specific CD19-directed CD3 T-cell engager single-agent immunotherapy approved by the FDA.

BLINCYTO® is administered as a continuous IV infusion delivered at a constant flow rate using an infusion pump. A single cycle of treatment consists of 28 days of continuous infusion, and each treatment cycle is followed by 2 weeks without treatment prior to administering any further treatments. A course of treatment would consist of two phases. Phase 1 consists of initial inductions or treatments intended to achieve remission followed by additional inductions and treatments to maintain consolidation; or treatments given after remission has been achieved to prolong the duration. During Phase 1 of a single treatment course, up to two cycles of BLINCYTO® are administered, and up to three additional cycles are administered during consolidation. The recommended dosage of BLINCYTO® administered during the first cycle of treatment is 9 mcg per day for the first 7 days of treatment. The dosage is then increased to 28 mcg per day for 3 weeks until reaching Phase 2 of the treatment course, all subsequent doses are administered as 28 mcg per day throughout the entire duration of the 28-day treatment period.

With regard to the newness criterion, the BLINCYTO® technology received FDA approval on December 3, 2014, for the treatment of patients diagnosed with Ph-R/R B-cell precursor ALL and the product gained entry onto the U.S. market on December 17, 2014.

After evaluation of the newness, costs, and substantial clinical improvement criteria for new technology add-on payments for BLINCYTO® and consideration of the public comments we received in response to the FY 2016 IPPS/LTCPPPS proposed rule, we approved BLINCYTO® for new technology add-on payments for FY 2016 (80 FR 49449). Cases involving BLINCYTO® that are eligible for new technology add-on payments are identified using one of the following ICD–10–PCS procedure codes: XW04351 (Introduction of Blinatumomab antineoplastic immunotherapy into peripheral vein, percutaneous approach, New Technology Group 1), or XW04351 (Introduction of Blinatumomab antineoplastic immunotherapy into central vein, percutaneous approach, New Technology Group 1).

As discussed in the FY 2016 IPPS/LTCPPPS final rule (80 FR 49449), the applicant recommended that CMS consider and use the cost of the full 28-day inpatient treatment cycle as the expected length of treatment when determining the maximum new technology add-on payment for cases involving the BLINCYTO®, rather than the average cost of lesser number of days used as other variables. For the reasons discussed, we disagreed with the applicant and established the maximum new technology add-on payment amount for a case involving the BLINCYTO® technology for FY 2016 using the weighted average of the cycle 1 and cycle 2 observed treatment length. Specifically, in the Phase II trial, the most recent data available, 92 patients received cycle 1 treatment for an average length of 21.2 days, and 52 patients received cycle 2 treatment for an average length of 10.2 days. The weighted average of cycle 1 and cycle 2 treatment length is 17 days. We noted that a small number of patients also received 3 to 5 treatment cycles. However, based on the data provided, these cases do not appear to be typical at this point and we excluded them from this calculation. We noted that, if we included all treatment cycles in this calculation, the weighted average number of days of treatment is much lower, 10 days. Using the clinical data provided by the applicant, we stated that we believe setting the maximum new technology add-on payment amount for a case involving the BLINCYTO® technology for FY 2016 based on a 17-day length of treatment cycle is representative of historical and current practice. We also stated that, for FY 2017, if new data on length of treatment are available, we would consider any such data in evaluating the maximum new technology add-on payment amount. However, we did not receive any new data from the applicant to evaluate for FY 2017.

In the application, the applicant estimated that the average Medicare beneficiary would require a dosage of 9mcg/day for the first 7 days under the first treatment cycle, followed by a dosage of 28mcg/day for the duration of the treatment cycle, as well as all days included in subsequent cycles. All vials contain 35mcg at a cost of $3,178.57 per vial. The applicant noted that all vials are single-use. Therefore, we determined that cases involving the use of the BLINCYTO® technology would incur an average cost per case of $54,035.69 (1 vial/day × 17 days × $3,178.57/vial). Under §412.88(a)(2), we limit new technology add-on payments to the lesser of 50 percent of the average cost of the technology or 50 percent of the costs in excess of the MS–DRG payment for the case. As a result, the maximum new technology add-on payment amount for a case involving the use of the BLINCYTO® is $27,017.85.

With regard to the newness criterion for BLINCYTO®, we consider the beginning of the newness period to commence when the product gained entry onto the U.S. market on December 17, 2014. As discussed previously in this section, in general, we extend new technology add-on payments for an additional year only if the 3-year anniversary date of the product’s entry onto the U.S. market occurs in the latter half of the upcoming fiscal year.

Because the 3-year anniversary date of the entry of the BLINCYTO® onto the U.S. market will occur in the first half of FY 2018 (December 17, 2017), we are proposing to discontinue new technology add-on payments for this technology for FY 2018. We are inviting public comments on this proposal.

6. FY 2018 Applications for New Technology Add-On Payments

We received nine applications for new technology add-on payments for FY 2018. In accordance with the regulations under §412.87(c), applicants for new technology add-on payments must have received FDA approval or clearance by July 1 of the year prior to the beginning of the fiscal year that the application is being considered. Three applicants withdrew their applications prior to the issuance of this proposed rule. We are addressing the remaining six applications below.

a. Bezlotoxumab (ZINPLAVATM)

Merck & Co., Inc. submitted an application for new technology add-on payments for ZINPLAVATM for FY 2018. ZINPLAVATM is indicated for use in adult patients who are receiving antibacterial drug treatment for a diagnosis of Clostridium difficile infection (CDI) who are at high risk for CDI recurrence. ZINPLAVATM is not indicated for the treatment of the presenting episode of CDI and is not an antibacterial drug.

Clostridium difficile (C-diff) is a disease-causing anaerobic, spore forming bacteria that can affect the gastrointestinal (GI) tract. Some people carry the C-diff bacterium in their intestines, but never develop symptoms
of an infection. The difference between asymptomatic colonization and pathogenicity is caused primarily by the production of an enterotoxin (Toxin A) and/or a cytoxin (Toxin B). The presence of either or both toxins can lead to symptomatic CDI, which is defined as the acute onset of diarrhea with a documented infection with toxigenic C-diff, or the presence of either toxin A or B. The GI tract contains millions of bacteria, commonly referred to as “normal flora” or “good bacteria,” which play a role in protecting the body from infection. Antibiotics can kill these good bacteria and allow the C-diff bacteria to multiply and release toxins that damage the cells lining the intestinal wall, resulting in a CDI. CDI is a leading cause of hospital-associated gastrointestinal illnesses.

Persons at increased risk for CDI include people who are treated with current or recent antibiotic use, people who have encountered current or recent hospitalization, people who are older than 65 years, immunocompromised patients, and people who have recently had a diagnosis of CDI. CDI symptoms include, but are not limited to, diarrhea, abdominal pain, and fever. CDI symptoms range in severity from mild (abdominal discomfort, loose stools) to severe (profuse, watery diarrhea, severe pain, and high fevers). Severe CDI can be life-threatening and, in rare cases, can cause bowel rupture, sepsis and organ failure. CDI is responsible for 14,000 deaths per year in the United States.

C-diff produces two virulent, pro-inflammatory toxins, Toxin A and Toxin B, which target host colonocytes (that is, large intestine endothelial cells) by binding to endothelial cell surface receptors via combined repetitive oligopeptide (CROP) domains. These toxins cause the release of inflammatory cytokines leading to intestinal fluid secretion and intestinal inflammation. The applicant asserted that ZINPLAVA™ targets Toxin B sites within the CROP domain rather than the C-diff organism itself. According to the applicant, by targeting C-diff Toxin B, ZINPLAVA™ neutralizes Toxin B, prevents large intestine endothelial cell inflammation, symptoms associated with CDI, and reduces the recurrence of CDI. ZINPLAVA™ binds to sites within the CROP domain, which prevents Toxin B from binding to the host cell, thereby preventing the inflammation and symptoms associated with CDI. ZINPLAVA™ is used concomitantly with standard of care (SOC) antibiotics. Typical treatment of CDI includes antibiotic therapy using vancomycin, metronidazole, fidaxomicin, or other antibiotics. Alternative therapies include fecal microbiota transplant (FMT) and the use of probiotics.

The primary goal of CDI treatment is resolving the infection. Antibacterial drug treatment remains the cornerstone of treatment of CDI. However, this treatment option alone may not be adequate for patients diagnosed with recurrent CDI. A major concern with respect to a CDI is that even when treatment with an antibacterial drug of a primary infection is successful, generally, 25 percent to 30 percent of patients experience a recurrence of the infection within days or weeks of the presenting episode’s symptom resolution. The risk of recurrence increases to 65 percent with subsequent CDI episodes. Disease recurrence results from continued disruption of the intestinal microbiota by SOC CDI antibiotics (or use of other antibiotics used to treat non-gastrointestinal conditions), combined with persistence of resistant C-diff spores (relapse) or acquisition of new spores from the environment (reinfection).

Antibacterial drug use may inhibit the intestinal microbiota from reestablishing itself, allowing C-diff spores potentially to germinate and colonize the intestines when the antibacterial drug is discontinued. If regrowth of C-diff overtakes the reestablishment of the intestinal microbiota, then spore germination and toxin production from vegetative C-diff may restart the cycle of CDI and the need for subsequent treatment. These challenges highlight the need for nonantibiotic therapies. ZINPLAVA™ targets Toxin B rather than the C-diff bacteria itself. According to the applicant, unlike antibacterial drugs, ZINPLAVA™ is a human monoclonal antibody and does not affect the microbiota. According to the applicant, ZINPLAVA™ neutralizes C-diff Toxin B and reduces recurrence of CDI. ZINPLAVA™ is given concomitantly during the course of SOC antibacterial treatment of a CDI.

With respect to the newness criterion, ZINPLAVA™ received FDA approval on October 21, 2016, for reduction of recurrence of CDI in patients receiving antibacterial drug treatment for CDI and who are at high risk of CDI recurrence. ZINPLAVA™ is anticipated to be commercially available as of February 2017. We note that the applicant anticipates submitting a request for a unique ICD–10–PCS code for the administration of ZINPLAVA™. Currently, cases that receive ICD–10–CM request to differentiate CDI recurrence. If approved, the codes will become effective on October 1, 2017 (FY 2018).

As discussed above, if a technology meets all three of the substantial similarity criteria, it would be considered substantially similar to an existing technology and would not be considered “new” for purposes of new technology add-on payments.

With regard to the first criterion, whether a product uses the same or a similar mechanism of action to achieve a therapeutic outcome, according to the applicant, ZINPLAVA™ is a human monoclonal antibody with an innovative mechanism of action. The applicant asserted that ZINPLAVA™ is a novel treatment, with a unique mechanism of action relative to SOC CDI antibiotics that target C-diff. The applicant explained that ZINPLAVA™ is the first human monoclonal antibody that targets and neutralizes C-diff/Toxin B because the technology specifically binds to and neutralizes C-diff/Toxin B (which is an exotoxin that contributes to intestinal tissue damage and immune system effects that underlie the symptoms of CDI) and inhibits binding of the toxin to mammalian cells. The applicant further asserted that the administration of ZINPLAVA™, in addition to standard of care antibacterial drug treatment, reduces CDI recurrence by providing passive immunity against Toxin B resulting from persistent or newly acquired C-diff spores. According to the applicant, ZINPLAVA™ is the only FDA-approved treatment indicated for reducing CDI recurrence as an adjunctive therapy in adult patients who are receiving antibacterial drug treatment for CDI and who are at high risk for CDI recurrence.

With respect to the second criterion, whether a product is assigned to the same or a different MS–DRG, the applicant maintained that patients who may be eligible to receive treatment using ZINPLAVA™ could be in an acute-care hospital setting for a wide variety of reasons and may develop a secondary CDI as a hospital-acquired infection and, therefore, cases representing patients that may be eligible for treatment using ZINPLAVA™ may map to a wide range of MS–DRGs. ZINPLAVA™ is indicated for patients receiving SOC treatment for CDI and who are at a high risk for CDI recurrence. In order to identify the range of MS–DRGs for which cases representing patients that may be eligible for treatment using ZINPLAVA™ may map to, the applicant identified all MS–DRGs containing cases that SOC antibacterials for CDI patients presenting with CDI as a primary or secondary diagnosis. The applicant used
FY 2015 MedPAR data to map the identified cases to 543 MS–DRGs, with 12 MS–DRGs accounting for approximately 40 percent of all cases. The applicant segmented these cases based on age because patients 65 years and older are at higher risk for CDI recurrence. Based on the FY 2015 MedPAR data, MS–DRG distribution was found to be similar, irrespective of CDI status (primary or secondary), for patients over 65 years of age and those under 65 years of age. The top 7 MS–DRGs across both age groups account for nearly 54 percent (over 65 years of age) and 49 percent (under 65 years of age). The applicant further segmented these cases to determine if status of CDI as a primary or secondary diagnosis influenced MS–DRG mapping. Regardless of age, when CDI is the primary diagnosis, approximately 98 percent of patient cases map to the same 3 MS–DRGs: MS–DRG 371 (Major Gastrointestinal Disorders and Peritoneal Infections with MCC); MS–DRG 372 (Major Gastrointestinal Disorders and Peritoneal Infections with CC); and MS–DRG 373 (Major Gastrointestinal Disorders and Peritoneal Infections without CC/MCC), respectively. Potential cases representing patients who may be eligible for treatment with ZINPLAVA™ would be assigned to the same MS–DRGs as cases representing patients who receive SOC treatment for a diagnosis of CDI.

With respect to the third criterion, whether the new use of the technology involves the treatment of the same or similar type of disease and the same or similar patient population, according to the applicant, ZINPLAVA™ is administered concomitantly or as adjunctive therapy with SOC antibacterial treatment for recurrent CDI. The applicant stated that ZINPLAVA™ is indicated to reduce recurrence of CDI in adult patients at high risk of CDI recurrence who are receiving antibacterial drug treatment for CDI. According to the applicant, the addition of ZINPLAVA™ to SOC antibacterial treatment reduces CDI recurrence by providing passive immunity against Toxin B resulting from persistent or newly acquired C. difficile spores. ZINPLAVA™ is used to treat the same or similar type of disease (recurrent CDI) and a similar patient population receiving SOC therapy for the treatment of recurrent CDI.

Based on the applicant’s statements presented above, because ZINPLAVA™ has a unique mechanism of action, we do not believe that the technology is substantially similar to existing technologies and, therefore, meets the newness criterion. We are inviting public comments on whether ZINPLAVA™ meets the newness criterion.

With regard to the cost criterion, the applicant conducted the following analysis to demonstrate that the technology meets the cost criterion. In order to identify the range of MS–DRGs that cases representing potential patients who may be eligible for treatment using ZINPLAVA™ may map to, the applicant identified all MS–DRGs for patients diagnosed with CDI as a primary or secondary diagnosis. Specifically, the applicant searched the FY 2015 MedPAR file for claims that included target patients over 65 years of age and identified cases reporting diagnoses of CDI by ICD–9–CM diagnosis code 008.45 (Intestinal infection due to Clostridium difficile) as a primary or secondary diagnosis. This resulted in 139,135 cases across 543 MS–DRGs, with approximately 40 percent of all cases mapping to the following 12 MS–DRGs: MS–DRG 177 (Respiratory Infections and Inflammations with MCC); MS–DRG 193 (Simple Pneumonia and Pleurisy with MCC); MS–DRG 291 (Heart Failure and Shock with MCC); MS–DRGs 371, 372, and 373 (Major Gastrointestinal Disorders and Peritoneal Infections with MCC, with CC, and without CC/MCC, respectively); MS–DRGs 682 and 683 (Renal Failure with MCC and with CC, respectively); MS–DRG 853 (Infectious and Parasitic Diseases with O.R. Procedure with MCC); MS–DRGs 870, 871, and 872 (Septicemia or Severe Sepsis with Mechanical Ventilation >96 Hours, with MCC, and without MCC, respectively).

Using the 139,135 identified cases, the average unstandardized case-weighted charge per case was $80,677. The applicant then standardized the charges. The applicant did not remove charges for the current treatment because, as discussed above, ZINPLAVA™ will be used concomitantly with SOC antibacterial treatments for the treatment of CDI as an additive, or adjunctive treatment option, to reduce the recurrence of CDI infection. The applicant then applied the 2-year inflation factor of 1.098446 from the FY 2017 IPPS/LTCH final rule (81 FR 57286) to inflate the charges from FY 2015 to FY 2017. The applicant noted that the anticipated price for ZINPLAVA™ has yet to be determined; therefore, no charges for ZINPLAVA™ were added in the analysis. Based on the FY 2017 IPPS/LTCH FFPS Table 10 thresholds, the average case-weighted threshold amount was $56,871. The inflated average case-weighted standardized charge per case was $78,929. Because the inflated average case-weighted standardized charge per case exceeds the average case-weighted threshold amount, the applicant maintained that the technology meets the cost criterion. The applicant noted that the inflated average case-weighted standardized charge per case exceeds the average case-weighted threshold amount without the average per patient cost of the technology. As such, the applicant anticipated that the inclusion of the cost of ZINPLAVA™, at any price point, will further increase charges above the average case-weighted threshold amount. We are inviting public comments on whether ZINPLAVA™ meets the cost criterion.

With respect to the substantial clinical improvement criterion, the applicant asserted that the addition of ZINPLAVA™ to SOC antibacterial drug treatment reduces CDI recurrence because it provides passive immunity against Toxin B resulting from persistent or newly acquired C. difficile spores.

The applicant conducted two Phase III studies, MODIFY I and MODIFY II. The primary endpoint of the studies was recurrent CDI within 12 weeks after completion of treatment with ZINPLAVA™. The first study design initially included actoxumab, an antitoxin A monoclonal antibody treatment arm that was later discontinued due to a high failure rate and increase in mortality compared to other treatment arms.3 Clinical data on ZINPLAVA™ is provided exclusively from the FDA briefing document available on the FDA Web site at: http://www.fda.gov/AdvisoryCommittees/CommitteesMeetingMaterials/Drugs/Anti-InfectiveDrugsAdvisoryCommittee. Information is also provided in the package insert by the manufacturer, Merck & Company, Inc. The FDA briefing provided data on the safety and efficacy of ZINPLAVA™. The FDA considered sustained clinical responses defined as clinical cure of the initial CDI episode and the absence of CDI recurrence as an appropriate endpoint to assess the efficacy of ZINPLAVA™ in the prevention of CDI recurrences. In MODIFY I trial, the clinical cure rate of the presenting CDI episode was lower in the ZINPLAVA™ arm as compared to the placebo arm, whereas in MODIFY II trial the clinical cure rate was lower in the placebo arm as compared to the ZINPLAVA™ arm. Additional analyses showed that, by 3 Wilcox MH et al. Bezlotoxumab for Prevention of Recurrent Clostridium difficile Infection. N Engl J Med. 2017 Jun 26;376(4):305–317.
weeks post study drug infusion, the clinical cure rates of the presenting CDI episode were similar between treatment arms.

In MODIFY I, the rate of sustained clinical response was numerically in favor of ZINPLAVAM (60.1 percent) compared to placebo (55.2 percent) with an adjusted difference and 95 percent CI of 4.8 percent (−2.1 percent; 11.7 percent). In MODIFY II, the proportion of subjects with sustained clinical response in the ZINPLAVAM arm (66.8 percent) was also higher than in the placebo arm (52.1 percent) with an adjusted difference of 14.6 percent and 95 percent CI (7.8 percent; 21.4 percent). The treatment did not significantly decrease mortality. Recurrence rates, including CDI-related hospital readmission rates, reportedly were between 10 and 25 percent. No clinically meaningful differences in the exposure of bezlotoxumab were found between patients 65 years of age and older and patients under 65 years of age. In the Phase III trials, the safety profile of ZINPLAVAM was similar overall to that of placebo. However, heart failure was reported more commonly in the two Phase III clinical trials of ZINPLAVAM-treated patients compared to placebo-treated patients. These adverse reactions occurred primarily in patients with underlying congestive heart failure (CHF). In patients with a history of CHF, 12.7 percent (15/118) of ZINPLAVAM-treated patients and 4.8 percent (5/104) of placebo-treated patients had the serious adverse reaction of heart failure during the 12-week study period. In addition, in patients with a history of CHF, there were more deaths in ZINPLAVAM-treated patients (19.5 percent (23/118)) than in placebo-treated patients (12.5 percent (13/104)) during the 12-week study period. We are concerned regarding the safety of ZINPLAVAM in patients diagnosed with CHF. In regard to safety, data from the MODIFY I and MODIFY II studies suggest few adverse events associated with ZINPLAVAM, with no significant differences in the number of serious adverse events, deaths or discontinuations of study drug that occurred between the ZINPLAVAM and the placebo groups. However, both the ZINPLAVAM and the ZINPLAVAM plus actoxumab treatment groups experienced more episodes of cardiac failure (defined as acute or chronic cardiac failure) than compared to the placebo group (2.2 percent versus 1 percent). We are unsure if the cardiac failure reported in the studies may be the result of a higher number of baseline patients with heart failure in the treatment arms or the result of an adverse effect to ZINPLAVAM.

Therefore, we are concerned with regard to the adverse event of cardiac failure of ZINPLAVAM.

We are inviting public comments on whether ZINPLAVAM meets the substantial clinical improvement criterion.

We did not receive any written public comments in response to the New Technology Town Hall meeting notice regarding the application of ZINPLAVAM for new technology add-on payments.

b. EDWARDS INTUITY EliteTM Valve System (INTUITY) and LivaNova Perceval Valve (Perceval)

Two manufacturers, Edwards Lifesciences and LivaNova, submitted applications for new technology add-on payments for FY 2018 for the INTUITY EliteTM Valve System (INTUITY) and the Perceval Valve (Perceval), respectively. Both of these technologies are prosthetic aortic valves inserted using surgical aortic valve replacement (AVR). We note that, while Edwards Lifesciences submitted an application for new technology add-on payments for FY 2017 for the INTUITY valve, FDA approval was not received by July 1, 2016, and, therefore, the device was not eligible for consideration for new technology add-on payments for FY 2017.

Aortic valvular disease is relatively common, primarily manifested by aortic stenosis. Most aortic stenosis is due to calcification of the valve, either on a normal tri-leaflet valve or on a congenitally bicuspid valve. The resistance to outflow of blood is progressive over time, and as the size of the aortic orifice narrows, the heart must generate increasingly elevated pressures to maintain blood flow. Symptoms such as angina, heart failure, and syncope eventually develop, and portend a very serious prognosis. There is no effective medical therapy for aortic stenosis, so the diseased valve must be replaced or, less commonly, repaired.

The INTUITY valve incorporates the expansion feature of a catheter implanted valve, but is designed to be placed during cardiac surgery. The manufacturer explained that the INTUITY valve requires fewer stitches to hold the device in place because of the balloon expanded design and, therefore, can be inserted more quickly than a standard valve, and also facilitates minimally invasive cardiac surgery; that is, use of a smaller incision to allow faster recovery. The manufacturer of the Perceval valve indicated that the device is comprised of: (1) A bovine pericardial aortic bioprosthetic valve; (2) a balloon expandable stainless steel frame; and (3) a textured sealing cloth. The manufacturer of the Perceval valve indicated that the Perceval valve device is comprised of: (1) Sizers used to determine the correct size of the prosthesis; (2) a dual holder used for positioning and deployment (available in two models, one for sternal approaches and one for MIS); (3) a “smart clip” to assist during assembly of the valve on the dual holder to prevent release during positioning; (4) a dual collapser used to evenly reduce the diameter of the prosthesis allowing it to mount onto the holder prior to implantation; (5) a dual collapser base used to allow proper positioning; and (6) a postdilation catheter used for in situ dilation of the prosthesis after implantation (available in two models, one for sternal approaches and one for MIS). According to both applicants, the INTUITY valve and the Perceval valve are the first sutureless, rapid deployment aortic valves that can be used for the treatment of patients who are candidates for surgical AVR. The applicants indicated that the two new device innovations facilitate MIS approaches through: (1) The device rapid deployment mechanisms; and (2) the design of the prosthetic valve that allows for markedly fewer to no sutures to securely fasten the prosthetic valve to the aortic orifice. The applicants explained that both of these aspects of their devices are credited with the reduction of operating time.

As noted, according to both applicants, the INTUITY valve and the Perceval valve are the first sutureless, rapid deployment aortic valves that can be used for the treatment of patients who are candidates for surgical AVR. Because potential cases representing patients who are eligible for treatment using the INTUITY and the Perceval aortic valve devices would group to the same MS–DRGs, and we believe that these devices are intended to treat the same or similar disease in the same or similar patient population, and are purposed to achieve the same therapeutic outcome using the same or similar mechanism of action, we believe these two devices are substantially similar to each other and that it is appropriate to evaluate both technologies as one application for new technology add-on payments under the IPPS.

With respect to the newness criterion, the INTUITY valve received FDA approval on August 12, 2016, and was commercially available in the U.S. market on August 19, 2016. The Perceval valve received FDA approval
on January 8, 2016, and was commercially available on the U.S. market on February 29, 2016. We believe that, in accordance with our policy, it is appropriate to use the earliest market availability date submitted as the beginning of the newness period. Therefore, based on our policy, with regard to both devices, if the technologies are approved for new technology add-on payments, we believe that the beginning of the newness period would be February 29, 2016. In addition, both applicants indicated that ICD–10–PCS code X2RF032 (Replacement of Aortic Valve using Zooplastic Tissue, Rapid Deployment Technique, Open Approach, New Technology Group 2) would identify procedures involving the use of the devices when surgically implanted.

We previously stated that, because we believe these two devices are substantially similar to each other, we believe it is appropriate to evaluate both technologies as one application for new technology add-on payment under the IPPS. The applicants submitted separate cost and clinical data, and we reviewed and discussed each set of data separately. However, we intend to make one determination regarding new technology add-on payments that will apply to both devices. We believe that this is consistent with our policy statements in the past regarding substantial similarity. Specifically, we have noted that approval of new technology add-on payments would extend to all technologies that are substantially similar (66 FR 69615), and we believe that continuing our current practice of extending new technology add-on payments without a further application from the manufacturer of the competing product, or a specific finding on cost and clinical improvement if we make a finding of substantial similarity among two products is the better policy because we avoid—

- Creating manufacturer-specific codes for substantially similar products;
- Requiring different manufacturers of substantially similar products to submit separate new technology applications;
- Having to compare the merits of competing technologies on the basis of substantial clinical improvement; and
- Bestowing an advantage to the first applicant representing a particular new technology to receive approval (70 FR 47351).

If these substantially similar technologies were submitted for review in different (and subsequent) years, rather than the same year, we would evaluate and make a determination on the first application and apply that same determination to the second application. However, because the technologies have been submitted for review in the same year, we believe that it is appropriate to consider both sets of cost data and clinical data in making a determination and we do not believe that it is possible to choose one set of data over another set of data in an objective manner.

As stated above, we believe that the INTUITY valve and the Perceval valve are substantially similar to each other for purposes of analyzing these two applications as one application. We also need to determine whether the INTUITY valve and the Perceval valve are substantially similar to existing technologies prior to their approval by the FDA and their release on the market. As discussed earlier, if a technology meets all three of the substantial similarity criteria, it would be considered substantially similar to an existing technology and would not be considered “new” for purposes of new technology add-on payments.

With respect to the first criterion, whether a product uses the same or a similar mechanism of action to achieve a therapeutic outcome, the applicant for the INTUITY valve asserted that its unique design, which utilizes features that were not previously included in conventional aortic valves, constitutes a new mechanism of action. The deployment mechanism allows for rapid delivery. The expandable frame can reshape the native valve’s orifice, creating a larger and more efficiently shaped effective orifice area. In addition, the expandable skirt allows for structural differentiation upon fixation of the valve requiring 3 permanent, guiding sutures rather than the 12 to 18 permanent sutures used to fasten standard prosthetic aortic valves. The applicant for the Perceval valve described the Perceval valve’s mechanism of action as including: (a) No permanent sutures; (b) a dedicated delivery system that increases the surgeon’s visibility; (c) an enabler of minimally invasive approach; (d) a complexity reduction and reproducibility of the procedure; and (e) a unique device assembly and delivery systems.

With respect to the second and third criteria, whether a product is assigned to the same or a different MS–DRG and whether the new use of the technology involves the treatment of the same or similar type of disease and the same or similar patient population, the applicant for the INTUITY valve indicated that the technology is used in the treatment of the same patient population and potential cases representing patients that may be eligible for treatment using the INTUITY valve would be assigned to the same MS–DRGs as cases involving the use of other prosthetic aortic valves (that is, MS–DRGs 216 (Cardiac Valve & Other Major Cardiothoracic Procedures with Cardiac Catheterization with MCC), 217 (Cardiac Valve & Other Major Cardiothoracic Procedures with Cardiac Catheterization with CC), 218 (Cardiac Valve & Other Major Cardiothoracic Procedures with Cardiac Catheterization without CC/MCC), 219 (Cardiac Valve & Other Major Cardiothoracic Procedures without Cardiac Catheterization with CC), and 221 (Cardiac Valve & Other Major Cardiothoracic Procedures without Cardiac Catheterization without CC/MCC). The applicant for the Perceval valve also indicated that the Perceval valve device is used in the treatment of the same patient population and potential cases representing patients that may be eligible for treatment using the technology would be assigned to the same MS–DRGs (MS–DRGs 216 through 221) as cases involving the use of other prosthetic aortic valves.

After considering the materials included with both applications, we remain concerned as to whether the mechanism of action described by the applicants represents an improvement to an existing surgical technique and technology or a new technology. While the INTUITY and Perceval valves address some of the challenges posed by implantation of existing valves, including improving the visibility of the orifice and the physiological function of the valves, we do not believe that their mechanisms of action are fundamentally different from that of other aortic valves. As one of the applicants stated in its application, the goal of the prosthetic aortic valve is to mimic the native valve that it has replaced via the incorporation of three leaflets that open and close in response to pressure gradients developed during the cardiac cycle. We believe that the INTUITY and Perceval valves are similar to other prosthetic aortic valves used to treat the same or similar diagnoses.

We are inviting public comments on whether the mechanisms of action of the sutureless, rapid deployment of the INTUITY and Perceval valves differs from the mechanism of action of standard AVR valves and whether the technologies meet the newness criterion.

As stated above, each applicant submitted separate analyses regarding the cost criterion for each of their devices, and both applicants maintained...
that their device meets the cost criterion. We summarize each analysis below.

With regard to the cost criterion, the INTUITY valve’s applicant researched the FY 2015 MedPAR claims data file to identify cases representing patients who may be potential recipients of treatment using the INTUITY valve. The applicant identified claims that reported an ICD–9–CM diagnosis code of 424.1 (Aortic valve disorder), in combination with an ICD–9–CM procedure code of 35.21 (Replacement of aortic valve with tissue) or 35.22 (Open and other replacement of aortic valve). The applicant also identified cases with or without a coronary artery bypass graft (CABG) using the ICD–9–CM procedure codes in the table below.

<table>
<thead>
<tr>
<th>ICD–9–CM code</th>
<th>Code description</th>
</tr>
</thead>
<tbody>
<tr>
<td>36.10</td>
<td>Aorto coronary bypass for heart revascularization, not otherwise specified.</td>
</tr>
<tr>
<td>36.11</td>
<td>(Aorto)coronary bypass of one coronary artery.</td>
</tr>
<tr>
<td>36.12</td>
<td>(Aorto)coronary bypass of two coronary arteries.</td>
</tr>
<tr>
<td>36.13</td>
<td>(Aorto)coronary bypass of three coronary arteries.</td>
</tr>
<tr>
<td>36.14</td>
<td>(Aorto)coronary bypass of four or more coronary arteries.</td>
</tr>
<tr>
<td>36.15</td>
<td>Single internal mammary-coronary artery bypass.</td>
</tr>
<tr>
<td>36.16</td>
<td>Double internal mammary-coronary artery bypass.</td>
</tr>
<tr>
<td>36.17</td>
<td>Abdominal-coronary artery bypass.</td>
</tr>
</tbody>
</table>

The applicant identified a total of 25,173 cases that mapped to MS–DRGs 216 through 221. Of these cases, the applicant identified 10,251 CABG cases and 14,922 non-CABG cases. According to the applicant, patients that undergo a procedure without need of a concomitant CABG are more likely to receive treatment with the INTUITY valve than patients in need of a concomitant CABG. Therefore, the applicant weighted the non-CABG cases at 90 percent of total cases and the CABG cases at 10 percent of total cases under each of the six MS–DRGs. The final case count is a weighted average of 14,455 cases.

The applicant calculated an average unstandardized charge per case of $192,506 for all cases. The applicant then removed 100 percent of the charges for pacemakers, investigational devices, and other implants that would not be required for patients receiving treatment using the INTUITY valve. The applicant standardized the charges and then applied an inflation factor of 1.098446, which is the 2-year inflation factor in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57286), to update the charges from FY 2015 to FY 2017. The applicant calculated the average expected charge for the INTUITY valve based on the current list price of the device.

Although the applicant submitted data related to the cost of the INTUITY valve, the applicant noted that the cost of the device is proprietary information. To add charges for the device, the applicant assumed a hospital mark-up of approximately 3.00 percent, based on the current average CCR for implantable devices (0.331) as reported in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56876). Based on the FY 2017 IPPS/LTCH PPS Table 10 thresholds, the average case-weighted threshold amount was $170,321. The applicant computed an inflated average case-weighted standardized charge per case of $194,291, which is $23,970 above the average case-weighted threshold amount. Because the inflated average case-weighted standardized charge per case exceeds the average case-weighted threshold amount, the applicant maintained that the technology meets the cost criterion.

We thank the applicant for the analysis above. However, we would like more information from the applicant regarding how it decided upon which cases to include in the sensitivity analysis, as well as further details about how and on what basis the applicant weighted CABG and non-CABG cases. We are inviting public comments on whether the INTUITY valve meets the cost criterion.

With regard to substantial clinical improvement for the INTUITY valve, the applicant asserted that several aspects of the valve system represent a substantial clinical improvement over existing technologies. The applicant believed that the flexible deployment arm allows improved surgical access and visualization, making the surgery less challenging for the surgeon, improving the likelihood that the surgeon can use a minimally invasive approach. According to the applicant, the assembly of the device only allows the correct valve size to be fitted, which ensures that the valve does not slip or
migrate, which prevents paravalvular leaks and patient prostatic mismatch. The applicant indicated that the device improves clinical outcomes for patients undergoing minimally invasive AVR and full-sternotomy AVR. The applicant stated that the rapid deployment technology enables reduced operative time, specifically cross-clamp time, thereby reducing the period of myocardial ischemia. In addition, the applicant indicated that the device offers a reduction in operative time for full-sternotomy AVR. The applicant noted that several clinical trials document significant patient outcome and utilization improvements, including improved patient satisfaction, faster return to normal activity, decreased post-operative pain, reduced mortality and decreased complications, including need for reoperation due to bleeding, reduced recovery time, reduced length of stay (both ICU and overall), more access to minimally invasive surgery, and improved hemodynamics.

The INTUITY valve has been tested clinically in several trials. In the TRITON trial (Kocher et al., 20134), 287 patients diagnosed with aortic stenosis underwent surgery in 1 of 6 European centers. The first 149 patients received the first generation Model 8300A valve, and the next 138 patients received the second generation Model 8300AB. The average age of the patients was 75.7 years. Early, 30-day mortality was 1.7 percent (5/287), the post-op valve gradient was low, and 75 percent of the patients improved functionally. A total of 4 valves were explanted in the final 30 days due to bleeding, and 3 were explanted later for paravalvular leak, endocarditis, and aortic root aneurysms. Follow-up extended to 3 years (mean 1.8 years).

Implantation of the INTUITY valve using minimally invasive surgery was compared with conventional aortic valve replacement via full sternotomy in the CADENCE–MIS randomized trial (Borger et al., 201525) of 100 patients treated in 1 of 5 centers in Germany. The authors found no significant difference in 30-day mortality, the need for pacemaker implantation, significant paravalvular regurgitation, and quality of life scores at 3 months. Aortic cross-


clamp time was significantly reduced from 54.0 to 41.3 minutes (p < 0.0001), and cardiopulmonary bypass time was reduced from 74.4 to 68.8 minutes (p = 0.21). Early clinical outcomes were similar: No significant differences in mortality, reoperation, or other clinical outcomes. The aortic valve gradient was significantly lower in the MIS group: 8.5 versus 10.3 mmHg.

The TRANSFORM trial (Barnhart et al. 20179) was a single-arm, non-randomized, multicenter trial, in which 839 patients underwent rapid deployment AVR surgery. The average age of the patients was 75.5 years. The mean cross-clamp time and cardiopulmonary bypass times for full sternotomy were 49.3 ± 26.9 min and 69.2 ± 34.7 min, respectively, and for MIS, 63.1 ± 25.4 min and 84.6 ± 33.5 min, respectively. The authors compared these times to STS database comparators: For full sternotomy, 76.3 minutes and 104.2 minutes, respectively, and for MIS, 82.9 minutes and 111.4 minutes, respectively. All cause early mortality was 0.8 percent, mean EOAn at 1 year was 1.7 cm2; mean gradient, 10.3 mmHg; and moderate and severe PVL, 1.2 percent and 0.4 percent, respectively. The authors indicated that the INTUITY valve “...may lead to a relative reduction in aortic cross-clamp time and cardiopulmonary bypass time” and “may confer benefits to patients, such as decreased mortality and morbidity.” The authors noted the possibility of potential bias resulting from the level of experience of the study surgeons relative to typical cardiac surgeons. In addition, long-term follow-up is not available, and study comparators from the Society of Thoracic Surgeons (STS) database were not matched.

In the FY 2017 IPPS/LTCH PPS proposed rule (81 FR 25057), after reviewing the studies provided by the applicant with its application for FY 2017, we expressed some specific concerns. We indicated that we were concerned that the INTUITY valve does not have sufficient advantages over alternative valve retained valves to constitute a substantial clinical improvement. We noted that, while some of the studies included with the application demonstrate reduced aortic cross-clamp time, conventional aortic valve replacement was used in the comparison group. Therefore, it is unclear whether the reduced aortic cross-clamp time is associated with the use of the INTUITY valve or as a result of the MIS surgery in general.

In response to these concerns, the INTUITY valve’s applicant stated that the INTUITY valve is associated with significant clinical benefits outside of the benefits achieved by use of an MIS approach. The applicant referenced the sub-study of the TRANSFORM trial, which compared the MISAVR with the INTUITY valve to MISAVR with a conventional valve, stating that the results indicated reduced cross-clamp time and other benefits that are not simply a function of the MIS approach. The applicant also referenced trials that indicated that the INTUITY valve had excellent hemodynamic performance (Haverich et al.,7 Borger et al.,8 Barnhart et al.),9 one of which found a significant improvement in functional status (Haverich et al.).

After considering the studies provided by the INTUITY valve applicant, we are concerned about the possibility of potential bias resulting from the level of experience of the study surgeons relative to typical cardiac surgeons, as well as the lack of long-term follow-up in these studies.

With regard to substantial clinical improvement for the Perceval valve, the applicant submitted several studies examining the Perceval valve. The following discussion summarizes some of these studies.

Pollari and colleagues10 (2014) utilized a propensity score analysis to examine 82 matched pairs as part of a larger trial that included 566 patients treated with bioprosthetic aortic valve replacement, 166 of which received treatment using the Perceval sutureless valve and 400 of which received treatment using a stented valve. Aortic cross-clamp, cardiopulmonary bypass, and operation times were significantly shorter in the group that received treatment using the Perceval sutureless

valve. The Perceval sutureless group also had shorter ICU stays, hospital stays, and intubation times, and lower incidence of postoperative atrial fibrillation and respiratory insufficiency. The authors noted that, despite the promising preliminary results, longer follow-up is warranted before drawing definite conclusions.

In a nonrandomized trial of 100 patients in a German hospital, Santarpino and colleagues 11 (2013) found that procedures completed using the Perceval valve were associated with significantly shorter cross-clamp and cardiopulmonary bypass times (40 ± 13.8 and 69 ± 19.1 versus 66 ± 20.4 and 105 ± 34.8) relative to conventional stented bioprosthetic valves, as well as less frequent use of blood transfusions, shorter ICU stays and shorter use of intubation. In contrast, Gilmanov and colleagues 12 (2013) found that a MIS approach resulted in improved outcomes, albeit longer aortic cross-clamp times. A meta-analysis by Hurley and colleagues 13 (2015) found reduced cross-clamp and cardiopulmonary bypass times, but found a significantly higher permanent pacemaker rate with the use of Perceval sutureless valves.

A study conducted by Dalen and colleagues 14 (2015) used propensity score matching to examine early postoperative outcomes and 2-year survival between 171 pairs of patients who underwent mininisterectomy using the Perceval device or a full sternotomy with stented prosthesis. There were no differences in 30-day mortality or 2-year survival between the groups. The aortic cross-clamp time and cardiopulmonary bypass time were shorter, and there were fewer blood transfusions in the group that received treatment using the Perceval device. However, this group was also at higher risk for post-operative permanent pacemaker implantation.

After reviewing the publications submitted by the applicant, we are concerned that the lack of randomization and blinded investigators may have influenced the outcomes in many of the studies provided. For example, in the discussion following Santarpino et al.’s 2013 study, one of the participants suggested that medical decision-making regarding ventilation times, ICU times, and blood transfusions may be affected by the knowledge of investigators as to which valve the patient received treatment using. Also, as indicated above with respect to the INTUITY valve, the experience of the surgeons in these studies may be confounding factors that may have influenced the length of surgical procedures and/or surgical outcomes. We are inviting public comments on whether rapid deployment valves, specifically the INTUITY and Perceval valves, meet the substantial clinical improvement criterion.

We did not receive any written public comments regarding the INTUITY and Perceval valves in response to the New Technology Town Hall meeting notice.

c. Ustekinumab (Stelara®)

Janssen Biotech submitted an application for a new technology add-on payments for the Stelara® induction therapy for FY 2018. Stelara® received FDA approval as an intravenous (IV) infusion treatment of Crohn’s disease (CD) on September 23, 2016, which added a new indication for the use of Stelara® and route of administration for this monoclonal antibody. IV infusion of Stelara® is indicated for the treatment of adult patients (18 years and older) diagnosed with moderately to severely active CD who have: (1) Failed or were intolerant to treatment using immunomodulators or corticosteroids, but never failed a tumor necrosis factor (TNF) blocker; or (2) Failed or were intolerant to treatment using one or more TNF blockers. Stelara® for IV infusion has only one purpose, induction therapy. Stelara® must be administered intravenously by a health care professional in either an inpatient hospital setting or an outpatient hospital setting.

Stelara® for IV infusion is packaged in single 130mg vials. Induction therapy consists of a single IV infusion dose using the following weight-based dosing regimen: Patients weighing less than (<) 55kg are administered 260mg of Stelara® (2 vials); patients weighing more than (>) 55kg, but less than (<) 85kg are administered 390mg of Stelara® (3 vials); and patients weighing more than (> ) 85kg are administered 520mg of Stelara® (4 vials). An average dose of Stelara® administered through IV infusion is 390mg (3 vials). Maintenance doses of Stelara® are administered at 90mg, subcutaneously, at 8-week intervals and may occur in the outpatient hospital setting.

CD is an inflammatory bowel disease of unknown etiology, characterized by transmural inflammation of the gastrointestinal (GI) tract. Symptoms of CD may include fatigue, prolonged diarrhea with or without bleeding, abdominal pain, weight loss and fever. CD can affect any part of the GI tract including the mouth, esophagus, stomach, small intestine, and large intestine.

Conventional pharmacologic treatments of CD include antibiotics, mesalamines, corticosteroids, immunomodulators, tumor necrosis alpha (TNFα) inhibitors, and anti-integrin agents. Surgery may be necessary for some patients diagnosed with CD in which conventional therapies have failed. The applicant asserted that use of Stelara® offers an alternative to conventional pharmacologic treatments, and has been shown to be successful in the treatment of patients who have failed treatment using the conventional agents currently being used for a diagnosis of CD, including TNFα inhibitors.

Although the precise cause of CD is unknown, the environment, genetics, and the patient’s immune system are thought to play a role in this form of inflammatory bowel disease (IBD). Conventional pharmacologic therapy is directed against many different inflammatory mediators that produce inflammation and ultimately lead to gastrointestinal damage. The applicant asserted that it is of paramount importance to have a variety of pharmacologic agents that can address the proper inflammatory mediator for a particular patient. The applicant also asserted that, while the currently available anti-inflammatory agents used in the treatment of a diagnosis of CD are excellent medications, these agents do not successfully treat all patients diagnosed with CD, nor do they reliably sustain disease remission once a response has been achieved. The applicant believed that the use of Stelara® offers an alternative to currently available treatment options.

With regard to the newness criterion, Stelara® is not a newly formulated drug. Stelara®, administered subcutaneously, received FDA approval in 2009 (September 25, 2009) for the treatment of moderate to severe plaque psoriasis and psoriatic arthritis in adults. Its IV use for the treatment of patients diagnosed with CD was approved by the FDA in 2016 (September 23, 2016). With regard to the new use of an existing technology, in the September 1, 2001 final rule (66 FR 46915), we stated that if the new use of an existing technology was for treating patients not expected to...
be assigned to the same MS–DRGs as the patients receiving the existing technology, it may be considered for approval, but it must also meet the cost and substantial clinical improvement criteria in order to qualify for the new technology add-on payment. We do not believe that potential cases representing patients that may be eligible for treatment with the new use of the Stelara® for IV treatment of a diagnosis of CD would be assigned to the same MS–DRGs as cases treated using the prior indications.

As discussed above, if a technology meets all three of the substantial similarity criteria, it would be considered substantially similar to an existing technology and would not be considered “new” for purposes of new technology add-on payments.

With regard to the first criterion, whether a product uses the same or a similar mechanism of action to achieve a therapeutic outcome, we are concerned that Stelara®’s mechanism of action may not appear to differ from the mechanism of action of other monoclonal antibodies, which also target unique gastrointestinal-selective cytokines. The applicant believed that the Stelara® uses a different mechanism of action than other medications currently available for the treatment of patients diagnosed with CD. However, we believe that the mechanism of action for the new use of the Stelara® may be similar to the mechanism of action of other cytokine-selective monoclonal antibodies that disrupt cytokine-mediated signals crucial to the inflammatory process in patients diagnosed with CD.

The applicant stated that the Stelara® is a human IgG1k monoclonal antibody that binds with specificity to the p40 protein subunit, which is common to both the interleukin-12 (IL-12) and interleukin (IL-23) cytokines. IL-12 and IL-23 are naturally occurring cytokines that are involved in inflammatory and immune responses, such as natural killer cell activation and CD4+ T-cell differentiation and activation. In vitro models, the Stelara® was shown to disrupt IL-12 and IL-23 mediated signaling and cytokine cascades by blocking the interaction of these cytokines with a shared cell-surface receptor chain, IL-12Rβ1. The cytokines IL-12 and IL-23 have been implicated as important contributors to chronic inflammation. According to the applicant, IV induction therapy quickly achieves optimal blood levels of Stelara® so that blockade of IL-12 and IL-23 is maintained. This level of blockade is not achieved with subcutaneous administration.

The applicant further stated that other available CD anti-inflammatory or immune modulator therapies do not target the IL-12/IL-23p40 substrate. Rather, these therapies may target other integrin pairs such as the alpha4-beta7 integrins. Therefore, the applicant believed that the Stelara® drug is not substantially similar to any other approved drug for the treatment of moderately to severely active CD. As previously noted, the applicant asserted that, while the currently available agents are excellent medications, these agents do not successfully treat all patients diagnosed with CD, nor do these agents reliably sustain remission once a clinical response has been achieved. According to the applicant, the new use of the Stelara® offers an alternative to currently available treatment options, and has been shown to be successful in the treatment of patients who have failed treatment with the conventional agents currently being used for a diagnosis of CD, including TNF blockers. We are concerned that the Stelara®’s mechanism of action is similar to that of other immune system suppressors used in the treatment of patients diagnosed with moderately to severely active CD because other cytokine-selective monoclonal antibodies also disrupt cytokine-mediated signals crucial to the inflammatory process in patients diagnosed with CD.

With respect to the second criterion, whether a product is assigned to the same or a different MS–DRG, the applicant maintained that MS–DRGs 386, 387, and 385 (Inflammatory Bowel Disease with CC, without CC/MCC, and with MCC, respectively) and MS–DRGs 330, 329 and 331 (Major Small and Large Bowel Procedures with CC, without CC/MCC, and with MCC, respectively) are used to identify cases representing patients who may potentially be eligible for treatment using the Stelara®. The applicant researched claims data from the FY 2015 MedPAR file and found 10,344 cases. About 85 percent of potentially eligible cases were mapped to MS–DRGs for inflammatory bowel disease and most of the remainder of cases mapped to MS–DRGs for bowel surgery. We believe that potential cases involving Stelara® induction therapy may be assigned to the same MS–DRGs as cases representing patients who have been treated using currently available treatment options.

With respect to the third criterion, whether the new use of the technology involves the treatment of the same or similar type of disease and the same or similar patient population, according to the applicant, currently available pharmacologic treatments include antibiotics, mesalamines, corticosteroids, immunomodulators, tumor necrosis alfa (TNFα) inhibitors and anti-integrins. The applicant stated that the new use of the Stelara® for IV infusion is indicated for the treatment of adults (18 years and older) diagnosed with moderately to severely active CD who have: (1) Failed or were intolerant to treatment with immunomodulators or corticosteroids, but never failed treatment using a TNF blocker; or (2) failed or were intolerant to treatment with one or more TNF blockers. The applicant asserted that Stelara® for induction therapy is not substantially similar to other treatment options because it does not involve the treatment of the same or similar type of patient population. Patients who are eligible for treatment using the Stelara® induction therapy have failed other CD treatment modalities. The applicant believed that the subset of primary and secondary nonresponder patients to TNF inhibitor treatments is a patient population unresponsive to, or ineligible for, currently available treatments for diagnoses of moderate to severe CD. Based on the indications for the use of Stelara®, there is a class of patients who failed, or were intolerant to, treatment using immunomodulators or corticosteroids, but never failed treatment using a TNF blocker. The applicant indicated that, for those patients who never failed treatment with a TNF blocker, this class of patients can be recognized as two separate patient populations: One population of patients who have never received treatment using a TNF blocker, or the other population of patients who have received and responded to treatment using a TNF blocker. We believe that, if the new use of the Stelara® has the same mechanism of action as other immune system suppressors such as TNF blockers, the patient population that did not receive treatment using a TNF blocker may not be a new patient population because those patients may be able to receive treatment using, and would successfully respond to treatment using, a TNF blocker. Moreover, if the mechanism of action is the same as other immune system suppressors, we believe that the new use of the Stelara® may be targeted at a new patient population in some circumstances and instances, but we are concerned that it may not be targeted at a new patient population in all circumstances and instances.
We are inviting public comments on whether the Stelara® meets the newness criterion.

With regard to the cost criterion, the applicant conducted the following analysis to demonstrate that Stelara® meets the cost criterion. The applicant searched claims from the FY 2015 MedPAR file for cases with a principal ICD–9–CM diagnosis of 555.x (Regional Enteritis), which are cases of a diagnosis of Crohn’s Disease that may be eligible for treatment using Stelara®.

The applicant identified 10,344 cases that mapped to MS–DRGs. Approximately 85 percent of cases mapped to the following Inflammatory Bowel MS–DRGs: MS–DRGs 385 (Inflammatory Bowel Disease with MCC), 386 (Inflammatory Bowel Disease with CC), and 387 (Inflammatory Bowel Disease without CC/MCC). Similarly, 11 percent of the cases mapped to the following MS–DRGs for bowel surgery: MS–DRGs 329 (Major Small and Large Bowel Procedures with MCC), 330 (Major Small and Large Bowel Procedures with CC), and 331 (Major Small and Large Bowel Procedures without CC/MCC). The remaining cases (4 percent) represented all other digestive system disorders.

Using the 10,344 identified cases, the average unstandardized case-weighted charge per case was $39,935. The applicant then standardized the charges. The applicant did not remove charges for the current treatment because as discussed above Stelara® is indicated for use in patients who fail other treatments. The applicant then applied the 2-year inflation factor of 1.098446 from the FY 2015 IPPS/LTC final rule (81 FR 57286) to inflate the charges from FY 2015 to FY 2017. The applicant then added charges for the Stelara® technology. Specifically, the applicant assumed that hospitals would mark up Stelara® IV to the same extent that they currently mark-up Stelara® SC (J3357, ustekinumab, 1 mg). The applicant used the actual hospital mark-up based on charges in the 2017 OPPS proposed rule file (OPPS claims incurred and paid in CY 2015). Based on the FY 2017 IPPS/LTC PPS Table 10 thresholds, the average case-weighted threshold amount was $55,023. The inflated average case-weighted standardized charge per case was $69,826. Because the inflated average case-weighted standardized charge per case exceeds the average case-weighted threshold amount, the applicant maintained that the technology meets the cost criterion. We are inviting public comments whether Stelara® meets the cost criterion.

With regard to the third criterion, whether a technology represents a substantial clinical improvement over existing technologies, according to the applicant, the new use of the Stelara® has been shown to produce clinical response and remission in patients diagnosed with moderate to severe CD who have failed treatment using conventional therapies, including antibiotics, mesalamine, corticosteroids, immunomodulators, and TNFα inhibitors. Stelara® has been commercially available on the U.S. market for the treatment of patients diagnosed with psoriasis (PsO) since 2009 and the treatment of patients diagnosed with psoriatic arthritis (PsA) since 2013, and the applicant has maintained a safety registry, which enrolled over 12,000 patients since 2007. According to the applicant, the drug has been extremely well-tolerated, and the safety profile in patients diagnosed with CD has been consistent with that experienced in cases representing patients diagnosed with PsO and PsA.

The applicant presented the results of three pivotal trials involving over 1,300 patients diagnosed with moderate to severe CD. All three trials utilized a multicenter, double-blind, placebo-controlled study design. There were two single-dose IV induction trials, which included patients who had failed treatment using one or more TNFα inhibitors (UNITI–1) (N = 741), and patients who had failed treatment using corticosteroids and/or immunomodulators (UNITI–2) (N = 628). Responders to the single IV induction dose were then eligible to be enrolled in a maintenance trial (IM–UNITI) (N = 397), which began 8 weeks after administration of the single IV induction dose. IM–UNITI patients were given subcutaneous Stelara® and were treated for 44 weeks. Over half of the patients treated with 90mg of Stelara® every 12 weeks were able to achieve remission; a highly significant response compared to placebo, according to the applicant. The results of these trials have been published by the New England Journal of Medicine and the applicant provided the published studies.

Responders to the single IV induction dose were then eligible to be enrolled in a maintenance trial (IM–UNITI) (N = 397), which began 8 weeks after administration of the single IV induction dose. IM–UNITI patients were given subcutaneous Stelara® and were treated for 44 weeks. Over half of the patients treated with 90mg of Stelara® every 12 weeks were able to achieve remission; a highly significant response compared to placebo, according to the applicant. The results of these trials have been published by the New England Journal of Medicine and the applicant provided the published studies.

The published study supported the applicant’s assertion that Stelara® single IV dose induces response and remission in patients diagnosed with moderately to severely active CD that is refractory to either TNF antagonists or conventional therapy. Of the patients in the IM–UNITI trial receiving subcutaneous Stelara® at 8 weeks or 12 weeks, 53.1 percent and 48 percent, respectively, were in remission at week 44 as compared with 35.9 percent of those patients receiving treatment using placebo.

The applicant submitted published results of a multicenter, double-blind, placebo controlled Phase III study of Stelara®. We are concerned that the study did not effectively establish the need for Stelara® induction therapy. Also, the median age of patients in the study was 37 years, and we are concerned that the study did not include a significant amount of older patients.

We also are concerned that we do not have enough information to determine that the new use of the Stelara® is a substantial clinical improvement over existing technologies for the treatment of moderate to severe CD. We note that the UNITI–1, UNITI–2, and IMUNITI trials were completed to evaluate efficacy and safety of Stelara®, not superiority of Stelara® to current conventional therapy. Our concerns are based on a lack of head-to-head trials comparing IV induction and maintenance Stelara® therapy with conventional therapy in patients diagnosed with moderate to severe CD that are also primary and secondary nonresponders to treatment using TNF alpha inhibitor therapy. We recognize the subset of primary and secondary nonresponder patients to TNF inhibitor treatments as a patient population unresponsive to, or ineligible for, currently available treatments for diagnoses of moderate to severe CD. However, we believe that this primary and secondary TNF inhibitor non-responder patient population represents patients that experience a gap in treatment for diagnoses of moderate to severe CD. Specifically, we recognize the nonresponder patient population as described by Simon et al. as those patients who are TNF inhibitor immunogenicity failures, pharmacokinetic failures, and/or pharmacodynamics failures. We also note the supplemental data in Feagan et al.’s publication summarized the primary and secondary nonresponders in UNITI–1. However, we are not clear how the inclusion of the TNF alpha

16 Ibid.
17 Ibid.
inhibitor intolerant patients with primary and secondary TNF alpha inhibitor failure patients impacts the final comparison of the placebo and treatment arms. In addition, we note that in the UNITI–1, UNITI–2, and IMUNITI studies all treatment arms were allowed to continue conventional treatments for diagnoses of CD throughout the study. We are concerned that it is difficult to determine whether the new use of the Stelara® represents a substantial clinical improvement over existing technologies with the concurrent use of other conventional CD medications throughout the duration of the UNITI–1, UNITI–2, and IMUNITI studies.

Also, as mentioned earlier, based on the indications for the use of the Stelara®, there is a class of patients who failed, or were intolerant to, treatment with immunomodulators or corticosteroids, but never failed treatment using a TNF blocker. According to the applicant, for those patients who never failed treatment using a TNF blocker, this patient population can be recognized as two separate patient populations: one patient population representing patients who never received treatment using a TNF blocker, or the other patient population representing patients who received and responded to treatment using a TNF blocker. In the patient population that did not receive treatment using a TNF blocker, we are unsure if the new use of the Stelara® represents a substantial clinical improvement because it is possible that some patients will have a positive response to treatment using a TNF blocker and will not respond successfully to treatment using Stelara®, or some patients may have a positive response to both treatment using a TNF blocker and using Stelara®, or some patients may not respond to treatment using a TNF blocker, but will have a positive response to treatment using Stelara®.

We are inviting public comments on whether the Stelara® meets the substantial clinical improvement criterion.

We did not receive any written public comments in response to the New Technology Town Hall meeting notice regarding the application of Stelara® for new technology add-on payments.

d. KTE–C19 (Axicabtagene Ciloleucel)

Kite Pharma, Inc. submitted an application for new technology add-on payments for KTE–C19 (axicabtagene ciloleucel) for FY 2018. The KTE–C19 technology has not received FDA approval as of the time of the development of this proposed rule. KTE–C19 is an engineered autologous T-cell immunotherapy used for the treatment of adult patients with relapsed/refractory aggressive B-cell non-Hodgkin lymphoma (NHL) who are ineligible for autologous stem cell transplant (ASCT). KTE–C19 is a single intravenous infusion of T-cell immunotherapy.

The applicant noted that KTE–C19 was granted Breakthrough Therapy Designation by the FDA on December 3, 2015, for the treatment of patients with refractory DLBCL, PMBCL, and TFL forms of aggressive B-cell NHL. The applicant submitted a request for priority review by the FDA in December 2016. The applicant stated that, when approved by the FDA, KTE–C19 would represent the only FDA-approved treatment for adult patients with relapsed refractory aggressive B-cell NHL who are ineligible for ASCT. Currently, there are no ICD–10–CM/PCS codes that describe the administration and use of KTE–C19. The applicant has submitted an application for a unique ICD–10–PCS procedure code to uniquely identify KTE–C19. If approved, the code will be effective October 1, 2017 (FY 2018). According to the applicant, adult NHL represents a heterogeneous group of B-cell malignancies with varying patterns of behavior and response to treatment. B-cell NHL can be classified as either aggressive, or indolent disease, with aggressive variants including diffuse large B-cell lymphoma (DLBCL); primary mediastinal large B cell lymphoma (PMBCL) and transformed follicular lymphoma (TFL). Within NHL, DLBCL is the most common subtype of NHL, accounting for approximately 30 percent of patients with NHL, and survival without treatment is measured in months.20 21

The applicant stated that, since the 1970s, cyclophosphamide, doxorubicin, vincristine, and prednisone (CHOP) has been the mainstay of therapy with more intensive regimens failing to show improved overall survival. The applicant further stated that the approval in 2006 of the anti-CD20 monoclonal antibody rituximab and its addition to the traditional CHOP regimen, R–CHOP, for patients with newly diagnosed aggressive NHL resulted in a dramatic improvement in NHL therapy. The combination of CHOP and R–CHOP is now first-line therapy for treatment of patients diagnosed with DLBCL with complete response rates upwards of 76 percent.22 Data from the Surveillance, Epidemiology and End Results (SEER) registries have reflected an observed increase of the median overall survival from 20 to 47 months over the last two decades. Despite the improved therapies, only 50 to 70 percent of newly diagnosed patients are cured by standard first-line therapy alone.23 Furthermore, relapsed or refractory (r/r) disease continues to carry a poor prognosis because only 50 percent of patients are eligible for more intensive second-line regimens, followed by high dose chemotherapy (HDT) and ASCT. Second-line chemotherapy regimens studied to date include rituximab, ifosfamide, carboplatin and etoposide (ICE) and rituximab, dexamethasone, cytarabine, and cisplatin (R–DHAP), followed by consolidative HDT/ASCT. Both regimens offer similar overall response rates (ORR) of 51 percent with 1 in 4 patients achieving long-term complete response (CR) at the expense of increased toxicity.24 Given the modest response to second line therapy and/or HDT/ASCT, the population of patients with the highest unmet need is those with chemorefractory disease, which include DLBCL, PMBCL and TFL. These patients are defined as either progressive disease (PD) as best response to chemotherapy, stable disease as best response following 4 cycles of first-line or 2 cycles of inter-line therapy, or relapse within 12 months of ASCT.25 26 Based on these definitions and available data from a multicenter retrospective study (SCHOLAR–1), chemorefractory disease treated with current and historical standards of care has consistently poor

outcomes with an ORR of 26 percent and median OS of 6.6 months.

According to the applicant, KTE–C19 is a different pathway to treat patients diagnosed with relapsed or refractory disease. KTE–C19 is supplied as a T-cell suspension for infusion. With KTE–C19 treatment, a patient’s own T-cells are harvested and engineered ex vivo by retroviral transduction of a chimeric antigen receptor (CAR) construct encoding an anti-CD19 CD28/CD3-zeta. The anti-CD19 CAR T-cells are expanded and infused back into the patient. The new anti-CD19 CAR T-cells can recognize and eliminate CD19 antigen expressing target cells, an antigen also expressed on the cell surface of B-cell lymphomas and leukemias. According to the applicant, prior to KTE–C19 immunotherapy, the patient would have received outpatient administration of a non-myeloablative conditioning chemotherapy regimen consisting of cyclophosphamide 500 mg/m2 IV and fludarabine 30 mg/m2 IV for 3 days at days -5, -4, and -3 before the infusion of KTE–C19 at Day 0. The applicant noted that, if KTE–C19 infusion is delayed more than 2 weeks, readministration of the conditioning chemotherapy regimen may be required. Hospitalization is recommended for the infusion of KTE–C19.

As discussed earlier, if a technology meets all three of the substantial similarity criteria, it would be considered substantially similar to an existing technology and would not be considered “new” for purposes of new technology add-on payments.

With regard to the first criterion, the applicant stated that KTE–C19 does not use the same or similar mechanism of action to achieve a therapeutic outcome as any other drug or therapy assigned to the same or a different MS–DRG. The applicant further stated that KTE–C19 is the first engineered autologous cellular immunotherapy comprised of CAR T-cells that recognizes CD19 express cancer cells and normal B-cells; therefore, the applicant believed that KTE–C19’s mechanism of action is distinct and unique from any other cancer drug or biologic that is currently approved for use in the treatment of aggressive B-cell NHL, namely single-agent or combination chemotherapy regimens.

With regard to the second criterion, whether a product is assigned to the same or a different MS–DRG, the applicant noted that based on the 2014 Standard Analytic files, cases potentially eligible for treatment using the KTE–C19 and representing the target patient population span 50 unique MS–DRGs and 73 percent of all of the cases within these 50 unique MS–DRGs that represent potentially eligible cases for treatment using KTE–C19 map to the following 4 MS–DRGs: MS–DRG 840 (Lymphoma & Non-Acute Leukemia with MCC); MS–DRG 841 (Lymphoma & Non-Acute Leukemia with CC); MS–DRG 846 (Chemotherapy without Acute Leukemia as Secondary Diagnosis with MCC); and MS–DRG 847 (Chemotherapy without Acute Leukemia as Secondary Diagnosis with CC). The applicant stated that, with the assignment of the unique KTE–C19-specific ICD–10–PCS code, patient cases where KTE–C19 is used will be distinguishable. However, patient cases where KTE–C19 is used and patient cases that are treated for DLBCL map to the same MS–DRGs.

With regard to the third criterion, whether the new use of the technology involves the treatment of the same or similar type of disease and the same or similar patient population, the applicant asserted that when approved by the FDA, KTE–C19 would represent the only FDA-approved treatment for adult patients diagnosed with relapsed or refractory aggressive B-cell NHL who are ineligible for ASCT. As a result, the applicant stated that KTE–C19 is not substantially similar to any existing technology and meets the newness criterion. CMS is concerned the CAR technology used in KTE–C19 may have a mechanism of action similar to that seen with the use of bispecific T cell engager (BiTE) technology.

We are inviting public comments on whether KTE–C19 meets the substantial similarity criteria and the newness criterion.

With respect to the cost criterion, the applicant provided an analysis to demonstrate that KTE–C19 meets the cost criterion. The applicant used the 2014 and 2015 100 Percent Inpatient Standard Analytic File (SAF) to assess the MS–DRGs that are most relevant to patients that may be potentially eligible for treatment using KTE–C19. The sample was restricted to patients discharged in FY 2015. The applicant searched for cases with an ICD–9–CM diagnosis code from the series of 200.7x (large cell lymphoma).

The applicant sought to ensure that claims included in the cost criterion analysis reflected charges for treating patients diagnosed with DLBCL and, therefore, minimized the chance that charges were related to other conditions. Therefore, the applicant searched for cases with the following criteria:

- A primary diagnosis with an ICD–9–CM diagnosis code from the series of 200.7x (large cell lymphoma) to identify cases of DLBCL with or without chemotherapy; or
- A secondary diagnosis with an ICD–9–CM diagnosis code from the series of 200.7x (large cell lymphoma) combined with an ICD–9–CM diagnosis code of V58.11, or V58.12, or ICD–9–CM procedure code 99.25, 99.28, 00.15 or 00.10 to identify cases of DLBCL that received chemotherapy during their hospitalization.

The applicant excluded claims where the MS–DRG was missing. Medicare was not the primary payer, there were zero covered charges or zero covered days, or the provider was not in the FY 2017 IPPS/LTCH PPS Final Rule Impact File. Additionally, patients under age 18 were excluded to align with the proposed label that is being prepared for submission with the KTE–C19 Biologics License Application (BLA). After applying the trims above, the results showed 762 cases that mapped to 50 MS–DRGs with 11 MS–DRGs containing more than 10 cases. The 11 MS–DRGs contained a total of 526 cases.

The applicant noted that MS–DRGs 840, 841, 846, and 847 accounted for 554 (73 percent) of the 762 cases in the cohort.

Using the 702 identified cases, the average unstandardized case-weighted charge per case was $71,725. The applicant then standardized the charges. The applicant noted that adult patients with relapsed/refractory aggressive B-cell NHL who are ineligible for ASCT would generally not be receiving treatment with both chemotherapy and KTE–C19. Therefore, all charges listed in the chemotherapy revenue centers (331, 332, and 335) were removed. The applicant then applied the 2-year inflation factor of 1.098446 from the FY 2017 IPPS/LTCH final rule (81 FR 57286) to inflate the charges from FY 2015 to FY 2017. Based on the FY 2017 IPPS/LTCH PPS Table 10 thresholds, the average case-weighted threshold amount was $55,023. The inflated average case-weighted standardized charge per case was $69,826. Because the inflated average case-weighted standardized charge per case exceeds the average case-weighted threshold amount, the applicant maintained that the technology meets the cost criterion. The applicant noted that it was not necessary to take into account the average per patient cost of the technology because the inflated average case-weighted standardized charge per case exceeds the average case-weighted threshold amount without the average per patient cost of the technology. The applicant provided the following three sensitivity analyses to further demonstrate that the technology meets
the cost criterion. The three sensitivity analyses consisted of: (1) cases representing patients identified with an ICD–9–CM diagnosis code 200.7x (large cell lymphoma) and cases representing patients identified with a secondary DLBCL diagnosis who did not receive chemotherapy; (2) cases representing patients identified with a primary or secondary ICD–9–CM diagnosis code from the series of 200.7x (large cell lymphoma) who received chemotherapy; and (3) cases representing patients under a broader ICD–9–CM diagnosis code range to capture other types of lymphoma. In all three of the sensitivity analyses, the inflated average case-weighted standardized charge per case exceeded the average case-weighted threshold amount. We are inviting public comments on whether KTE–C19 meets the cost criterion.

According to the applicant, KTE–C19 represents a substantial clinical improvement over existing technologies used in the treatment of patients with aggressive B-cell NHL. The applicant asserted that KTE–C19 can benefit the patient population with the highest unmet need, patients with refractory or relapsed disease after failure of first-line or second-line therapy, and patients who have failed or are ineligible for ASCT. These patients otherwise have adverse outcomes as demonstrated by historical control data. Regarding clinical data for KTE–C19, the applicant stated that historical control data was the only ethical and feasible comparison information for these chemorrefractory, aggressive NHL patients who have no other available treatment options and have a very short lifespan without therapy. According to the applicant, based on meta-analysis of outcomes in chemorrefractory DLBCL, there are no curative options for aggressive B-cell NHL patients regardless of refractory subgroup, line of therapy, and disease stage with their median overall survival being 6.6 months.

The applicant provided clinical data from the pivotal Study 1 (ZUMA–1, KTE–C19–101), Phase I and II. The applicant also provided supportive evidence from Study 2 (NCI 009–C–0082). Study 1 is a Phase I–II multicenter, open label study evaluating the safety and efficacy of the use of KTE–C19 in patients diagnosed with aggressive refractory NHL. The trial consists of two distinct phases designed as Phase I (n=7) and Phase II (n=92). Phase II is a multi-cohort open label study evaluating efficacy of KTE–C19. Study 1 subjects were treated with cyclophosphamide and fludarabine conditioning chemotherapy, followed by a target dose of 2 × 10^10 anti-CD19 CAR T-cells per kg body weight. Study 2 subjects were treated with cryopreserved autologous anti-CD19 CAR T-cells, which were manufactured by a similar, but different process than that used for KTE–C19. The applicant noted that, as of the analysis cutoff date for the interim analysis, the results of Study 1 demonstrated rapid and substantial improvement in objective, or overall response rate. The overall response rate was 7% percent (49 responders among 62 subjects), with 76 percent overall response rate in Cohort 1 (39 responders among 51 subjects) and 91 percent in Cohort 2 (10 responders among 11 subjects) versus historical control of 26 percent. According to the applicant, Study 1 overall response rates were consistent across all age groups, with those patients greater than 65 years of age responding at the rates consistent with those under age 65 years and consistent with earlier, positive results from Study 2. The applicant further stated that pre-specified criteria for demonstration of early efficacy were met and an independent safety monitoring board (DSMB) confirmed the efficacy results and found no additional safety signals.

The applicant further stated that evidence of substantial improvement regarding the efficacy of KTE–C19 for the treatment of chemorrefractory, aggressive B-cell NHL is supported by the complete response rates of KTE–C19 in Study 1 (52 percent) versus the historical control (9 percent). Additionally, the applicant noted that the results of Study 1 have demonstrated that treated patients experienced a rapid response to KTE–C19 with 52 percent showing complete response at 3 months, and 41 percent at 1 month.

As noted above, the applicant cited data results from Study 2, which is an ongoing Phase I safety and efficacy study in which anti-CD19 CAR T-cells were manufactured using a process similar to, but different from, KTE–C19 to yield cryopreserved autologous anti-CD19 CAR T-cells. From Study 2, a subset of 13 patients with a diagnosis of DLBCL/PMBCL was noted to be comparable to those treated in Study 1. The applicant noted that all patients were diagnosed with refractory DLBCL, received similar doses of conditioning chemotherapy, and were infused with the cryopreserved autologous anti-CD19 CAR T-cells (which have been shown to result in an immunotherapy comparable to KTE–C19). The applicant noted that the results from Study 2 demonstrated the following: (a) an overall response rate of 69 percent (9 responders among 13 patients) (95 percent CI 38.6, 90.9); (b) 47 percent of patients had complete response at month 3 (ongoing 6+ to 20+ months); and (c) complete response was observed as early as 1 month in 57 percent of patients in Study 2.

According to the applicant, further results will be reported in February 2017.

The applicant also cited safety results from the pivotal Study 1, Phase II. According to the applicant, almost all patients in Study 1 (95 percent) experienced Grade 3 or higher adverse events with onset on or after commencement of conditioning chemotherapy, including cytopenias (Grade 3 and 4 anemia, neutropenia, thrombocytopenia, and lymphopenia were 40 percent, 40 percent, 29 percent, and 5 percent respectively), and infection (Grade 3 or worse urinary tract infection, clostridium difficile colitis and lung infection were 5 percent, 5 percent, and 6 percent respectively). All patients were treated according to standard of care. The clinical trial protocol stipulated that patients were infused with KTE–C19 in the hospital inpatient setting and were monitored in the inpatient setting for at least 7 days for early identification and treatment of KTE–C19 related toxicities, which primarily include cytokine release syndrome and neurotoxicities. The applicant stated that KTE–C19 is expected to be administered in the hospital inpatient setting to assure appropriate monitoring of patient adverse events. The applicant noted that the interim analysis of Study 1 showed the following: length of stay following KTE–C19 infusion was a median of 15 days; cytokine release syndrome (Grade 3 or higher, 18 percent) and neurotoxicity (Grade 3 or higher, 34 percent) were self-limiting and generally reversible; two patients died from KTE–C19 related adverse events (hemophagocytic lymphohistiocytosis and cardiac arrest in the setting of cytokine release syndrome). The medications most often used to treat KTE–C19 clinical trial complications included growth factors, blood products, anti-infectives, steroids, tocilizumab, and vasopressors. In the majority of patients (92 percent), the applicant noted that predominant toxicities associated with the use of KTE–C19, cytokine release syndrome and neurologic events, resolved by data cutoff. Median days to resolution of cytokine release syndrome complications post-KTE–C19 infusion was 9 days, with median days to resolution of KTE–C19-related
neurologic events post-KTE–C19 infusion of 18 days. According to the applicant, there were no clinically important differences in adverse event rates across age groups (younger than 65; 65 or older), including cytokine release syndrome and neurotoxicity, and KTE–C19-related adverse events in Study 1 were consistent with the earlier Study 2 experience.

The applicant further noted that by the cutoff date for the interim analysis of Study 1, among all KTE–C19 treated patients, 12 patients in Study 1, Phase II, including 10 from Cohort 1 and 2 from Cohort 2, died. Eight of these deaths were due to disease progression. One subject had disease progression after KTE–C19 treatment and subsequently had ASCT. After ASCT, the subject died due to sepsis. Two subjects (3 percent) died due to KTE–C19 related AEs (Grade 5 hemophagocytic lymphohistiocytosis event and Grade 5 anoxic brain injury), and one died due to an AE deemed unrelated to KTE–C19 (Grade 5 pulmonary embolism), without disease progression.

We are concerned that there are no published results showing any survival benefit from the treatment. We also are concerned with the limited number of subjects (n=82) that were studied after infusion of KTE–C19 T-cell immunotherapy. Although the applicant references Study 2, we are concerned that the applicant has included data on DLBL/PMBCL patients that did not specifically receive KTE–C19.

Additionally, we are concerned that Study 2 was based on 13 patients which can result in skewed outcomes due to a small patient population. Finally, we note that, for Study 1 and Study 2, the data on overall survival are not reported.

We are inviting public comments on whether KTE–C19 meets the substantial clinical improvement criterion.

Comment: The applicant stated that it has been notified by the United States Adopted Names Council (USAN Council) that the technology’s name for KTE–C19 has been revised from ‘‘axicabtagene ciloleucel’’ to ‘‘axicabtagene cileoleucel.’’ In addition, the applicant requested that all references by CMS to the technology’s name of KTE–C19 use this final naming convention of ‘‘axicabtagene cileoleucel.’’

Response: We appreciate the applicant’s updated information and have corrected the name of the technology throughout the discussion above.

e. VYXEOSTM (Cytarabine and Daunorubicin Liposome for Injection)

Celator Pharmaceuticals, Inc. submitted an application for new technology add-on payments for VYXEOSTM for FY 2018. The proposed indication for the use of VYXEOSTM, which has not received FDA approval as of the time of the development of this proposed rule, is the treatment of adult patients diagnosed with acute myeloid leukemia (AML).

AML is a type of cancer in which the bone marrow makes abnormal myeloblasts (immature bone marrow white blood cells), red blood cells, and platelets. If left untreated, AML progresses rapidly. Normally, the bone marrow makes blood stem cells that develop into mature blood cells over time. Stem cells have the potential to develop into many different cell types in the body. Stem cells can act as an internal repair system, dividing, essentially without limit, to replenish other cells. When a stem cell divides, each new cell has the potential to either remain a stem cell or become a specialized cell, such as a muscle cell, a red blood cell or a brain cell, etc. A myeloid stem cell may become a myeloid stem cell or a lymphoid stem cell. Lymphoid stem cells become white blood cells. A myeloid stem cell becomes one of three types of mature blood cells: (1) red blood cells that carry oxygen and other substances to body tissues; (2) white blood cells that fight infection; or (3) platelets that form blood clots and help to control bleeding. In patients diagnosed with AML, the myeloid stem cells usually become a type of myeloblast. The myeloblasts in patients diagnosed with AML are abnormal and do not become healthy white blood cells. Sometimes in patients diagnosed with AML, too many stem cells become abnormal red blood cells or platelets. These abnormal cells are called leukemia cells or blasts.

AML is defined by the World Health Organization (WHO) as >20 percent blasts in the bone marrow or blood. AML can also be diagnosed if the blasts are found to have a chromosome change that occurs only in a specific type of AML, even if the blast percentage does not reach 20 percent. Leukemia cells can build up in the bone marrow and blood, resulting in less room for healthy white blood cells, red blood cells, and platelets. When this occurs, infection, anemia, or increased risk for bleeding may result. Leukemia cells can spread outside the bone marrow to other parts of the body, including the central nervous system (CNS), skin, and gums.

Treatment of AML diagnoses usually consists of two phases: remission induction and post-remission therapy. Phase one, remission induction, is aimed at eliminating as many myeloblasts as possible. The most common used remission induction regimens for AML diagnoses are the ‘‘7+3’’ regimens using an antineoplastic and an anthracycline. Cytarabine and daunorubicin are two commonly used drugs for ‘‘7+3’’ remission induction therapy. Cytarabine is continuously administered intravenously over the course of 7 days, while daunorubicin is intermittently administered intravenously for the first 3 days. The ‘‘7+3’’ regimen typically achieves a 70 to 80 percent complete remission (CR) rate in most patients under 60 years of age.

High rates of CR are not generally seen in older patients for a number of reasons, such as different leukemia biology, much higher incidence of adverse cytogenetic abnormalities, higher rate of multidrug resistant leukemic cells, and comparatively lower patient performance status (the standard criteria for measuring how the disease impacts a patient’s daily living abilities). Intensive induction therapy has worse outcomes in this patient population.

The applicant asserted that many older adults diagnosed with AML have a poor performance status presentation and multiple medical comorbidities that make the use of intensive induction therapy quite difficult or contraindicated altogether. Moreover, the CR rates of poor-risk patients diagnosed with AML are substantially higher in patients >60 years old; owing to a higher proportion of secondary AML, disease developing in the setting of a prior myeloid disorder, or prior cytotoxic chemotherapy. Therefore, less than half of older adults diagnosed with AML achieve CR with combination induction regimens.

The combination of cytarabine and an anthracycline, either as ‘‘7+3’’ regimens or as part of a different regimen incorporating other cytotoxic agents, may be used as so-called ‘‘salvage’’ induction therapy in the treatment of adults diagnosed with AML who experience relapse in an attempt to
achieve CR. According to the applicant, while CR rates of success vary widely depending on underlying disease biology and host factors, there is a lower CR rate overall in achievement of CR with “7+3” regimens compared to VYXEOS™ therapy. In addition, “7+3” regimens produce a CR rate of approximately 50 percent in younger adults who have relapsed, but were in CR for at least 1 year.  

VYXEOS™ is a nano-scale liposomal formulation containing a fixed combination of cytarabine and daunorubicin in a 5:1 molar ratio. This formulation was developed by the applicant using a proprietary system known as CombiPlex. According to the applicant, CombiPlex addresses several fundamental shortcomings of conventional combination regimens, specifically the conventional “7+3” free drug dosing, as well as the challenges inherent in combination drug development, by identifying the most effective synergistic molar ratio of the drugs being combined in vitro, and fixing this ratio in a nano-scale drug delivery complex to maintain the optimized combination after administration and ensuring exposure of this ratio to the tumor.

Cytarabine and daunorubicin are co-encapsulated inside the VYXEOS™ liposome at a fixed ratiometrically, optimized 5:1 cytarabine:daunorubicin molar ratio. According to the applicant, encapsulation maintains the synergistic ratios, reduces degradation, and minimizes the impact of drug transporters and the effect of known resistant mechanisms. The applicant stated that the 5:1 molar ratio has been shown, in vitro, to maximize synergistic antitumor activity across multiple leukemic and solid tumor cell lines, including AML, and in animal model studies to be optimally efficacious compared to other cytarabine:daunorubicin ratios. In addition, the applicant stated that in clinical studies, the use of VYXEOS™ has demonstrated consistently more efficacious results than the conventional “7+3” free drug dosing. VYXEOS™ is intended for intravenous administration after reconstitution with 19 mL sterile water for injection. VYXEOS™ is administered as a 90-minute intravenous infusion on days 1, 3, and 5 (induction therapy), as compared to the “7+3” free drug dosing, which consists of two individual drugs administered on different days, including 7 days of continuous infusion.

With regard to the “newness” criterion, the applicant indicated that the rolling New Drug Application (NDA) submission to the FDA for VYXEOS™ began on September 30, 2016. The applicant stated that it intends to request Priority Review from the FDA. VYXEOS™ is currently available in the United States only on an investigational basis, under an Investigational New Drug (IND) designation. Breakthrough Therapy designation was granted on May 19, 2016, for the treatment of adults diagnosed with therapy-related AML (t-AML) or AML with myelodysplasia-related changes (AML–MRC). Fast Track designation was granted by the FDA in January 2015 for the treatment of elderly patients diagnosed with secondary AML. Orphan Drug designation was granted by the FDA on August 22, 2008, for the treatment of acute AML. VYXEOS™ had not received pre-market (PMA) approval from the FDA at the time of development of this proposed rule. However, the applicant anticipates receiving approval from the FDA by July 1, 2017. The applicant also has submitted a request for a unique ICD–10–PCS code, beginning with FY 2018.

As discussed earlier, if a technology meets all three of the substantial similarity criteria, it would be considered substantially similar to an existing technology and would not be considered “new” for purposes of new technology add-on payments.

With regard to the first criterion, whether a product uses the same or a similar mechanism of action to achieve a therapeutic outcome, the applicant asserted that VYXEOS™ does not use the same or similar mechanism of action to achieve a therapeutic outcome as any other drug assigned to the same or a different DRG. The applicant stated that no other AML treatment is designed, nor is able, to deliver a fixed, ratiometrically optimized and synergistic drug:drug ratio of 5:1 cytarabine to daunorubicin, and selectively target and accumulate at the site of malignancy, while minimizing unwanted exposure, which the applicant based on the data results of preclinical and clinical studies of the use of VYXEOS™. The applicant indicated that VYXEOS™ is a nano-scale liposomal formulation of a fixed combination of cytarabine and daunorubicin. Further, the applicant stated that the rationale for the development of VYXEOS™ is based on prolonged delivery of synergistic drug ratios utilizing the applicant’s proprietary targeted drug delivery technology. According to the applicant, conventional “7+3” free drug dosing has no delivery complex, and these individual drugs are administered without regard to their ratio dependent interaction. According to the applicant, enzymatic inactivation and imbalanced drug efflux and transporter expression reduce drug levels in the cell. Decreased cytotoxicity leads to cell survival, emergence of drug resistant cells, and decreased overall survival. The applicant provided the results of clinical studies to demonstrate that the CombiPlex technology and the ratiometric dosing of VYXEOS™ represent a shift in anticancer agent delivery, whereby the fixed, optimized dosing provides less drug to achieve improved efficacy, while maintaining a favorable risk-benefit profile. The results of this ratiometric dosing approach are in contrast to the typical combination chemotherapy development that establishes the recommended dose of one agent and then adds subsequent drugs to the combination at increasing concentrations until the aggregate effects of toxicity are considered to be limiting (the “7+3” drug regimen). According to the applicant, this current approach to combination chemotherapy development assumes that maximum therapeutic activity will be achieved with maximum dose intensity for all drugs in the combination, and ignores the possibility that more subtle concentration-dependent drug interactions could result in frankly synergistic outcomes.

The applicant maintained that, while VYXEOS™ contains no novel active agents, its innovative drug delivery mechanism appears to be a superior way to deliver the two active compounds in an effort to optimize their efficacy in killing leukemic blasts. However, we are concerned it is possible that VYXEOS™ may use a similar mechanism of action compared to current treatment because both the current treatment regimen and VYXEOS™ are used in the treatment of AML by intravenous administration of cytarabine and daunorubicin.

With respect to the second criterion, whether a product is assigned to the same or a different MS–DRG, the applicant maintained that based on the 2014 and 2015 100 Percent Inpatient Standard Analytic files, cases representing patients potentially eligible for treatment using VYXEOS™ and the target patient population span 134 unique MS–DRGs, and 78 percent of all of the cases within these 134 unique MS–DRGs map to the following 4 MS–DRGs: 834 (Acute Leukemia Without Major O.R. Procedure With MCC), 837 (Chemotherapy With Acute Leukemia as SDX or With High Dose Chemotherapy...
Agent with MCC), 838 (Chemotherapy With Acute Leukemia as SDX With CC or High Dose Chemotherapy Agent), and 839 (Chemotherapy With Acute Leukemia as SDX Without CC/MCC). We believe that these are the same MS–DRGs that identify cases representing patients who are treated for AML.

With respect to the third criterion, whether the new use of the technology involves the treatment of the same or similar type of disease and the same or similar patient population, the applicant asserted that VYXEOS™ is indicated for the use in patients diagnosed with high-risk AML. However, we believe that VYXEOS™ involves the treatment of the same patient population as other AML treatment therapies.

We are inviting public comments on whether VYXEOS™ is substantially similar to existing technology, including whether the mechanism of action of VYXEOS™ differs from the mechanism of action of the current treatment regimen. We also are inviting public comments on whether VYXEOS™ meets the newness criterion.

With regard to the cost criterion, the applicant conducted the following analysis. The applicant used the 2014 and 2015 100 Percent Inpatient Standard Analytic Files (SAFs) to assess the MS–DRGs assigned for hospitalizations most likely to represent the MS–DRGs assigned for analysis. The applicant used the 2014 applicant conducted the following ICD–9–CM procedure codes:

- Bone marrow transplant, not otherwise specified;
- Autologous bone marrow transplant without purging;
- Allogeneic bone marrow transplant with purging;
- Allogeneic hematopoietic stem cell transplant without purging;
- Allogeneic hematopoietic stem cell transplant with purging;
- Cord blood cell transplant;
- Autologous hematopoietic stem cell transplant with purging;
- Allogeneic hematopoietic stem cell transplant;
- and Bone marrow transplant with purging.

According to the applicant, the eligible cases span 134 unique MS–DRGs, 14 of which contain more than 10 cases. The most common MS–DRGs are MS–DRGs 834, 837, 838, and 839. These 4 MS–DRGs account for 3,601 (78 percent) of the 4,613 potential eligible cases.

Using the 4,613 identified cases, the average unstandardized case-weighted charge per case was $203,234. The applicant then standardized the charges. The applicant noted that the average case-weighted threshold amount, the average case-weighted standardized charge per case exceeded the average case-weighted threshold amount. We are inviting public comments whether VYXEOS™ meets the cost criterion.

With regard to substantial clinical improvement, according to the applicant, clinical data results have shown that the use of VYXEOS™ represents a substantial clinical improvement for the treatment of AML in newly diagnosed high-risk, older (60 years and older) patients, marked by statistically significant improvements in overall survival, event free survival and response rates, and in relapsed patients aged 18 to 65 years of age, where a statistically significant improvement in overall survival was documented for the poor-risk subset of patients as defined by the European Prognostic Index. In both groups of patients, the applicant stated that there was significant improvement in survival for the high-risk patient group. The applicant provided the following specific clinical data results.

- The applicant stated the clinical data results show that treatment with VYXEOS™ in older patients (60 years of age and older) diagnosed with untreated, high-risk AML will result in superior survival rates, as compared to patients treated with conventional "7+3" free drug dosing. The applicant provided a summary of the pivotal Phase III Study 301 in which 309 patients were enrolled, with 153 patients randomized to the VYXEOS™ arm and 156 to the "7+3" free drug dosing arm. Among patients aged 60 to 69 years, there were 96 patients in the VYXEOS™ arm and 102 in the "7+3" free drug dosing arm; for patients aged 70 to 75 years, there were 57 and 54 patients in each arm, respectively. The applicant noted that the data results from the Phase III Study 301 demonstrated that first-line treatment of patients diagnosed with high-risk AML in the VYXEOS™ arm resulted in substantially greater median overall survival of 9.56 months versus 5.95 months in the "7+3" free drug dosing arm (hazard ratio of 0.69; p = 0.005).
The applicant further asserted that high-risk, older patients (60 years of age and older) previously untreated for diagnoses of AML will have a lower risk of early death when treated with VYXEOS™ than those treated with the conventional “7+3” free drug dosing. The applicant cited Medeiros, et al. 2015,31 which reported a large observational study of Medicare beneficiaries and noted the following: The data result of the study showed that 50 to 60 percent of elderly patients diagnosed with AML remain untreated following diagnosis; treated patients were more likely younger, male, and married, and less likely to have secondary diagnoses of AML, poor performance indicators, and poor comorbidity scores compared to untreated patients; and in multivariate survival analyses, treated patients exhibited a significant 33 percent lower risk of death compared to untreated patients.

Based on data from the Phase III Study 301,32 the applicant cited the following results: The rate of 60-day mortality was less in the VYXEOS™ arm (13.7 percent) versus the “7+3” free drug dosing arm (21.2 percent); the reduction in early mortality was due to fewer deaths from refractory AML (3.3 percent versus 11.3 percent), with very similar rates of 60-day mortality due to adverse events (10.4 percent versus 9.9 percent); there were fewer deaths in the VYXEOS™ arm versus the “7+3” free drug dosing arm during the treatment phase (7.8 percent versus 11.3 percent); and there were fewer deaths in the VYXEOS™ arm during the follow-up phase than in the “7+3” free drug dosing arm (59.5 percent versus 71.5 percent).

The applicant asserted that high-risk, older patients (60 years of age and older) previously untreated for a diagnosis of AML exhibited statistically significant improvements in response rates after treatment with VYXEOS™ versus treatment with the conventional “7+3” free drug chemotherapy dosing, suggesting that the use of VYXEOS™ is a superior pre-transplant induction treatment versus “7+3” free drug dosing. Restoration of normal hematopoiesis is the ultimate goal of any therapy for AML diagnoses. The first phase of treatment consists of induction chemotherapy, in which the goal is to “empty” the bone marrow of all hematopoietic elements (both benign and malignant), and to allow repopulation of the marrow with normal cells, thereby yielding remission. According to the applicant, post-induction response rates were significantly higher following the use of VYXEOS™, which elicited a 47.7 percent total response rate and a 37.3 percent rate for CR, whereas the total response and CR rates for the “7+3” free drug dosing arm were 33.3 percent and 25.6 percent, respectively. The CR + CRi rates for patients aged 60 to 69 years were 50.0 percent in the VYXEOS™ arm and 36.3 percent in the “7+3” free drug dosing arm, with an odds ratio of 1.76 (95 percent CI, 1.00–3.10). For patients aged 70 to 75, the rates of CR + CRi were 43.9 percent in the VYXEOS™ arm and 27.8 percent in the “7+3” free drug dosing arm. The applicant asserted that VYXEOS™ treatment will enable high-risk, older patients (60 years of age and older) to bridge to allogeneic transplant, and VYXEOS™ responding patients will have markedly better outcomes following transplant. The applicant stated that diagnoses of secondary AML are considered incurable with standard chemotherapy approaches and, as with other high-risk hematological malignancies, transplantation is a useful treatment alternative. The applicant further stated that autologous HSCT has limited effectiveness and at this time, only allogeneic HSCT with full intensity conditioning has been reported to produce long-term remissions. However, the applicant stated that the clinical study by Medeiros et al., 2015, reported that, while the use of allogeneic HSCT is considered a potential cure for AML, its use is limited in older patients because of significant baseline comorbidities and increased transplant-related morbidity and mortality. Patients in either arm of the Phase III Study 301 responding to induction with a CR or CR+CRi (n=125) were considered for allogeneic hematopoietic cell transplant (HCT) when possible. In total, 91 patients were transplanted: 52 (34 percent) from the VYXEOS™ arm and 39 (25 percent) from the “7+3” free drug dosing arm. Patient and AML characteristics were similar according to randomized arm, including percentage of patients in each arm that underwent transplant in CR+CRi status. However, the applicant noted that the VYXEOS™ arm contained a higher percentage of older patients (aged 70 or greater) who were transplanted (VYXEOS™, 31 percent; “7+3” free drug dosing, 15 percent).33 According to the applicant, patient outcome following transplant strongly favored patients in the VYXEOS™ arm. The Kaplan-Meier analysis of the 91 transplanted patients showed that the VYXEOS™ arm had markedly better overall survival (hazard ratio 0.46; p=0.0046). The time-dependent Adjustment Model (Cox proportional hazard ratio) was used to evaluate the contribution of VYXEOS™ to overall survival rate after adjusting for transplant and showed that VYXEOS™ remained a significant contributor, even after adjusting for transplant. The time-dependent Adjustment Model (Cox proportional hazard ratio) was used to evaluate the contribution of VYXEOS™ to overall survival rate after adjusting for transplant. The time-dependent Adjustment Model (Cox proportional hazard ratio) was used to evaluate the contribution of VYXEOS™ to overall survival rate after adjusting for transplant and showed that VYXEOS™ remained a significant contributor, even after adjusting for transplant.

The applicant cited Gordon et al., 2016,34 which reported on the significant anti-leukemic activity of VYXEOS™ in AML blasts exhibiting high-risk characteristics, including FLT3–ITD, that are typically associated with poor outcomes when treated with conventional “7+3” free drug dosing. To determine whether the improved complete remission and overall survival rates of VYXEOS™ as compared to conventional “7+3” free drug dosing are attributable to liposome-mediated altered drug PK or direct cellular interactions with specific AML blast samples, the authors evaluated cytotoxicity in 53 AML patient specimens. Cytotoxicity results were correlated with patient characteristics.

as well as VYXEOS™ cellular uptake and molecular phenotype status including FLT3-ITD, which is a predictor of poor patient outcomes to conventional “7+3” free drug dosing. The applicant stated that a notable result from this research was the observation that AML blasts exhibiting the FLT3-ITD phenotype exhibited some of the lowest IC₅₀ (the 50 percent inhibitory concentration) values and, as a group, were five-fold more sensitive to VYXEOS™ than those with wild type FLT3. In addition, there was evidence that increased sensitivity to VYXEOS™ was associated with increased uptake of the drug-laden liposomes by the patient-derived AML blasts. The applicant noted that Gordon, et al. 2016, concluded taken together, the data are consistent with clinical observations where VYXEOS™ retains significant anti-leukemic activity in AML patients exhibiting high-risk characteristics. The applicant also noted that a sub analysis of Phase III Study 301 identified 22 patients diagnosed with FLT3 mutation in the VYXEOS™ arm and 20 in the “7+3” free drug dosing arm, which resulted in the following response rates of FLT3 mutated patients, which were higher with VYXEOS™ (15 of 22, 68.2 percent) versus “7+3” free drug dosing (5 of 20, 25.0 percent); and the Kaplan-Meier analysis of the 42 FLT3 mutated patients showed that patients in the VYXEOS™ arm had a trend towards better overall survival rates (hazard ratio 0.57; p=0.093).

- The applicant asserted that younger patients (18 to 65 years of age) with poor risk first relapse AML have shown higher response rates with VYXEOS™ versus conventional “salvage” chemotherapy. Overall, the applicant stated that the use of VYXEOS™ had an acceptable safety profile in this patient population based on 60-day mortality data. Study 205 was a randomized study comparing VYXEOS™ against the investigator’s choice of first “salvage” chemotherapy in patients diagnosed with relapsed AML after a first remission lasting greater than 1 month (VYXEOS™ arm, n=81 and “7+3” free drug dosing arm, n=44; ages 18 to 65 year of age). Investigator’s choice was almost always based on cytarabine + anthracyclines, usually with the addition of one or two new agents. According to the applicant, VYXEOS™ demonstrated a higher rate of morphological leukemia clearance among all patients, 43.2 percent versus 40.0 percent, and the advantage was most apparent in poor-risk patients, 78.7 percent versus 44.4 percent, as defined by the European Prognostic Index (EPI). In the subset analysis of this EPI poor-risk patient subset, the applicant stated there was a significant improvement in survival rate (6.6 versus 4.2 months median, hazard ratio=0.55, p=0.02) and improved response rate (39.3 percent versus 27 percent). The applicant also noted the following; the safety profile for the use of VYXEOS™ was qualitatively similar to that of control “salvage” therapy, with nearly identical 60-day mortality rates (14.8 percent versus 15.9 percent); among VYXEOS™ treated patients, those with no history of prior HSCT (n=59) had higher response rates (54.2 percent versus 37.8 percent) and lower 60-day mortality (10.2 percent versus 16.2 percent); overall, the use of VYXEOS™ had acceptable safety based on 60-day mortality data, with somewhat higher frequency of neutropenia and thrombocytopenia-related grade 3–4 adverse events. Even though these patients are younger (18 to 65 years of age) than the population studied in Phase III Study 301 (60 years and older), Study 205 patients were at a later stage of disease and almost all had responded to first-line therapy (cytarabine + anthracyclines) and had relapsed. The applicant also cited Cortes, et al. 2015, which reported that patients diagnosed with first relapse AML have limited likelihood of response and short expected survival following “salvage” treatment with the results from showing that:

* Mitoxantrone, etopoide, and cytarabine induced response in 23 percent of patients, with median overall survival of only 2 months.
* Modulation of deoxycytidine kinase by fludarabine led to the combination of fludarabine and cytarabine, resulting in a 36 percent CR rate with median remission duration of 39 weeks.
* First salvage gemtuzumab ozogamicin induced CR+CRp (or CR+CRi) response in 30 percent of patients with CD3+ AML and, for patients with short first CR durations, appeared to be superior to cytarabine-based therapy.

The applicant noted that Study 205 results showed the use of VYXEOS™ retained greater anti-leukemic efficacy in patients diagnosed with poor-risk first relapse AML, and produced higher morphological leukemia clearance rates (78.7 percent) compared to conventional “salvage” therapy (44 percent). The applicant further noted that, overall, the use of VYXEOS™ had acceptable safety profile in this patient population based on 60-day mortality data.

Based on all of the data presented above, the applicant concluded that VYXEOS™ represents a substantial clinical improvement over existing technologies. However, we are concerned that, although there was an improvement in a number of outcomes in Phase III Study 301, specifically overall survival rate, lower risk of early death, improved response rates, better outcomes following transplant, increased response rate and overall survival in patients diagnosed with FLT3 mutation, and higher response rates versus conventional “salvage” chemotherapy in younger patients diagnosed with poor-risk first relapse, the improved outcomes may not be statistically significant. Furthermore, we are concerned that the overall improvement in survival from 5.95 months to 9.56 months may not represent a substantial clinical improvement. In addition, the rate of adverse events in both arms of Study 205, given the theoretical benefit of reduced toxicity with the liposomal formulation, was similar for both the VYXEOS™ and “7+3” free drug treatment groups. Therefore, we also are concerned that there is a similar rate of adverse events, such as febrile neutropenia (68 percent versus 71 percent), pneumonia (20 percent versus 19 percent), and hypoxia (13 percent versus 15 percent), with the use of VYXEOS™ as compared with the conventional “7+3” free drug regimen.

We are inviting public comments on whether the VYXEOS™ meets the substantial clinical improvement criterion.

Below we summarize and respond to comments submitted on VYXEOS™ during the open comment period in response to the New Technology Town Hall meeting notice.

Comment: The applicant provided a written response regarding the definition of “free drug” as “Unbound drug pharmacology;” an active drug or other compound that is not bound to a carrier protein—for example, albumin or alpha-1-acid glycoprotein. The applicant explained that the term “free drug dosing” is used to describe the two different non-encapsulated, separately administered drugs in the “7+3” free drug regimen (cytarabine and daunorubicin), each an unrestricted uniform aqueous solution of the drug in water for continuous administration of cytarabine and separate intravenous dosing.
administration of daunorubicin according to the "7+3" dosing schedule. The applicant then stated that the fixed molar drug ratio delivered by VYXEOS™ is not relevant to the conventional dosing of the two free drugs, cytarabine and daunorubicin. The applicant explained that the doses of cytarabine and daunorubicin used in the conventional "7+3" free drug dosing regimen were based on the maximum tolerated dose of the two agents, not on any concept related to a drug ratio that provides optimal synergy. Finally, the ratio of cytarabine and daunorubicin administered in free (non-liposomal) form is irrelevant because the administered ratio cannot be maintained when these drugs are infused separately. This is because the drugs will be distributed and eliminated differentially and independently of one another and the ratio will change rapidly and continuously. Consequently, according to the applicant, the inability to control drug ratios following administration in conventional dosage forms likely results in exposure of tumor cells to antagonistic drug ratios with a corresponding loss of therapeutic activity.

Response: We appreciate the applicant’s comments. We will take these comments into consideration when deciding whether to approve new technology add-on payments for VYXEOS™.

f. GammaTile™

Isoray Medical, Inc. & GammaTile, LLC submitted an application for a new technology add-on payments for FY 2018 for the GammaTile™. The GammaTile™ is a brachytherapy technology for use in the treatment of patients diagnosed with brain tumors using cesium-131 radioactive sources embedded in a collagen matrix. GammaTile™ is designed to provide adjuvant radiation therapy to eliminate remaining tumor cells in patients who required surgical resection of brain tumors. According to the applicant, the GammaTile™ is a new vehicle of delivery for and inclusive of cesium-131 brachytherapy sources embedded within the product. The applicant stated that the technology has been manufactured for use in the setting of a craniotomy resection site where there is a high chance of local recurrence of a CNS or dual-based tumor. The applicant asserted that the use of GammaTile™ provides a new, unique modality for treating patients who require radiation therapy to augment surgical resection of malignancies of the brain. By offsetting the radiation sources with a 3mm gap of a collagen matrix, the applicant asserted that the use of GammaTile™ resolves issues with “hot” and “cold” spots associated with brachytherapy, improves safety, and potentially offers a treatment option for patients with limited, or no other, available options. The GammaTile™ is biocompatible and bioabsorbable, and is left in the body permanently without need for future surgical removal. The applicant asserted that the commercial manufacturing of the product will significantly improve on the process of constructing customized implants with greater speed, efficiency, and accuracy than is currently available, and require loss surgical expertise in placement of the radioactive sources, allowing a greater number of surgeons to utilize brachytherapy techniques in a wider variety of hospital settings.

The applicant for GammaTile™ has applied for FDA approval and anticipated FDA approval by the spring of 2017. In its application, the applicant indicated that it anticipated that the product would be approved by the FDA for use in both the primary and salvage treatment of radiosensitive malignances of the brain. However, the applicant had not received FDA approval at the time of development of this proposed rule. In subsequent discussions with the applicant, the applicant indicated that it is only seeking FDA approval for use in the salvage treatment of recurrent radiosensitive malignances of the brain. The applicant submitted a request for a unique ICD–10–PCS code for the administration of GammaTile™. If approved, the effective dates will be effective October 1, 2017 (FY 2018).

As discussed earlier, if a technology meets all three of the substantial similarity criteria, it would be considered substantially similar to an existing technology and would not be considered “new” for purposes of new technology add-on payments.

With regard to the first criterion, whether a product uses the same or a similar mechanism of action to achieve a therapeutic outcome, the applicant stated that when compared to treatment using external beam radiation therapy, GammaTile™ uses a new and unique mechanism of action to achieve a therapeutic outcome. The applicant explained that the GammaTile™ is fundamentally different in structure, function, and safety from all external beam radiation therapies, and delivers treatment through a different mechanism of action. In contrast to external beam radiation modalities, the applicant further explained that the GammaTile™ can deliver therapeutic radiation termed brachytherapy. Brachytherapy treatments are performed using radiation sources positioned very close to the area requiring radiation treatment and only deliver radiation to the tissues that are immediately adjacent to the margin of the surgical resection. For this reason, brachytherapy is a current standard of care treatment for many non-central nervous system tumors, including breast, cervical, and prostate cancers. Due to the custom positioning of the radiological sources and the use of the cesium-131 isotope, the applicant noted that the GammaTile™ can target the margin of the excision with greater precision than any alternative treatment option, while sparing healthy brain tissue from unnecessary and potentially damaging radiation exposure.

The applicant also stated that, when compared to other types of brain brachytherapy, GammaTile™ uses a new and unique mechanism of action to achieve a therapeutic outcome. The applicant explained that cancerous cells at the margins of a tumor resection cavity can also be irradiated with the placement of brachytherapy sources in the tumor cavity. However, the applicant asserted that the GammaTile™ is a pioneering form of brachytherapy for the treatment of brain tumors that uses the isotope cesium-131 embedded in a collagen implant that is customized to the geometry of the brain cavity. According to the applicant, use of cesium-131 and the custom distribution of seeds in a three-dimensional collagen device result in a unique and highly effective delivery of radiation therapy to brain tissue.

With regard to the second criterion, whether a product is assigned to the same or a different MS–DRG, GammaTile™ is a treatment option for patients diagnosed with brain tumors that progress locally after initial treatment with external beam radiation therapy, and cases representing patients that may be eligible for treatment involving this technology are assigned to the same MS–DRGs. MS–DRGs 25, 26, and 27 (MS–DRGs 25, 26, and 27 (Craniotomy & Endovascular Intracranial Procedure with MCC, with CC, and without CC/MCC), respectively)
as other current treatment forms of brachytherapy and external beam radiation therapy.

With regard to third criterion, whether the new use of the technology involves the treatment of the same or similar type of disease and the same or similar patient population, the applicant stated that the GammaTile™ offers a treatment option for a patient population with limited, or no other, available treatment options. The applicant explained that treatment options for patients diagnosed with brain tumors that progress locally after initial treatment with external beam radiation therapy are limited, and there is no current standard of care in this setting. According to the applicant, surgery alone for recurrent tumors may provide symptom relief, but does not remove all of the cancer cells. The applicant further stated that repeating external beam radiation therapy for adjuvant treatment is hampered by an increasing risk of brain injury because additional external beam radiation therapy will increase the total dose of radiation to brain tissue, as well as increase the total volume of irradiated brain tissue. Secondary treatment with external beam radiation therapy is often performed with a reduced and, therefore, less effective dose. The applicant asserted that brachytherapy with GammaTile™ may be the only effective treatment option for these patients.

Based on the above, the applicant concluded that the GammaTile™ is not substantially similar to other existing technologies and meets the newness criterion. However, we are concerned that the mechanism of action for this device may be the same or similar to current forms of radiation or brachytherapy. Specifically, while the placement of the cesium-131 source (or any radioactive source) in a collagen matrix offset may constitute a new delivery vehicle, we are concerned that this sort of improvement in brachytherapy for use in the salvage treatment of radiosensitive malignancies of the brain may not represent a new mechanism of action. We also have concerns as to whether GammaTile™ would represent the first approved use of offset radioactive material in brachytherapy for recurrent brain malignancies. The applicant cited studies that used a similar predicate device, but did not indicate whether these researchers or institutions are seeking separate FDA approval.

We are inviting public comments on whether GammaTile™ meets the substantial similarity criteria and the newness criterion.

With regard to the cost criterion, the applicant conducted the following analysis. The applicant worked with the Barrow Neurological Institute at St. Joseph’s Hospital and Medical Center (St. Joseph’s) to obtain actual claims for cranioanatomies using a prototype brain brachytherapy device of stranded cesium-131 seeds held in place with a collagen tile. The application found a total of 23 claims from FY 2001 through FY 2016 data that used a cesium-131 brachytherapy predicate device. All 23 claims were assigned to MS–DRGs 25 through 27. Of the 23 cases, 13 cases were assigned to MS–DRG 25, 4 cases were assigned to MS–DRG 26, and 6 cases were assigned to MS–DRG 27. Using hospital data, the applicant estimated and then subtracted all charges for the predicate device and all charges for ancillary services associated with the device delivery for each case. The applicant standardized the remaining charges for each case and inflated each case’s charges by applying the FY 2017 IPPS/LTC FFS final rule outlier charge inflation factor of 1.043957 by the age of each case. The factor was applied to FY 2011 claims six times, to FY 2012 claims five times, etc., the applicant then calculated the average inflated standardized charges for the cases assigned to MS–DRG 25 ($124,064), MS–DRG 26 ($131,677) and MS–DRG 27 ($90,615). The applicant then calculated an estimate for ancillary charges associated with placement of the GammaTile™ device, as well as standardized charges for the GammaTile™ device. The applicant determined it meets the cost criterion because the final average case-weighted standardized charge per case (including the charges associated with the GammaTile™ device) of $226,741 exceeds the average case-weighted threshold amount of $95,783.

We are concerned that the applicant submitted a small sample of cases to determine it meets the cost criterion. A small sample size may not be statistically significant to determine if the GammaTile™ meets the cost criterion. We also note that, while the applicant has attributed reduced operating room times as a significant benefit to the GammaTile™, a reduction in the associated costs does not appear to be reflected in its calculations. We are inviting public comments on whether the GammaTile™ meets the cost criterion.

With regard to substantial clinical improvement, the applicant stated that the GammaTile™ offers a treatment option for a patient population unresponsive to, or ineligible for, currently available treatments and significantly improves clinical outcomes when compared to currently available treatment options. The applicant explained that therapeutic options for patients diagnosed with large or recurrent brain metastases are limited. However, according to the applicant, the GammaTile™ provides a treatment option for patients diagnosed with radiosensitive recurrent brain tumors that are not eligible for treatment with any other currently available treatment option. Specifically, the applicant stated that GammaTile™ may provide the only radiation treatment option for patients diagnosed with tumors located close to sensitive vital brain sites (for example, brain stem); patients diagnosed with recurrent brain tumors may not be eligible for additional treatment involving the use of external beam radiation therapy. There is a lifetime limit for the amount of radiation therapy a specific area of the body can receive. Patients whose previous treatment includes external beam radiation therapy may be precluded from receiving high doses of radiation associated with subsequent external beam radiation therapy, and the GammaTile™ can also be used to treat tumors that are too large for treatment with external beam radiation therapy. These large tumors are not eligible for treatment with external beam radiation therapy because the radiation dose to healthy brain tissue would be too high.

The applicant described how the GammaTile™ improves clinical outcomes compared to other existing treatment options, including external beam radiation therapy and other forms of brain brachytherapy. To demonstrate that the GammaTile™ represents a substantial clinical improvement over existing technologies, the applicant submitted data from three abstracts, with one associated paper demonstrating feasibility or superior progression-free survival compared to the patient’s own historical control rate. In a presentation at the Society for Neuro-Oncology in November 2014 (Dardis, Christopher; Surgery and permanent intraoperative brachytherapy improves time to progression of recurrent intracranial neoplasms), the outcomes of 20 patients diagnosed with 27 tumors covering a variety of histological types treated with the GammaTile™ prototype were presented. The applicant noted the following with regard to the patients: (1) All tumors were intracranial, supratentorial masses and included low and high-grade meningiomas, metastases from various primary cancers, high-grade gliomas, and others;
(2) all treated masses were recurrent following treatment with surgery and or radiation and the group averaged two prior craniotomies and two prior courses of external beam radiation treatment; and (3) following surgical excision, prototype GammaTilesTM were placed in the resection cavity to deliver a dose of 60 Gray to a depth of 5 mm of tissue; and all patients had previously experienced re-growth of their tumors at the site of treatment and the local control rate of patients entering the study was 0 percent.

With regard to outcomes, the applicant stated that, after their initial treatment, patients had a median progression-free survival time of 5.8 months; post treatment with prototype GammaTilesTM, at the time of this analysis, only one patient had progressed at the treatment site, for a local control rate of 96 percent; and median progression-free survival time, a measure of how long a patient lives without recurrence of the treated tumor, has not been reached (as this value can only be calculated when more than 50 percent of treated patients have failed the prescribed treatment).

A second set of outcomes on prototype GammaTilesTM was presented at the Society for Neuro-Oncology Conference on Meningioma in June 2016 (Brachman, David; Surgery and permanent intraoperative brachytherapy improves time to progress of recurrent intracranial neoplasms). This study enrolled 16 patients with 20 recurrent grade 2 or 3 meningiomas, who had undergone prior surgical excision for re-operation compared to external beam radiation therapy. These patients underwent surgical excision of the tumor, followed by adjuvant radiation therapy with prototype GammaTilesTM. The applicant noted the following outcomes: (1) Of the 20 treated tumors, 19 showed no evidence of radiographic progression at last follow-up, yielding a local control rate of 95 percent; two of the 20 patients exhibited radiation necrosis (one asymptomatic, one asymptomatic); and (2) the median time to failure from the prior treatment with external beam radiation therapy was 10.3 months and after treatment with prototype GammaTilesTM only one patient failed at 18.2 months. Therefore, the median time to same site failure after prototype GammaTilesTM treatment has not yet been reached (average follow up of 16.7 months, range 1–37 months).

A third prospective study was accepted for presentation at the November 2016 Society for Neuro-Oncology annual meeting (Youssef, Emad; Cs131 implants for salvage therapy of recurrent high grade gliomas). In this study, 13 patients diagnosed with recurrent high-grade gliomas (9 with glioblastoma and 4 with grade 3 astrocytoma) were treated in an identical manner to the cases described above. Previously, all patients had failed the international standard treatment for high-grade glioma, a combination of surgery, radiation therapy, and chemotherapy referred to as the “Stupp regimen.” For the prior therapy, the median time to failure was 9.2 months (range 1–40 months). After therapy with a prototype GammaTilesTM, the applicant noted the following: (1) The median time to same site local failure has not been reached and one failure was seen at 18 months (local control 92 percent); and (2) with a median follow-up time of 8.1 months (range 1–23 months) one symptomatic patient (8 percent) and two asymptomatic patients (15 percent) had radiation-related MRI changes. However, no patients required re-operation for radiation necrosis or wound breakdown.

The applicant asserted that, when compared in total, the data reported in these three studies support the conclusion that a significant therapeutic effect results from the addition of GammaTilesTM radiation therapy to the site of surgical removal. According to the applicant, the fact that these patients had failed prior best available treatments (aggressive surgical and adjuvant radiation management) presents the unusual scenario of a salvage therapy outperforming the current standard-of-care. The applicant noted that follow-up data continues to accrue on these patients. The applicant further noted that, although these reported experiences with the GammaTilesTM are as a salvage therapy in patients who currently have no standard treatment options, it is anticipated GammaTilesTM will also be used as first-line therapy due to these promising results.

The applicant stated that the use of GammaTilesTM reduces rates of mortality compared to alternative treatment options. The applicant explained that the rate of symptomatic radiation necrosis in the GammaTilesTM clinical studies of 5 to 8 percent is substantially lower than the 26 percent to 57 percent rate of symptomatic radiation necrosis requiring re-operation historically associated with brain brachytherapy, and lower than the rates reported for initial treatment of similar tumors with modern external beam and stereotactic radiation techniques. The applicant indicated that this is consistent with the customized and ideal distribution of radiation therapy provided by GammaTilesTM.

The applicant also asserted that the use of GammaTilesTM reduces the need for re-operation compared to alternative treatment options. The applicant explained that patients receiving a craniotomy, followed by external beam radiation therapy or brachytherapy, could require re-operation in the following three scenarios:

- Tumor recurrence at the excision site could require additional surgical removal;
- Symptomatic radiation necrosis could require excision of the affected tissue; and
- Certain forms of brain brachytherapy require the removal of brachytherapy sources after a given period of time.

However, according to the applicant, because of the high local control rates, low rates of symptomatic radiation necrosis, and short half-life of cesium-131, GammaTilesTM will reduce the need for re-operation compared to external beam radiation therapy and other forms of brain brachytherapy.

Additionally, the applicant stated that the use of GammaTilesTM reduces the need for additional hospital visits and procedures compared to alternative treatment options. The applicant noted that the GammaTilesTM is placed during surgery, and does not require any additional visits or procedures. The applicant contrasted this improvement with external beam radiation therapy, which is often delivered in multiple fractions that must be administered over multiple days. The applicant provided an example where WBRT is delivered over 2 to 3 weeks, while the placement of GammaTilesTM occurs during the craniotomy and does not add any time to a patient’s recovery.

The applicant further stated that the GammaTilesTM has high local control rates and low rates of symptomatic radiation necrosis will reduce the need for...
additional hospital visits and procedures, and provides a more rapid initiation and complement of the treatment compared to alternative treatment options.

Based on consideration of all of the data presented above, the applicant believed that the use of GammaTile™ represents a substantial clinical improvement over existing technologies. The studies were limited to patients diagnosed with recurrent tumors after previous surgical rescission. As previously discussed, the applicant explained that it is seeking FDA approval for the use of the GammaTile™ in the treatment of recurrent malignancies.

We are inviting public comments on whether GammaTile™ meets the substantial clinical improvement criterion.

We did not receive any written public comments in response to the New Technology Town Hall meeting notice regarding the application of GammaTile™ for new technology add-on payments.

III. Proposed Changes to the Hospital Wage Index for Acute Care Hospitals

A. Background

1. Legislative Authority

Section 1886(d)(3)(E) of the Act requires that, as part of the methodology for determining prospective payments to hospitals, the Secretary adjust the standardized amounts for area differences in hospital wage levels by a factor (established by the Secretary) reflecting the relative hospital wage level in the geographic area of the hospital compared to the national average hospital wage level. We currently define hospital labor market areas based on the delineations of statistical areas established by the Office of Management and Budget (OMB). A discussion of the proposed FY 2018 hospital wage index based on the statistical areas appears under sections III.A.2. and C. of the preamble of this proposed rule.

Section 1886(d)(3)(E) of the Act requires the Secretary to update the wage index annually and to base the update on a survey of wages and wage-related costs of short-term, acute care hospitals. (CMS collects these data on the Medicare cost report, CMS Form 2552–10, Worksheet S–3, Parts II, III, and IV. The OMB control number for approved collection of this information is 0938–0050.) This provision also requires that any updates or adjustments to the wage index be made in a manner that ensures that aggregate payments to hospitals are not affected by the change in the wage index. The proposed adjustment for FY 2018 is discussed in section II.B. of the Addendum to this proposed rule.

As discussed in section III.J. of the preamble of this proposed rule, we also take into account the geographic reclassification of hospitals in accordance with sections 1886(d)(8)(B) and 1886(d)(10) of the Act when calculating IPPS payment amounts.

Under section 1886(d)(8)(D) of the Act, the Secretary is required to adjust the standardized amounts so as to ensure that aggregate payments under the IPPS after implementation of the provisions of sections 1886(d)(6)(B), 1886(d)(8)(C), and 1886(d)(10) of the Act are equal to the aggregate prospective payments that would have been made absent these provisions. The proposed budget neutrality adjustment for FY 2018 is discussed in section II.A.4.b. of the Addendum to this proposed rule.

Section 1886(d)(3)(E) of the Act also provides for the collection of data every 3 years on the occupational mix of employees for short-term, acute care hospitals participating in the Medicare program, in order to construct an occupational mix adjustment to the wage index. A discussion of the occupational mix adjustment that we are proposing to apply to the FY 2018 wage index, appears under sections III.E.3. and F. of the preamble of this proposed rule.

2. Core-Based Statistical Areas (CBSAs) for the Proposed FY 2018 Hospital Wage Index

The wage index is calculated and assigned to hospitals on the basis of the labor market area in which the hospital is located. Under section 1886(d)(3)(E) of the Act, beginning with FY 2005, we delineate hospital labor market areas based on OMB-established Core-Based Statistical Areas (CBSAs). The current statistical areas (which were implemented beginning with FY 2015) are based on revised OMB delineations issued on February 28, 2013, in OMB Bulletin No. 13–01. OMB Bulletin No. 13–01 established revised delineations for Metropolitan Statistical Areas, Micropolitan Statistical Areas, and Combined Statistical Areas in the United States and Puerto Rico based on the 2010 Census, and provided guidance on the use of the delineations of these statistical areas using standards published on June 28, 2010 in the Federal Register (75 FR 37246 through 37252). We refer readers to the FY 2015 IPPS Final Rule (80 FR 49957 through 49963) for a full discussion of our implementation of the OMB labor market area delineations beginning with the FY 2015 wage index.

Generally, OMB issues major revisions to statistical areas every 10 years, based on the results of the decennial census. However, OMB occasionally issues minor updates and revisions to statistical areas in the years between the decennial censuses through OMB Bulletins. On July 15, 2015, OMB issued OMB Bulletin No. 15–01, which provides updates to and supersedes OMB Bulletin No. 13–01 that was issued on February 28, 2013. The attachment to OMB Bulletin No. 15–01 provides detailed information on the update to statistical areas since February 28, 2013. The updates provided in OMB Bulletin No. 15–01 are based on the application of the 2010 Standards for Delineating Metropolitan and Micropolitan Statistical Areas to Census Bureau population estimates for January 1, 2012 and July 1, 2013. In the FY 2017 IPPS/LTCH PPS final rule (81 FR 56913), we adopted the updates set forth in OMB Bulletin No. 15–01 effective October 1, 2016, beginning with the FY 2017 wage index. For a complete discussion of the adoption of the updates set forth in OMB Bulletin No. 15–01, we refer readers to the FY 2017 IPPS/LTCH PPS final rule.

For FY 2018, we are continuing to use the OMB delineations that we adopted beginning with FY 2015 to calculate the area wage indexes, with updates as reflected in OMB Bulletin No. 15–01 specified in the FY 2017 IPPS/LTCH PPS final rule.

3. Codes for Constituent Counties in CBSAs

CBSAs are made up of one or more constituent counties. Each CBSA and constituent county has its own unique identifying codes. There are two different lists of codes associated with counties: Social Security Administration (SSA) codes and Federal Information Processing Standard (FIPS) codes. Historically, CMS has listed and used SSA and FIPS county codes to identify and crosswalk counties to CBSA codes for purposes of the hospital wage index. We have learned that SSA county codes are no longer being maintained and updated. However, the FIPS codes continue to be maintained by the U.S. Census Bureau. The Census Bureau’s most current statistical area information is derived from ongoing census data received since 2010; the most recent data are from 2015. For the purposes of crosswalking counties to CBSAs, we are proposing to continue the use of SSA county codes and begin using only the FIPS county codes.
The Census Bureau maintains a complete list of changes to counties or county equivalent entities on the Web site at: https://www.census.gov/geo/reference/county-changes.html. In our proposed transition to using only FIPS codes for counties for the hospital wage index, we are proposing to update the FIPS codes used for crosswalking counties to CBSAs for the hospital wage index to incorporate changes to the counties or county equivalent entities included in the Census Bureau’s most recent list. Based on information included in the Census Bureau’s Web site, since 2010, the Census Bureau has made the following updates to the FIPS codes for counties or county equivalent entities:

- Petersburgh Borough, AK (FIPS State County Code 02–195), CBSA 02, was created from part of former Petersburgh Census Area (02–195) and part of Hoonah-Angoon Census Area (02–105). The CBSA code remains 02.
- The name of La Salle Parish, LA (FIPS State County Code 22–059), CBSA 14, is now LaSalle Parish, LA (FIPS State County Code 22–059). The CBSA code remains as 14.
- The name of Shannon County, SD (FIPS State County Code 46–113), CBSA 43, is now Oglala Lakota County, SD (FIPS State County Code 46–102). The CBSA code remains as 43.

We believe that it is important to use the latest counties or county equivalent entities in order to properly crosswalk hospitals from a county to a CBSA for purposes of the hospital wage index used under the IPPS. In addition, we believe that using the latest FIPS codes will allow us to maintain a more accurate and up-to-date payment system that reflects the reality of population shifts and labor market conditions. Therefore, we are proposing to implement these FIPS code updates, effective October 1, 2017, beginning with the FY 2018 wage indexes. We are proposing to use these update changes to calculate area wage indexes in a manner that is generally consistent with the CBSA-based methodologies finalized in the FY 2005 IPPS final rule and the FY 2015 IPPS/LTCH PPS final rule. We note that while the county update changes listed earlier changed the county names, the CBSAs to which these counties map did not change from the prior counties. Therefore, there is no impact or change to hospitals in these counties; they continue to be considered rural for the hospital wage index under these changes. For FY 2018, Tables 2 and 3 associated with this proposed rule and the County to CBSA Crosswalk File and Urban CBSAs and Constituent Counties for Acute Care Hospitals File posted on the CMS Web site reflect these county changes. We are inviting public comments on our proposals.

**B. Worksheet S–3 Wage Data for the Proposed FY 2018 Wage Index**

The proposed FY 2018 wage index values are based on the data collected from the Medicare cost reports submitted by hospitals for cost reporting periods beginning in FY 2014 (the FY 2017 wage indexes were based on data from cost reporting periods beginning during FY 2013).

1. Included Categories of Costs

The proposed FY 2018 wage index includes all of the following categories of data associated with costs paid under the IPPS (as well as outpatient costs):

- Salaries and hours from short-term, acute care hospitals (including paid lunch hours and hours associated with military leave and jury duty);
- Home office costs and hours;
- Certain contract labor costs and hours, which include direct patient care, certain top management, pharmacy, laboratory, and non-teaching physician Part A services, and certain contract indirect patient care services (as discussed in the FY 2008 final rule with comment period (72 FR 47315 through 47317)); and
- Wage-related costs, including pension costs (based on policies adopted in the FY 2012 IPPS/LTCH PPS final rule (76 FR 51586 through 51590)) and other deferred compensation costs.

2. Excluded Categories of Costs

Consistent with the wage index methodology for FY 2017, the proposed wage index for FY 2018 also excludes the direct and overhead salaries and hours for services not subject to IPPS payment, such as skilled nursing facility (SNF) services, home health services, costs related to GME (teaching physicians and residents) and certified registered nurse anesthetists (CRNAs), and other subprovider components that are not paid under the IPPS. The proposed FY 2018 wage index also excludes the salaries, hours, and wage-related costs of hospital-based rural health clinics (RHCs), and Federally qualified health centers (FQHCs) because Medicare pays for these costs outside of the IPPS (68 FR 45385). In addition, salaries, hours, and wage-related costs of CAHs are excluded from the wage index for the reasons explained in the FY 2004 IPPS final rule (68 FR 45397 through 45398).

3. Use of Wage Index Data by Suppliers and Providers Other Than Acute Care Hospitals Under the IPPS

Data collected for the IPPS wage index also are currently used to calculate wage indexes applicable to suppliers and other providers, such as SNFs, home health agencies (HHAs), ambulatory surgical centers (ASCs), and hospices. In addition, they are used for prospective payments to IRFs, IPFs, and LTCHs, and for hospital outpatient services. We note that, in the IPPS rules, we do not address comments pertaining to the wage indexes of any supplier or provider except IPPS providers and LTCHs. Such comments should be made in response to separate proposed rules for those suppliers and providers.

**C. Verification of Worksheet S–3 Wage Data**

The wage data for the proposed FY 2018 wage index were obtained from Worksheet S–3, Parts II and III of the Medicare cost report (Form CMS–2552–10) for cost reporting periods beginning on or after October 1, 2013, and before October 1, 2014. For wage index purposes, we refer to cost reports during this period as the “FY 2014 cost report,” the “FY 2014 wage data,” or the “FY 2014 data.” Instructions for completing the wage index sections of Worksheet S–3 are included in the Provider Reimbursement Manual (PRM), Part 2 (Pub. No. 15–2), Chapter 40, Sections 4005.2 through 4005.4. The data file used to construct the proposed FY 2018 wage index includes FY 2014 data submitted to us as of February 10, 2017. As in past years, we performed an extensive review of the wage data, mostly through the use of edits designed to identify aberrant data.

We asked our MACs to revise or verify data elements that result in specific edit failures. For the proposed FY 2018 wage index, we identified and excluded 51 providers with aberrant data that should not be included in the wage index, although if data elements for some of these providers are corrected, we intend to include data from those providers in the final FY 2018 wage index. We also adjusted certain aberrant data and included these data in the proposed wage index. For example, in situations where a hospital did not have documentable salaries, wages, and hours for housekeeping and dietary services, we imputed estimates, in accordance with policies established in the FY 2015 IPPS/LTCH PPS final rule (70 FR 49965 through 49967). We instructed MACs to complete their data verification of questionable data elements and to transmit any changes to
the wage data no later than March 24, 2017. The revised data will be reflected in the FY 2018 IPPS/LTCH PPS final rule.

In constructing the proposed FY 2018 wage index, we included the wage data for facilities that were IPPS hospitals in FY 2014, inclusive of those facilities that have since terminated their participation in the program as hospitals, as long as those data did not fail any of our edits for reasonableness. We believed that including the wage data for these hospitals is, in general, appropriate to reflect the economic conditions in the various labor market areas during the relevant past period and to ensure that the current wage index represents the labor market area’s current wages as compared to the national average of wages. However, we excluded the wage data for CAHs as discussed in the FY 2004 IPPS final rule (68 FR 45397 through 45398). For the this proposed rule, we removed 7 hospitals that converted to CAH status on or after January 22, 2016, the cut-off date for CAH exclusion from the FY 2017 wage index, and through and including January 23, 2017, the cut-off date for CAH exclusion from the FY 2018 wage index. After excluding CAHs and hospitals with aberrant data, we calculated the proposed wage index using the Worksheet S–3, Part II and III wage data of 3,325 hospitals.

For the proposed FY 2018 wage index, we allotted the wages and hours data for a multicampus hospital among the different labor market areas where its campuses are located in the same manner that we allotted such hospitals’ data in the FY 2017 wage index (81 FR 56915). Table 2, which contains the proposed FY 2018 wage index associated with proposed rule (available via the Internet on the CMS Web site), includes separate wage data for the campuses of 9 multicampus hospitals.

D. Method for Computing the Proposed FY 2018 Unadjusted Wage Index

1. Proposed Methodology for FY 2018

The method used to compute the proposed FY 2018 wage index without an occupational mix adjustment follows the same methodology that we used to compute the proposed wage indexes without an occupational mix adjustment since FY 2012 (76 FR 51591 through 51593).

As discussed in the FY 2012 IPPS/LTCH PPS final rule, in “Step 5,” for each hospital, we adjust the total salaries plus wage-related costs to a common period to determine total adjusted salaries plus wage-related costs. To make the wage adjustment, we estimate the percentage change in the employment cost index (ECI) for compensation for each 30-day increment from October 14, 2013, through April 15, 2015, for private industry hospital workers from the BLS’s Compensation and Working Conditions. We have consistently used the ECI as the data source for our wages and salaries and other price proxies in the IPPS market basket, and we are not proposing any changes to the usage of the ECI for FY 2018. The factors used to adjust the hospital’s data were based on the midpoint of the cost reporting period, as indicated in the following table.

**Midpoint of Cost Reporting Period**

<table>
<thead>
<tr>
<th>After</th>
<th>Before</th>
<th>Adjustment factor</th>
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<tbody>
<tr>
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<td>1.02310</td>
</tr>
<tr>
<td>11/14/2013</td>
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<td>1.02155</td>
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<td>1.02004</td>
</tr>
<tr>
<td>01/14/2014</td>
<td>02/15/2014</td>
<td>1.01866</td>
</tr>
<tr>
<td>02/14/2014</td>
<td>03/15/2014</td>
<td>1.01740</td>
</tr>
<tr>
<td>03/14/2014</td>
<td>04/15/2014</td>
<td>1.01615</td>
</tr>
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<td>0.99845</td>
</tr>
</tbody>
</table>

For example, the midpoint of a cost reporting period beginning January 1, 2014, and ending December 31, 2014, is June 30, 2014. An adjustment factor of 1.01193 would be applied to the wages of a hospital with such a cost reporting period.

Using the data as previously described, the proposed FY 2018 national average hourly wage (unadjusted for occupational mix) is $42.0043. Previously, we also would provide a Puerto Rico overall average hourly wage. As discussed in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56915), prior to January 1, 2016, Puerto Rico hospitals were paid based on 75 percent of the national standardized amount and 25 percent of the Puerto Rico-specific standardized amount. As a result, we calculated a Puerto Rico-specific wage index that was applied to the labor share of the Puerto Rico-specific standardized amount. Section 601 of the Consolidated Appropriations Act, 2016 (Pub. L. 114–113) amended section 1886(d)(9)(E) of the Act to specify that the payment calculation with respect to operating costs of inpatient hospital services of a subsection (d) Puerto Rico hospital for inpatient hospital discharges on or after January 1, 2016, shall use 100 percent of the national standardized amount. As we stated in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56915 through 56916), because Puerto Rico hospitals are no longer paid with a Puerto Rico-specific standardized amount as of January 1, 2016, under section 1886(d)(9)(E) of the Act, as amended by section 601 of the Consolidated Appropriations Act, 2016, there is no longer a need to calculate a Puerto Rico-specific average hourly wage and wage index. Hospitals in Puerto Rico are now paid 100 percent of the national standardized amount and, therefore, are subject to the national average hourly wage (unadjusted for occupational mix) (which is $42.0043 for this FY 2018 proposed rule) and the national wage index, which is applied to the national labor share of the national standardized amount. For FY 2018, we are not proposing a Puerto Rico-specific overall average hourly wage or wage index.

2. Clarification of Other Wage Related Costs in the Wage Index

Section 1886(d)(3)(E) of the Act requires the Secretary to update the wage index based on a survey of hospitals’ costs that are attributable to wages and wage-related costs. In the September 1, 1994 IPPS final rule (59 FR 45356), we developed a list of “core” wage-related costs that hospitals may report on Worksheet S–3, Part II of the Medicare hospital cost report in order to include those costs in the wage index. Core wage-related costs include categories of retirement cost, plan administrative costs, health and insurance costs, taxes, and other specified costs such as tuition reimbursement. In addition to these categories of core wage-related costs, we allow hospitals to report wage-related costs other than those on the core list if the other wage-related costs meet certain criteria. The criteria for including other wage-related costs in the wage index are discussed in the September 1, 1994 IPPS final rule (59 FR 45357) and also are listed in the Provider Reimbursement Manual (PRM), Part II, Chapter 40, Sections 4005.2 through 4005.4, Line 18 of the Medicare cost report (Form CMS–2552–10, OMB control number 0936–0050).

Specifically, “other” wage-related costs are allowable for the wage index if the cost for employees wages are paid under the IPPS exceeds 1 percent of the total adjusted salaries net of
excluded area salaries, is a fringe benefit as defined by the IRS and has been reported to the IRS (as income to the employees or contractors), is not being furnished for the convenience of the provider, and is not listed on Worksheet S–3, Part IV.

We note that other wage-related costs are not to include benefits already included in Line 1 salaries on Worksheet S–3, Part II (refer to the cost report instructions for Worksheet S–3, Part II, Line 18, which state, “Other wage-related costs do not include wage-related costs reported on line 1 of this worksheet.”). We also note that the 1-percent test is conducted by dividing each individual category of the other wage-related cost (that is, the numerator) by the sum of the following lines on the Medicare hospital cost report (Form CMS–2552–10): Worksheet S–3, Part II, Lines 11, 12, 13, and 14, Column 4, and Worksheet S–3, Part III, Line 3, Column 4 (that is, the denominator). The other wage-related costs associated with contract labor and home office-related organization personnel are included in the numerator because these other wage-related costs are allowed in the wage index (in addition to other wage-related costs for direct employees), assuming the requirements for inclusion in the wage index are met. For example, if a hospital is trying to include a parking garage as an other-wage related cost that is reported on the W–2 or 1099 form, when running the 1-percent test, include in the numerator all the parking garage other-wage related cost for direct salary employees, contracted employees, and home office employees and divide by the sum of Worksheet S–3, Part II, Lines 11, 12, 13, and 14, Column 4, and Worksheet S–3, Part III, Line 3, Column 4. For the category of parking other-wage-related costs, the 1-percent test would be run only one time, inclusive of other wage related costs for employee salaries, contracted employees, and home office employees. We intend to clarify the hospital cost report instructions to reflect that contract labor and home office/related organization salaries should be added to the subtotal of salaries on Worksheet S–3, Part III, Line 3, Column 4 (Line 3 is the difference of net salaries minus excluded area salaries) for purposes of performing the 1-percent test. If a hospital has more than one other wage-related cost (for example, parking and cafeteria separately; do not sum all the different types of other wage-related costs together and then run the 1-percent test). If the 1-percent test is met for a particular type of other wage-related costs, and the other criteria listed earlier are met as well, the other wage-related cost may be reported on Worksheet S–3, Part II, Line 18 of the hospital cost report.

We originally allowed for the inclusion of wage-related costs other than those on the core list because we were concerned that individual hospitals might incur unusually large wage-related costs that are not reflected on the core list but that may represent a significant wage-related cost. However, we are reconsidering allowing other wage-related costs to be included in the wage index because recent internal reviews of the FY 2018 wage data show that only a small minority of hospitals are reporting other wage-related costs that meet the 1-percent test described earlier. In the calculation of the proposed FY 2018 wage index, for each hospital reporting other wage-related costs on Line 18 of Worksheet S–3, we performed the 1-percent test. We then made internal edits removing other wage-related costs on Line 18 where hospitals reported data that failed to meet the mathematical requirement that other wage-related costs must exceed 1 percent of total adjusted salaries net of excluded area salaries. After this review, only approximately 80 hospitals of approximately 3,320 hospitals had other wage-related costs on Line 18 meeting the 1-percent test. We believe that such a limited number of hospitals nationally reporting and meeting the 1-percent test may indicate that other wage-related costs might not constitute an appropriate part of a relative measure of wage costs in a particular labor market area, a longstanding tenet of the wage index. In other words, while other wage-related costs may represent costs that may have an impact on an individual hospital’s average hourly wage, we do not believe that costs reported by only a very small minority of hospitals accurately reflect the economic conditions of the labor market areas in which those hospitals are located. Therefore, it is possible that inclusion of other wage-related costs in the wage index in such a limited manner may distort the average hourly wage of a particular labor market area so that its wage index does not accurately represent that labor market area’s current wages relative to national wages.

Furthermore, the open-ended nature of the types of other wage-related costs that may be included on Line 18 of Worksheet S–3, in contrast to the core wage-related costs, may hinder consistent and proper reporting of fringe benefits. Our internal review indicates widely divergent types of costs that hospitals are reporting as other wage-related costs on Line 18. We are concerned that inconsistent reporting of other wage-related costs on Line 18 further compromises the accuracy of the wage index as a representation of the relative average hourly wage for each labor market area. Our intent in creating a core list of wage-related costs in the September 1, 1994 IPPS final rule was to promote consistent reporting of fringe benefits, and we are increasingly concerned that inconsistent reporting of wage-related costs on Line 18 of Worksheet S–3 undermines this effort. Specifically, we expressed in the September 1, 1994 IPPS final rule that since we began including fringe benefits in the wage index, we have been concerned with the inconsistent reporting of fringe benefits, whether because of a lack of provider proficiency in identifying fringe benefit costs or varying interpretations across fiscal intermediaries of the definition for fringe benefits in PRM–I, Section 2144.1 (59 FR 45356).

We believe that the limited and inconsistent use of Line 18 of Worksheet S–3 for reporting wage-related costs other than the core list might indicate that including other wage-related costs in the wage index compromises the accuracy of the wage index as a relative measure of wages in a given labor market area. Therefore, we are seeking public comments on whether we should, in future rulemaking, propose to only include the wage-related costs on the core list in the calculation of the wage index and not to include any other wage-related costs in the calculation of the wage index.

Meanwhile, in this FY 2018 IPPS/LTCH PPS proposed rule, we are clarifying that, under our current policy, an other-wage-related cost (which we define as the value of a benefit) must be a fringe benefit as described by the IRS (refer to IRS Publication 15–B) and must be reported to the IRS on employees’ or contractors’ W–2 or 1099 forms as taxable income in order to be considered an other-wage-related cost on Line 18 of Worksheet S–3 and for the wage index. That is, other-wage-related costs that are not reported to the IRS on employees’ or contractors’ W–2 or 1099 forms as taxable income, even if not required to be reported to the IRS according to IRS requirements, will not be included in the wage index. This is consistent with current cost report instructions for Line 18 of Worksheet S–3, Part II of the Medicare cost report, Form CMS–2552–10, whereby that, to be considered an allowable other-wage-related costs, the cost “has been
reported to the IRS.” We will apply this policy to the process for calculating the wage index for FY 2019, including the FY 2019 desk reviews beginning in September 2017.

We believe this clarification is necessary because some hospitals have incorrectly interpreted prior manual and existing preamble language to mean that a cost could be considered an other wage-related cost if the provider’s reporting (or not reporting) of the cost was in accordance with IRS requirements, rather than if the cost was actually reported on an employee’s or contractor’s W–2 or 1099 form as taxable income. We believe that such interpretation of our policy would require an analysis of whether the reporting or not reporting of the cost to the IRS was done properly in accordance with IRS regulations and guidance in order to allow the cost as an other wage-related cost. We believe that the determinations regarding the proper or improper reporting of certain other wage-related costs to the IRS for the purposes of inclusion in the Medicare wage index are impractical for CMS and the MACs because we do not have the expertise and fluency in IRS regulations and tax law sufficient to perform such technical reviews of hospital wage-related costs. In contrast, our current policy of including an amount as an other wage-related cost for wage index purposes only if the amount was actually reported to the IRS on employees’ or contractors’ W–2 or 1099 forms as taxable income is a straightforward policy that we believe provides clarity to all involved parties. The brightline test of allowing an other wage-related cost to be included in the wage index only if it has been reported on an employee’s or contractor’s W–2 or 1099 form as taxable income helps ensure consistent treatment of other wage-related costs for all hospitals. Considering the variety of types of costs that may be included on Line 18 of Worksheet S–3 of the Medicare cost report for other wage-related costs (assuming the 1-percent test is met and other criteria are met), we believe that a straightforward policy that is simple for hospitals and CMS to apply is particularly important.

In addition, we believe the policy we are clarifying in this proposed rule, that an other wage-related cost can be included in the wage index only if it was reported to the IRS as taxable income on the employee’s or contractor’s W–2 or 1099, is consistent with CMS’ longstanding position that a fringe benefit is not furnished for the convenience of the employer or otherwise excludable from income as a fringe benefit (such as a working condition fringe) and that inappropriate types of costs may not be included in the wage index. In response to a comment when we finalized the criteria for other wage-related costs in the September 1, 1994 IPPS final rule (59 FR 45359), we stated that “items such as the unrecovered cost of employee meals, tuition reimbursement, and auto allowances will only be allowed as a wage-related cost for purposes of the wage index if properly reported to the IRS on an employee’s W–2 form as a fringe benefit.” (We note that the September, 1 1994 IPPS final rule does not mention the 1099 form for contractors, as contract labor was not allowed at that time in the wage index. Consistent with our treatment of costs for contract labor similar to that of employees for the wage index, we are clarifying that the requirement that a cost be reported to the IRS to be allowed as a wage-related cost for the wage index also applies to contract labor, which must be reported on the contractor’s 1099 to be allowed as a wage-related cost for the wage index.) We believe that requiring other wage-related costs to be reported on employees’ or contractors’ W–2 or 1099 forms to be allowable for Line 18 of Worksheet S–3 of the Medicare cost report is consistent with the requirement that the cost is not being furnished for the convenience of the employer. A cost reported on an employee’s or contractor’s W–2 or 1099 form as taxable income is clearly a wage-related cost that is provided solely for the benefit of the employee. We believe that the requirement that other wage-related costs be a benefit to the employee also guarantees that administrative costs such as overhead and capitalized costs are excluded from other wage-related costs in the wage index.

Therefore, for the reasons discussed above, we are clarifying that a cost must be a fringe benefit as described by the IRS and reported to the IRS on an employee’s or contractor’s W–2 or 1099 form as taxable income.

E. Proposed Occupational Mix Adjustment to the FY 2018 Wage Index

As stated earlier, section 1886(d)(3)(E) of the Act provides for the collection of data every 3 years on the occupational mix of employees for each short-term, acute care hospital participating in the Medicare program, in order to construct an occupational mix adjustment to the wage index, for application beginning October 1, 2004 (the FY 2005 wage index). The purpose of the occupational mix adjustment is to control for the effect of hospitals’ employment choices on the wage index. For example, hospitals may choose to employ different combinations of registered nurses, licensed practical nurses, nursing aides, and medical assistants for the purpose of providing nursing care to their patients. The varying labor costs associated with these choices reflect hospital management decisions rather than geographic differences in the costs of labor.

1. Use of 2013 Occupational Mix Survey for the FY 2018 Wage Index

Section 304(c) of the Consolidated Appropriations Act, 2001 (Pub. L. 106–554) amended section 1886(d)(3)(E) of the Act to require CMS to collect data every 3 years on the occupational mix of employees for each short-term, acute care hospital participating in the
Medicare program. We collected data in 2013 to compute the occupational mix adjustment for the FY 2016, FY 2017, and FY 2018 wage indexes. A new measurement of occupational mix is required for FY 2019.

The 2013 survey included the same data elements and definitions as the previous 2010 survey and provided for the collection of hospital-specific wages and hours data for nursing employees for calendar year 2013 (that is, payroll periods ending between January 1, 2013 and December 31, 2013). We published the 2013 survey in the Federal Register on February 28, 2013 (78 FR 13679 through 13680). This survey was approved by OMB on May 14, 2013, and is available on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Wage-Index-Files-Items/Medicare-Wage-Index-Occupational-Mix-Survey2013.html. The 2013 Occupational Mix Survey Hospital Reporting Form CMS–10079 for the Wage Index Beginning FY 2016 (in Excel format) is available on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Wage-Index-Files-Items/Medicare-Wage-Index-Occupational-Mix-Survey2013.html. Hospitals were required to submit their completed 2013 surveys to their MACs by July 3, 2017.

3. Calculation of the Proposed Occupational Mix Adjustment for FY 2018

For FY 2018, we are proposing to calculate the occupational mix adjustment factor using the same methodology that we have used since the FY 2012 wage index (76 FR 51582 through 51586) and to apply the occupational mix adjustment to 100 percent of the FY 2018 wage index. Because the statute requires that the Secretary measure the earnings and paid hours of employment by occupational category not less than once every 3 years, all hospitals that are subject to payments under the IPPS, or any hospital that would be subject to the IPPS if not greater, must complete the occupational mix survey, unless the hospital has no associated cost report wage data that are included in the FY 2018 wage index. For the proposed FY 2018 wage index, we are using the Worksheet S–3, Parts II and III cost report wage data, which we compared to prior years.

2. Use of the 2016 Medicare Wage Index Occupational Mix Survey for the FY 2019 Wage Index

As stated earlier, a new measurement of occupational mix is required for FY 2019. The FY 2019 occupational mix adjustment will be based on a new calendar year (CY) 2016 survey. The CY 2016 survey (CMS Form CMS–10079) received OMB approval on September 27, 2016. The final CY 2016 Occupational Mix Survey Hospital Reporting Form is available on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Wage-Index-Files-Items/2016-Occupational-Mix-Survey-Hospital-Reporting-Form-CMS–10079-for-the-Wage-Index-Beginning-FY–2019.html. Hospitals are required to submit their completed 2016 surveys to their MACs by July 3, 2017. The preliminary, unaudited CY 2016 survey data will be posted on the CMS Web site in mid-July 2017. As with the Worksheet S–3, Parts II and III cost report wage data, as part of the FY 2019 desk review process, the MACs will revise or verify data elements in hospitals’ occupational mix surveys that result in certain edit failures.

F. Analysis and Implementation of the Proposed Occupational Mix Adjustment and the Proposed FY 2018 Occupational Mix Adjusted Wage Index

As discussed in section III.E. of the preamble of this proposed rule, for FY 2018, we are proposing to apply the occupational mix adjustment to 100 percent of the FY 2018 wage index. We calculated the proposed occupational mix adjustment using data from the 2013 occupational mix survey data, using the methodology described in the FY 2012 IPPS/LTC PPS final rule (76 FR 51582 through 51586) using the occupational mix survey data and applying the occupational mix adjustment to 100 percent of the FY 2017 wage index results in a proposed national average hourly wage of $41.9599.

The proposed FY 2018 average occupational wage index will be posted on the CMS Web site in mid-July 2017.
percent, and no rural areas’ proposed wage index values would increase by 5 percent or more. However, the proposed wage index values for 184 (45.1 percent) urban areas and 24 (51.1 percent) rural areas would decrease. The proposed wage index values for 85 (20.8 percent) urban areas would decrease by greater than or equal to 1 percent but less than 5 percent, and no urban areas’ final wage index value would decrease by 5 percent or more. The largest proposed positive impacts would be 17.4 percent for an urban area and 2.9 percent for a rural area. The largest proposed negative impacts would be 4.9 percent for an urban area and 2.3 percent for a rural area. One urban area’s proposed wage index, but no rural area wage indexes, would remain unchanged by application of the occupational mix adjustment. These results indicate that a larger percentage of urban areas (54.7 percent) would benefit from the occupational mix adjustment than would rural areas (48.9 percent).

G. Proposed Application of the Rural, Imputed, and Frontier Floors

1. Proposed Rural Floor

Section 4410(a) of Public Law 105–33 provides that, for discharges on or after October 1, 1997, the area wage index applicable to any hospital that is located in an urban area of a State may not be less than the area wage index applicable to hospitals located in rural areas in that State. This provision is referred to as the “rural floor.” Section 3141 of Public Law 111–148 also requires that a national budget neutrality adjustment be applied in implementing the rural floor. Based on the proposed FY 2018 wage index associated with this proposed rule (which is available via the Internet on the CMS Web site), we estimated that 366 hospitals would receive an increase in their FY 2018 proposed wage index due to the application of the rural floor.

2. Proposed Expiration of the Imputed Floor Policy

In the FY 2005 IPPS final rule (69 FR 49109 through 49111), we adopted the “imputed floor” policy as a temporary 3-year regulatory measure to address concerns from hospitals in all-urban States that have argued that they are disadvantaged by the absence of rural hospitals to set a wage index floor for those States. Since its initial implementation, we have extended the application of the rural floor policy for FY 2013. Under the alternative methodology, we first determined the average percentage difference between the post-reclassified, pre-floor area wage index and the post-reclassified, rural floor wage index (without rural floor budget neutrality applied) for all CBSAs receiving the rural floor. (Table 4D associated with the FY 2013 IPPS/LTCH PPS final rule (which is available via the Internet on the CMS Web site) included the CBSAs receiving a State’s rural floor wage index.) The lowest postreclassified wage index assigned to a hospital in an all-urban State having a range of such values then is increased by this factor, the result of which establishes the State’s alternative imputed floor. We amended § 412.64(h)(4) of the regulations to add new paragraphs to incorporate the finalized alternative methodology, and to make reference and date changes. In summary, for the FY 2013 wage index, we did not make any changes to the original imputed floor methodology at § 412.64(h)(4) and, therefore, made no changes to the New Jersey imputed floor calculation for FY 2013. Instead, for FY 2013, we adopted a second, alternative methodology for use in cases where an all-urban State has a range of wage indexes assigned to its hospitals, but the State cannot benefit under the original methodology. In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50589 through 50590), we extended the imputed floor policy (both the original methodology and the alternative methodology) for 1 additional year through September 30, 2014, while we continued to explore potential wage index reforms. In the FY 2015 IPPS/LTCH PPS final rule (79 FR 49969 through 49970), for FY 2015, we adopted a policy to extend the imputed floor policy (both the original methodology and alternative methodology) for another year, through September 30, 2015, as we continued to explore potential wage index reforms. In that final rule, we revised the regulations at § 412.64(h)(4) and (h)(4)(vi) to reflect the 1-year extension of the imputed floor.

As discussed in section III.B. of the preamble of that FY 2015 final rule, we adopted the new OMB labor market area delineations beginning in FY 2015. Under the new OMB delineations, Delaware became an all-urban State, along with New Jersey and Rhode Island. Under the new OMB delineations, Delaware has three CBSAs, New Jersey has seven CBSAs, and Rhode Island continues to have only one CBSA (Providence-Warwick, RI–MA). We refer readers to a detailed discussion of our adoption of the new...
OMB labor market area delineations in section III.B. of the preamble of the FY 2015 IPPS/LTCH PPS final rule. Therefore, under the adopted new OMB delineations discussed in section III.B. of the preamble of the FY 2015 IPPS/LTCH PPS final rule, Delaware became an all-urban State and was subject to an imputed floor as well for FY 2015.

In the FY 2016 IPPS/LTCH PPS final rule (80 FR 49497 through 49498), for FY 2016, we extended the imputed floor policy (under both the original methodology and the alternative methodology) for 1 additional year, through September 30, 2016. In that final rule, we revised the regulations at § 412.64(h)(4) and (h)(4)(vi) to reflect this additional 1-year extension.

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 56921 through 56922), for FY 2017, we extended the imputed floor policy (under both the original methodology and the alternative methodology) for 1 additional year, through September 30, 2017. In that final rule, we revised the regulations at § 412.64(h)(4) and (h)(4)(vi) to reflect this additional 1-year extension.

The imputed floor is set to expire effective October 1, 2017, and we are not proposing to extend the imputed floor policy. In the FY 2005 IPPS final rule (69 FR 49110), we adopted the imputed floor policy for all-urban States under the authority of section 1886(d)(3)(E) of the Act, which gives the Secretary broad authority to adjust the proportion (as estimated by the Secretary from time to time) of hospitals’ costs which are attributable to wages and wage-related costs of the DRG prospective payment rates for area differences in hospital wage levels by a factor (established by the Secretary).

However, we have expressed reservations about establishment of an imputed floor, considering that the imputed rural floor methodology creates a disadvantage in the application of the wage index to hospitals in States with rural hospitals but no urban hospitals receiving the rural floor (72 FR 24786 and 72 FR 47322). As we discussed in the FY 2008 IPPS final rule (72 FR 47322), the application of the rural and imputed floors requires transfer of payments from hospitals in States with rural hospitals but where the rural floor is not applied to hospitals in States where the rural or imputed floor is applied. For this reason, in this proposed rule, we are proposing not to apply an imputed floor to wage index calculations and payments for hospitals in all-urban States for FY 2018 and subsequent fiscal years.

Therefore, only States containing both rural areas and hospitals located in such areas (including any hospital reclassified as rural under the provisions of § 412.103 of the regulations) would benefit from the rural floor, in accordance with section 4410 of Public Law 105–33. In addition, we would no longer include the imputed floor as a factor in the national budget neutrality adjustment. Therefore, the proposed wage index and impact tables associated with this FY 2018 IPPS/LTCH PPS proposed rule (which are available via the Internet on the CMS Web site) do not reflect the imputed floor policy, and there is no proposed national budget neutrality adjustment for the imputed floor for FY 2018. We are inviting public comments on our proposal not to extend the imputed floor for FY 2018 and subsequent years.

3. Proposed State Frontier Floor for FY 2018

Section 10324 of Public Law 111–148 requires that hospitals in frontier States cannot be assigned a wage index of less than 1.000. (We refer readers to the regulations at 42 CFR 412.64(m) and to a discussion of the implementation of this provision in the FY 2011 IPPS/LTCH PPS final rule (75 FR 50160 through 50161.) Fifty-two hospitals would receive the frontier floor value of 1.0000 for their FY 2018 wage index in this proposed rule. These hospitals are located in Montana, Nevada, North Dakota, South Dakota, and Wyoming. We are not proposing any changes to the frontier floor policy for FY 2018. The areas affected by the proposed rural and frontier floor policies for the proposed FY 2018 wage index are identified in Table 3E associated with this proposed rule, which is available via the Internet on the CMS Web site.

H. Proposed FY 2018 Wage Index Tables

In the FY 2016 IPPS/LTCH PPS final rule (80 FR 49498 and 49807 through 49808), we finalized a proposal to streamline and consolidate the wage index tables associated with the IPPS proposed and final rules for FY 2016 and subsequent fiscal years. Prior to FY 2016, the wage index tables had consisted of 12 tables (Tables 2, 3A, 3B, 4A, 4B, 4C, 4D, 4E, 4F, 4J, 9A, and 9C) that were made available via the Internet on the CMS Web site. Effective beginning FY 2016, with the exception of Table 3A, we streamlined and consolidated 11 tables (Tables 2, 3A, 3B, 4A, 4B, 4C, 4D, 4F, 4J, 9A, and 9C) into 2 tables (Tables 2 and 3). We refer readers to section VI. of the Addendum to this proposed rule for a discussion of the proposed wage index tables for FY 2018.

1. Revisions to the Wage Index Based on Hospital Redesignations and Reclassifications

a. General Policies and Effects of Reclassification and Redesignation

Under section 1886(d)(10) of the Act, the Medicare Geographic Classification Review Board (MGCRB) considers applications by hospitals for geographic recategorization for purposes of payment under the IPPS. Hospitals must apply to the MGCRB to reclassify not later than 13 months prior to the start of the fiscal year for which recategorization is sought (usually by September 1). Generally, hospitals must be proximate to the labor market area to which they are seeking reclassification and must demonstrate characteristics similar to hospitals located in that area. The MGCRB issues its decisions by the end of February for recategorizations that become effective for the following fiscal year (beginning October 1). The regulations applicable to recategorizations by the MGCRB are located in 42 CFR 412.280. (We refer readers to a discussion in the FY 2002 IPPS final rule (66 FR 39874 and 39875) regarding how the MGCRB defines mileage for purposes of the proximity requirements.) The general policies for recategorizations and redesignations and the policies for the effects of hospitals’ recategorizations and redesignations on the wage index are discussed in the FY 2012 IPPS/LTCH PPS final rule for the FY 2012 final wage index (76 FR 51595 and 51596). In addition, in the FY 2012 IPPS/LTCH PPS final rule, we discussed the effects on the wage index of urban hospitals reclassifying to rural areas under 42 CFR 412.103. Hospitals that are geographically located in States without any rural areas are ineligible to apply for rural reclassification in accordance with the provisions of 42 CFR 412.103.

On April 21, 2016, we published an interim final rule with comment period (IFC) in the Federal Register (81 FR 23428 through 23438) that included provisions amending our regulations to allow hospitals nationwide to have simultaneous § 412.103 and MGCRB recategorizations. For recategorizations effective beginning FY 2018, a hospital may acquire rural status under § 412.103 and subsequently apply for a Table 4F, we streamlined and consolidated 11 tables (Tables 2, 3A, 3B, 4A, 4B, 4C, 4D, 4F, 4J, 9A, and 9C) into...
addition, we provided that a hospital that has an active MGCRB reclassification and is then approved for redesignation under § 412.103 will not lose its MGCRB reclassification; such a hospital receives a reclassified urban wage index during the years of its active MGCRB reclassification and is still considered rural under section 1886(d) of the Act and for other purposes.

We discussed that when there is both a § 412.103 redesignation and an MGCRB reclassification, the MGCRB reclassification controls for wage index calculation and payment purposes. We exclude hospitals with § 412.103 redesignations from the calculation of the reclassified rural wage index if they also have an active MGCRB reclassification to another area. That is, if an application for urban reclassification through the MGCRB is approved, and is not withdrawn or terminated by the hospital within the established timelines, we consider the hospital's geographic CBSA and the urban CBSA to which the hospital is reclassified under the MGCRB for the wage index calculation. We refer readers to the April 21, 2016 IFC (81 FR 23428 through 23438) and the FY 2017 IPPS/LTCH PPS final rule (81 FR 56922 through 56930) for a full discussion of the effect of simultaneous reclassifications under both the § 412.103 and the MGCRB processes on wage index calculations.

2. MGCRB Reclassification and Redesignation Issues for FY 2018

a. FY 2018 Reclassification Requirements and Approvals

As previously stated, under section 1886(d)(10) of the Act, the MGCRB considers applications by hospitals for geographic reclassification for purposes of payment under the IPPS. The specific procedures and rules that apply to the geographic reclassification process are outlined in regulations under 42 CFR 412.230 through 412.260. At the time this proposed rule was constructed, the MGCRB had completed its review of FY 2018 reclassification requests. Based on such reviews, there are 375 hospitals approved for wage index reclassifications by the MGCRB starting in FY 2018. Because MGCRB wage index reclassifications are effective for 3 years, for FY 2018, hospitals reclassified beginning in FY 2016 or FY 2017 are eligible to continue to be reclassified to a particular labor market area based on such prior reclassifications for the remainder of their 3-year period. There were 257 hospitals approved for wage index reclassifications in FY 2016 that will continue for FY 2018, and 274 hospitals approved for wage index reclassifications in FY 2017 that will continue for FY 2018. Of all the hospitals approved for reclassification for FY 2016, FY 2017, and FY 2018, based upon the review at the time of this proposed rule, 906 hospitals are in a MGCRB reclassification status for FY 2018.

Under the regulations at 42 CFR 412.273, hospitals that have been reclassified by the MGCRB are permitted to withdraw their applications if the request for withdrawal is received by the MGCRB within 45 days of the publication of CMS' annual notice of proposed rulemaking concerning changes to the inpatient hospital prospective payment system and proposed payment rates for the fiscal year for which the application has been filed. (We note that in section III.I.4. of the preamble of this proposed rule, we are proposing to revise the above described regulation text to specify that written notice to the MGCRB must be provided within 45 days from the date of public display of the proposed rule at the Office of the Federal Register. If finalized, that proposal would be effective beginning with the FY 2019 IPPS/LTCH PPS proposed rule.) For information about withdrawing, terminating, or canceling a previous withdrawal or termination of a 3-year reclassification for wage index purposes, we refer readers to § 412.273, as well as the FY 2002 IPPS final rule (66 FR 39887 through 39888) and the FY 2003 IPPS final rule (67 FR 50065 through 50066). Additional discussion on withdrawals and terminations, and clarifications regarding reinstating reclassifications and “fallback” reclassifications were included in the FY 2008 IPPS final rule (72 FR 47333).

Changes to the wage index that result from withdrawals of requests for reclassification, terminations, wage index corrections, appeals, and the Administrator's review process for FY 2018 will be incorporated into the wage index values published in the FY 2018 IPPS/LTCH PPS final rule. These changes affect not only the wage index value for specific geographic areas, but also the wage index value that redesignated/reclassified hospitals receive; that is, whether they receive the wage index that includes the data for both the hospitals already in the area and the redesignated/reclassified hospitals. Further, the wage index value for the area from which the hospitals are redesignated/reclassified may be affected.

Applications for FY 2019 reclassifications are due to the MGCRB by September 1, 2017 (the first working day of September 2017). We note that this is also the deadline for canceling a previous wage index reclassification, withdrawal, or termination under 42 CFR 412.273(d). Applications and other information about MGCRB reclassifications may be obtained, beginning in mid-July 2017, via the Internet on the CMS Web site at: https://www.cms.gov/Regulations-and-Guidance/Review-Boards/MGCRB/index.html, or by calling the MGCRB at (410) 786–1174. The mailing address of the MGCRB is: 2520 Lord Baltimore Drive, Suite L, Baltimore, MD 21244–2670.

Under previous regulations at 42 CFR 412.256(a)(1), applications for reclassification were required to be mailed or delivered to the MGCRB, with a copy to CMS, and were not allowed to be submitted through the facsimile (FAX) process or by other electronic means. Because we believed this previous policy was outdated and overly restrictive and to promote ease of application submission, we took steps to: (1) Modernize the MGCRB application process, and would reduce the overall burden upon hospitals.

b. Extension of PRA Information Collection Requirement Approval for MGCRB Applications

As stated earlier, under section 1886(d)(10) of the Act, the MGCRB considers applications by hospitals for geographic reclassification for purposes of payment under the IPPS. The specific
procedures and rules that apply to the geographic reclassification process are outlined in the regulations under 42 CFR 412.230 through 412.280. The current information collection requirements for the MGCRB procedures and criteria and supporting regulations in 42 CFR 412.256 subject to the Paperwork Reduction Act provisions are currently approved under OMB Control Number 0938–0573 and expired on February 28, 2017. An extension of the currently approved collection is required in time for applications due to the MGCRB September 1, 2017 for FY 2019 reclassifications. As discussed in section XIII.B. of the preamble of this proposed rule, a request for an extension of the current information collection requirements for the MGCRB procedures and criteria and supporting regulations is currently awaiting approval by OMB and can be accessed at: https://www.reginfo.gov/public/do/PRA VIEWCR?ref_nbr=201612–0938–023.

c. Proposed Deadline for Submittal of Documentation of Sole Community Hospital (SCH) and Rural Referral Center (RRC) Classification Status to the MGCRB

The regulations at 42 CFR 412.230(a)(3), consistent with section 1886(d)(10)(D)(i)(III) of the Act, set special rules for sole community hospitals (SCHs) and rural referral centers (RRCs) that are reclassifying under the MGCRB. Specifically, a hospital that is an RRC or an SCH, or both, does not have to demonstrate a close proximity to the area to which it seeks redesignation. If a hospital that is an RRC or an SCH, or both, qualifies for urban redesignation, it is redesignated to the urban area that is closest to the hospital. If the hospital is closer to another rural area than to any urban area, it may seek redesignation to either the closest rural or the closest urban area.

In addition, section 1886(d)(10)(D)(i)(III) of the Act, as implemented in the regulations at § 412.230(d)(3)(i), provides an exception to certain wage comparison criteria for RRCs and former RRCs reclassifying under the MGCRB. Under § 412.230(d)(3)(i), if a hospital was ever an RRC, it does not have to demonstrate that it meets the average hourly wage criterion at § 412.230(d)(1)(iii), which would require that the hospital’s average hourly wage be at least 106 percent for rural hospitals and at least 108 percent for urban hospitals of the average hourly wage of all other hospitals in the area in which the hospital is located. Rather, as codified at § 412.230(d)(3)(ii), consistent with our authority under section 1886(d)(10)(D)(i) of the Act, if a hospital was ever an RRC, it is required to meet only the criterion for rural hospitals at § 412.230(d)(1)(iv), which requires that the hospital’s average hourly wage is equal to at least 82 percent of the average hourly wage of hospitals in the area to which it seeks redesignation. The regulations at § 412.96 set forth the criteria that a hospital must meet in order to qualify as an RRC.

For a hospital to use the special rules at § 412.230(a)(3) for SCHs and RRCs, the existing regulation at § 412.230(a)(3) requires that the hospital be an active SCH or an RRC as of the date of the MGCRB’s review. In addition, for a hospital to use the RRC exceptions at § 412.230(d)(3), a hospital must either be an RRC at the time of the MGCRB’s review or have previously been classified as an RRC in the past. In other words, under the existing regulations, if a hospital is approved by CMS as an SCH or an RRC but the approval is not yet effective at the time of the MGCRB’s review, the hospital’s status as an SCH or an RRC would not be considered in the MGCRB’s decision, unless the hospital was a former RRC, in which case it would be able to use the RRC exceptions at § 412.230(d)(3).

The MGCRB currently accepts supporting documentation of SCH and RRC classification (the CMS approval letter) up until the date of MGCRB’s review, which varies annually. A hospital may apply at any time for classification as an SCH, and the classification is effective 30 days after the date of CMS’ written notification of approval, in accordance with § 412.92. Considering that the MGCRB usually meets in early February, hospitals typically seek to obtain SCH approval letters no later than early January (30 days prior to the date of MGCRB review) for the SCH status to be effective as of the date of the MGCRB’s review. However, consistent with section 1886(d)(5)(C)(i) of the Act, a hospital must submit its application for RRC status during the quarter before the first quarter of the hospital’s cost reporting period, to be effective at the beginning of the next cost reporting period. The existing regulation at § 412.230(a)(3), combined with the statutory timeframe for RRC classification, require that a hospital’s cost reporting period as an RRC begin on or before the date of the MGCRB’s review in order to be considered an RRC by the MGCRB for purposes of the special rules under § 412.230(a)(3). Similarly, in order to use the RRC exceptions under § 412.230(d)(3) the hospital had previously been classified as an RRC in the past.

The current practice of accepting SCH and RRC approvals up until the date of MGCRB review does not ensure adequate time for the MGCRB to include SCH and RRC approvals in its review. We note that many hospitals now obtain SCH or RRC status based on a § 412.103 reclassification in order to reclassify using the special rules and exceptions under the MGCRB following the April 21, 2016 IFC (81 FR 23428), which revised the regulations to allow hospitals nationwide to reclassify based on acquired rural status. We believe that the additional volume of SCH and RRC approvals submitted to the MGCRB increases the need for an earlier deadline for documentation of SCH and RRC classifications to be submitted to the MGCRB for purposes of the special rules at § 412.230(a)(3) and the exception for RRCs at § 412.230(d)(3). In addition, because the date of the MGCRB’s review varies annually, we believe hospitals would be relieved from the uncertainty of a set date by which documentation of SCH or RRC status must be submitted in order to have that status considered by the MGCRB under § 412.230(a)(3) and § 412.230(d)(3).

Therefore, to ensure sufficient time for the MGCRB to include SCH and RRC status approvals in its review and increase clarity for hospitals, while allowing as much time and flexibility as possible for hospitals applying for SCH and RRC status to be considered in the MGCRB review, we are proposing to revise the regulations at § 412.230(a)(3) and § 412.230(d)(3). We are proposing to revise the regulations at § 412.230(a)(3) in two ways. First, we are proposing to establish a deadline of the first business day after January 1 for hospitals to submit to the MGCRB documentation of SCH or RRC status approval (the CMS approval letter) in order to take advantage of the special rules under § 412.230(a)(3) when reclassifying under the MGCRB. We believe that this date of the first business day after January 1 would provide sufficient time for the MGCRB to consider documentation of
SCH or RRC status approval in its review, without negatively affecting hospitals seeking to obtain SCH or RRC status, as explained below. Second, we are proposing to revise § 412.230(a)(3) to require hospitals to submit documentation of SCH or RRC status approval (the CMS approval letter) by the deadline above, rather than to have SCH or RRC classification that is effective as of the date of MGCRB review, in order to use the special rules for SCHs and RRCs under § 412.230(d)(3). Likewise, we are proposing to revise the regulations at § 412.230(d)(3) so that a hospital qualifies for these RRC exceptions if it was ever approved as a RRC. In other words, the exceptions at § 412.230(d)(3) would continue to apply to hospitals that were ever classified as RRCs, but consistent with our authority under section 1886(d)(10)(D)(i) of the Act to publish guidelines to be utilized by the MGCRB, we would also extend these exceptions to hospitals that were ever approved as RRCs. Similar to § 412.230(a)(3), we also are proposing to establish a deadline of the first business day after January 1 for hospitals to submit documentation of RRC status approval (the CMS approval letter) in order to take advantage of the exception under § 412.230(d)(3) when reclassifying under the MGCRB. These proposed revisions would more appropriately allow the MGCRB to prepare for its review and would allow hospitals obtaining SCH or RRC status approval as late as the first business day after January 1 to have these classifications considered by the MGCRB under § 412.230(a)(3) and (d)(3), irrespective of the effective date of these classifications. These proposals would not substantially affect hospitals seeking SCH classification for purposes of reclassifying under the MGCRB because a hospital must obtain SCH status approval by early January under the existing regulations in order to have that classification effective 30 days later by the time the Board usually meets in early February. For hospitals seeking RRC classification for purposes of reclassifying under the MGCRB, however, the proposed deadline of no later than the first business day after January 1, in concert with our proposal to accept documentation of approval (the CMS approval letter) instead of requiring the hospital to be an active RRC at the time of the MGCRB review in order to take advantage of the special rules and exceptions under § 412.230(d)(3), is beneficial. The proposed revisions to the regulations at § 412.230(a)(3) and (d)(3) accommodate more hospitals with various cost reporting year ends by allowing hospitals with cost reporting periods beginning soon after the MGCRB’s decision to have RRC status approvals included in the MGCRB’s review. Under the proposals, the MGCRB would consider an RRC status approval obtained as late as the first business day after January 1 instead of requiring the RRC classification to be effective by the time the Board meets, which has been in February in past years. For example, a hospital with a cost reporting period beginning as late as March, which could apply for RRC status approval in accordance with the statutory timeframe starting in December, would be considered an RRC by the MGCRB if it submits documentation of approval of RRC status no later than the first business day after January 1, even though the approval would not be effective until after the MGCRB’s decision.

For the reasons discussed above, consistent with our authority under section 1886(d)(10)(D)(i) of the Act to publish guidelines to be utilized by the MGCRB, we are proposing to revise the regulations at § 412.230(a)(3) to specify that, to be reclassified under the special rules in that paragraph, the hospital must submit documentation of the approval of SCH or RRC status to the MGCRB no later than the first business day after January 1. In addition, we are proposing confining revisions to paragraphs (a)(3)(i) and (ii) of § 412.230 to reflect that these paragraphs apply to hospitals with RRC approval as specified above (and not only effective status). Specifically, we are proposing to revise § 412.230(a)(3)(i) to specify that a hospital that is approved as an RRC or SCH, or both, does not have to demonstrate a close proximity to the area to which it seeks redesignation; and to revise § 412.230(a)(3)(ii) to specify that this paragraph applies if a hospital that is approved as an RRC or SCH, or both, qualifies for urban SCH or RRC classification that is effective as of the date of MGCRB review, in order to use the special rules in that paragraph, the hospital must submit documentation of the approval of SCH or RRC status to the MGCRB no later than the first business day after January 1. In addition, we are proposing to revise § 412.230(a)(3)(ii) to specify that if a hospital was ever approved as an RRC, it is required to meet only the criterion that applies to rural hospitals under § 412.230(d)(1)(iv), regardless of its actual location in an urban or rural area.

We are inviting public comments on these proposals.

d. Clarification of Special Rules for SCHs and RRCs Reclassifying to Geographic Home Area

Following issuance of the April 21, 2016 IFC (81 FR 23428), hospitals may simultaneously be redesignated as rural under § 412.103 and reclassified under the MGCRB. An urban hospital seeking benefits of rural status, such as rural payments for disproportionate share hospitals (DSH) and eligibility for the 340B Drug Pricing Program administered by HRSA, without the associated rural wage index may be redesignated as rural under § 412.103 (if it meets the applicable requirements) and also reclassify under the MGCRB to an urban area (again, if it meets the applicable requirements). As discussed earlier and in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56922 through 56927), a hospital with simultaneous § 412.103 redesignation and MGCRB reclassification receives the wage index of the CBSA to which it is redesignated under the MGCRB while still maintaining § 412.103 reclassified rural status for other purposes.

Hospitals that are redesignated under § 412.103 may seek MGCRB reclassification to their geographic home area. Such hospitals automatically meet the criteria for proximity, but must still demonstrate that they meet the wage comparison requirements using the criteria for rural hospitals at § 412.230(d). Specifically, a hospital with a § 412.103 redesignation seeking reclassification under the MGCRB must demonstrate that its average hourly wage is at least 106 percent of the average hourly wage of all other hospitals in the area in which the hospital is located in accordance with § 412.230(d)(1)(iii), and the hospital’s average hourly wage is equal to at least 82 percent of the average hourly wage of hospitals in the area to which it seeks redesignation, in accordance with § 412.230(d)(1)(iv). In this case, both the area in which the hospital is located and
the area to which it seeks redesignation are the geographic home area.

If a hospital with a § 412.103 rural redesignation also has SCH or RRC status based on its acquired rural status, the hospital may use the exception at § 412.230(d)(3) for RRCs seeking reclassification under the MGCRB and the special reclassification rules at § 412.230(a)(3) for SCHs and RRCs. Specifically, under § 412.230(d)(3)(ii), an RRC or former RRC must only demonstrate that its average hourly wage is equal to at least 82 percent of the average hourly wage of hospitals in the area to which it seeks redesignation. In other words, a hospital with RRC status based on a § 412.103 rural redesignation that is seeking additional reclassification under the MGCRB to its geographic home area must only demonstrate that its average hourly wage is equal to at least 82 percent of the average hourly wage of hospitals in its geographic home area. The proximity requirement is waived under § 412.230(a)(3) for SCHs and RRCs, and SCHs and RRCs are redesignated to the urban area that is closest to the hospital (or if the hospital is closer to another rural area than to any urban area, it may seek redesignation to either the closest rural area or the closest urban area).

The existing regulation at § 412.230(a)(3)(ii) states that if an SCH or RRC qualifies for urban redesignation, it is redesignated to the urban area that is closest to the hospital. As currently worded, we believe it is unclear how this provision would apply to hospitals with a § 412.103 rural redesignation and SCH or RRC status. If the urban area that is closest to the hospital is interpreted to mean the hospital’s geographic home area, a hospital with a § 412.103 rural redesignation and SCH or RRC status would not be able to reclassify to any closest area outside of the hospital’s geographic home area, but would only be allowed to reclassify to the geographic home area. Alternatively, if the urban area that is closest to the hospital is interpreted to mean the closest urban area to the hospital’s geographic home area, the hospital would seem to be precluded from reclassifying under the MGCRB to its geographic home area. In other words, under the existing language of this regulation, the urban area that is closest to the hospital can either be interpreted to mean the hospital’s geographic home area, or the closest area outside of the hospital’s geographic home area.

We believe it would be appropriate to revise § 412.230(d)(3)(ii) to clarify that it allows for redesignation to either the hospital’s geographic home area or to the closest area outside of the hospital’s geographic home area. Prior to the April 21, 2016 interim final rule with comment period (IFC) (81 FR 23423), it was not possible for a hospital with § 412.103 rural redesignation to seek reclassification to its geographic home area or to the closest area outside its geographic home area under the MGCRB because dual reclassification under § 412.103 and under the MGCRB was not permitted. However, the IFC allowed dual § 412.103 and MGCRB reclassifications, so a hospital may now reclassify to a rural area under § 412.103 and then reclassify back to its geographic home area or another area under the MGCRB for wage index purposes (if it meets all criteria).

Therefore, we are proposing to revise the regulations at § 412.230(a)(3)(ii) to clarify that a hospital with a § 412.103 rural redesignation and SCH or RRC approval may reclassify under the MGCRB to its geographic home area or to the closest area outside of its geographic home area. Specifically, we are proposing to revise § 412.230(a)(3)(ii) to state that if a hospital that is approved as an RRC or an SCH, or both, qualifies for urban redesignation, it is redesignated to the urban area that is closest to the hospital or to the hospital’s geographic home area. If the hospital is closer to another rural area than to any urban area, it may seek redesignation to either the closest rural or the closest urban area.

3. Redesignations Under Section 1886(d)(6)(B) of the Act

In the FY 2012 IPPS/LTCH PPS final rule (76 FR 51599 through 51600), we adopted the policy that, beginning with FY 2012, an eligible hospital that waives its Lugar status in order to receive the out-migration adjustment has effectively waived its deemed urban status and, thus, is rural for all purposes under the IPPS effective for the fiscal year in which the hospital receives the out-migration adjustment. In addition, we adopted a minor procedural change that would allow a Lugar hospital that qualifies for and accepts the out-migration adjustment (through written notification to CMS within 45 days from the publication of the proposed rule) to waive its urban status for the full 3-year period for which its out-migration adjustment is effective. We further clarified that if a hospital wishes to reinstate its urban status for any fiscal year within this 3-year period, it must send a request to CMS within 45 days of publication of the proposed rule for that particular fiscal year. We indicated that such reinstatement requests may be sent electronically to wageindex@cms.hhs.gov. We wish to further clarify that both requests to waive and to reinstate “Lugar” status may be sent to this mailbox. To ensure proper accounting, we request hospitals to include their CCN, and either “waive Lugar” or “reinstate Lugar”, in the subject line of these requests.

4. Proposed Changes to the 45-Day Notification Rules

Certain Medicare regulations specify that hospitals have 45 days from the publication of the annual proposed rule for the hospital inpatient prospective payment system to inform CMS or the MGCRB of certain requested reclassification/redesignation and out-migration adjustment changes relating to the development of the hospital wage index. Specifically, 42 CFR 412.64(i)(iii), which provides for adjusting the wage index to account for commuting patterns of hospital workers, and 42 CFR 412.211(f)(3)(iii), which provides for the same adjustment for hospitals in Puerto Rico, state that a hospital may waive the application of this wage index adjustment by notifying CMS in writing within 45 days after the publication of the annual notice of proposed rulemaking for the hospital inpatient prospective payment system. The regulations at § 412.273(c)(ii) concerning withdrawing an MGCRB application, terminating an approved 3-year reclassification, or canceling a previous withdrawal or termination, also state (specifically § 412.273(c)(1)(ii)
and (2) that a request for withdrawal or termination must be received by the MGCRB within 45 days of publication of CMS' annual notice of proposed rulemaking concerning changes to the inpatient hospital prospective payment system and proposed payment rates. Similarly, the policy outlined in the FY 2012 IPPS/LTCH PPS final rule (76 FR 51599 through 51600) allows a Lugar hospital that qualifies for and accepts the out-migration adjustment, or that no longer wishes to accept the out-migration adjustment and instead elects to return to its deemed urban status to notify CMS within 45 days from the date of public display of the proposed rule at the Office of the Federal Register. We are inviting public comments on these proposals.

J. Proposed Out-Migration Adjustment Based on Commuting Patterns of Hospital Employees

In accordance with section 1886(d)(13) of the Act, as added by section 505 of Pub. L. 108-173, beginning with the FY 2005, we established a process to make adjustments to the hospital wage index based on commuting patterns of hospital employees (the "out-migration" adjustment). The process, outlined in the FY 2005 IPPS final rule (69 FR 49061), provides for an increase in the wage index for hospitals located in certain counties that have a relatively high percentage of hospital employees who reside in the county but work in a different county (or counties) with a higher wage index.

Section 1886(d)(13)(B) of the Act requires the Secretary to use data the Secretary determines to be appropriate to establish the qualifying counties. When the provision of section 1886(d)(13) of the Act was implemented for the FY 2005 wage index, we analyzed commuting data compiled by the U.S. Census Bureau that were derived from a special tabulation of the 2000 Census journey-to-work data for all industries (CMS extracted data applicable to hospitals). These data were compiled from responses to the "long-form" survey, which the Census Bureau used at the time and which contained questions on where residents in each county worked (69 FR 49062). However, the 2010 Census was "short form" only; information on where residents in each county worked was not collected as part of the 2010 Census. The Census Bureau worked with CMS to provide an alternative dataset based on the latest available data on where residents in each county worked in 2010, for use in developing a new out-migration adjustment based on new commuting patterns developed from the 2010 Census data beginning with FY 2016.

To determine the out-migration adjustments and applicable counties for FY 2016, we analyzed commuting data compiled by the Census Bureau that were derived from a custom tabulation of the American Community Survey (ACS), an official Census Bureau survey, utilizing 2008 through 2012 (5-Year) Microdata. The data were compiled from responses to questions regarding the county where workers reside and the county to which workers commute. As we discussed in the FY 2016 and FY 2017 IPPS/LTCH PPS final rules (80 FR 49501 and 81 FR 56930, respectively), the same policies, procedures, and computations that were used for the FY 2012 out-migration adjustment were applicable for FY 2016 and FY 2017, and we are proposing to use them again for FY 2018. We have applied the same policies, procedures, and computations since FY 2012, and we believe they continue to be appropriate for FY 2018. We refer readers to the FY 2016 IPPS/LTCH PPS final rule (80 FR 49500 through 49502) for a full explanation of the revised data source.

For FY 2018, until such time that CMS finalizes out-migration adjustments based on the next Census, the out-migration adjustment continues to be based on the data derived from the custom tabulation of the ACS utilizing 2008 through 2012 (5-Year) Microdata. For FY 2018, we are not proposing any changes to the methodology or data source that we used for FY 2016 (81 FR 25071). We refer readers to a full discussion of the out-migration adjustment, including rules on deeming hospitals reclassified under section 1886(d)(8) or section 1886(d)(10) of the Act to have waived the out-migration adjustment, in the FY 2012 IPPS/LTCH PPS final rule (76 FR 51601 through 51602). Table 2 associated with this proposed rule (which is available via the Internet on the CMS Web site) includes the proposed out-migration adjustments for the FY 2018 wage index.

K. Reclassification From Urban to Rural Under Section 1886(d)(8)(E) of the Act, Implemented at 42 CFR 412.103

Under section 1886(d)(8)(E) of the Act, a qualifying prospective payment hospital located in an urban area may apply for rural status for payment purposes separate from reclassification through the MGCRB. Specifically, section 1886(d)(8)(E) of the Act provides that, not later than 60 days after the receipt of an application (in a form and manner determined by the Secretary) from a subsection (d) hospital that satisfies certain criteria, the Secretary shall treat the hospital as being located in the rural area (as defined in paragraph (2)(D) of the State in which the hospital is located. We refer readers to the regulations at 42 CFR 412.103 for the general criteria and application requirements for a subsection (d) hospital to reclassify from urban to rural status in accordance with section 1886(d)(8)(E) of the Act. The FY 2012 IPPS/LTCH PPS final rule (76 FR 51595 through 51596) includes our policies.
regarding the effect of wage data from reclassified or redesignated hospitals. Hospitals must meet the criteria to be reclassified from urban to rural status under § 412.103, as well as fulfill the requirements for the application process. There may be one or more reasons that a hospital applies for the urban to rural reclassification, and the timeframe that a hospital submits an application is often dependent on those reasons(s). Because the wage index is part of the methodology for determining the prospective payments to hospitals for each fiscal year, we believe there should be a definitive timeframe within which a hospital should apply for rural status in order for the reclassification to be reflected in the next Federal fiscal year’s wage data used for setting payment rates.

Therefore, after notice of proposed rulemaking and consideration of public comments, in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56931 through 56932), we revised § 412.103(b) by adding (c) to specify that, in order for a hospital to be treated as rural in the wage index and budget neutrality calculations under § 412.64(e)(1)(ii), (e)(2), (e)(4), and (h) for payment rates for the next Federal fiscal year, the hospital’s filing date must be no later than 70 days prior to the second Monday in June of the current Federal fiscal year and the application must be approved by the CMS Regional Office in accordance with the requirements of § 412.103. We refer readers to the FY 2017 IPPS/LTCH PPS final rule for a full discussion of this policy. We clarified that the lock-in date does not affect the timing of payment changes occurring at the hospital-specific level as a result of reclassification from urban to rural under § 412.103. This lock-in date also does not change the current regulation that allows hospitals that qualify under § 412.103(a) to request, at any time during a cost reporting period, to reclassify from urban to rural. A hospital’s rural status and claims payment reflecting its rural status continue to be effective on the filing date of its reclassification application, which is the date the CMS Regional Office receives the application, in accordance with § 412.103(d). The hospital’s IPPS claims will be paid reflecting its rural status on the filing date (the effective date) of the reclassification, regardless of when the hospital applies.

L. Clarification of Application Deadline for Rural Referral Center (RRC) Classification

Section 1886(d)(5)(C)(i) of the Act, implemented at 42 CFR 412.96, provides for the classification and special treatment of rural referral centers (RRCs). The regulations at § 412.96 set forth the criteria that a hospital must meet in order to qualify as an RRC. Under § 412.96(b)(1)(ii), a hospital may qualify as an RRC if it is located in a rural area and has 275 or more beds during its most recently completed cost reporting period. The hospital also can obtain RRC status by showing that at least 50 percent of its Medicare patients are referred from other hospitals or from physicians not on the staff of the hospital, and at least 60 percent of the hospital’s Medicare patients live more than 25 miles from the hospital, and at least 60 percent of all the services that the hospital furnishes to Medicare beneficiaries are furnished to beneficiaries who live more than 25 miles from the hospital (§ 412.96(b)(2)), or by showing that the hospital meets the alternative criteria at § 412.96(c). We refer readers to 42 CFR 412.96 for a full description of the criteria for classification as an RRC.

Consistent with section 1886(d)(5)(C)(i) of the Act, the hospital must submit its application for RRC status during the last quarter of the hospital’s cost reporting period, to be effective with the beginning of the next cost reporting period. Specifically, section 1886(d)(5)(C)(i) of the Act provides that an appeal allowed under this paragraph must be submitted to the Secretary (in such form and manner as the Secretary may prescribe) during the quarter before the first quarter of the hospital’s cost reporting period (or, in the case of a cost reporting period beginning during October 1984, during the first quarter of that period), and the Secretary must make a final determination with respect to such appeal within 60 days after the date the appeal was submitted. Any payment adjustments necessitated by a reclassification based upon the appeal will be effective at the beginning of such cost reporting period. Therefore, in this proposed rule, we are clarifying that applications for RRC status must be submitted during the timeframe that is, applications for RRC status must be submitted during the last quarter of the cost reporting period before the first quarter of a hospital’s cost reporting year. If approved, the RRC status is effective with the beginning of the hospital’s cost reporting period occurring after the last quarter of the hospital’s cost reporting year in which the hospital submits an application.

Because a hospital may only apply for RRC status during the last quarter of its cost reporting year in accordance with section 1886(d)(5)(C)(i) of the Act, hospitals seeking RRC status, in order to reclassify through the MCCRB using the special rules for SCHs and RRCs at § 412.230(a)(3) and the exceptions at § 412.230(d)(3) for RRCs, may be disadvantaged due to their cost reporting year end. As discussed in section III.I.2. of the preamble of this proposed rule, we are proposing to revise the regulations at § 412.230(a)(3) and (d)(3) to allow hospitals to submit documentation of the approval of SCH or RRC status (as applicable) to the MCCRB no later than the first business day after January 1. We believe our proposal to accept documentation of approval of RRC classification, instead of requiring that the hospital be classified as an RRC at the time of Board review, would accommodate more hospitals with various cost reporting period endings. We refer readers to section III.I.2. of the preamble of this proposed rule for further discussion of this proposal.

M. Process for Wage Index Data Corrections

1. Process for Hospitals To Request Wage Index Data Corrections

The preliminary, unaudited Worksheet 5–3 wage data files for the proposed FY 2018 wage index were made available on May 16, 2016, and the preliminary CY 2013 occupational
mix data files on May 16, 2016, through the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Wage-Index-Files-Items/FY2018-Wage-Index-Home-Page.html.

On January 30, 2017, we posted a public use file (PUF) at https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Wage-Index-Files-Items/FY2018-Wage-Index-Home-Page.html containing FY 2018 wage index data available as of January 29, 2017. This PUF contains a tab with the Worksheet S–3 wage data (which includes Worksheet S–3, Parts II and III wage data from cost reporting periods beginning on or after October 1, 2013 through September 30, 2014; that is, FY 2014 wage data), a tab with the occupational mix data (which includes data from the CY 2013 occupational mix survey, Form CMS–10079), a tab containing the Worksheet S–3 wage data of hospitals deleted from the January 30, 2017 wage data PUF, and a tab containing the CY 2013 occupational mix data (if any) of the hospitals deleted from the January 30, 2017 wage data PUF. In a memorandum dated January 27, 2017, we instructed all MACs to inform the IPPS hospitals that they service of the availability of the January 30, 2017 wage index data PUFs, and the process and timeframe for requesting revisions in accordance with the FY 2018 Wage Index Timetable.

In the interest of meeting the data needs of the public, beginning with the proposed FY 2009 wage index, we post an additional PUF on our Web site that reflects the actual data that are used in computing the proposed wage index. The release of this file does not alter the current wage index process or schedule. We notify the hospital community of the availability of these data as we do with the current public use wage data files through our Hospital Open Door Forum. We encourage hospitals to sign up for automatic notifications of information about hospital issues and the dates of the Hospital Open Door Forums at the CMS Web site at: http://www.cms.gov/Outreach-and-Education/Outreach/OpenDoorForums/index.html.

In a memorandum dated May 16, 2016, we instructed all MACs to inform the IPPS hospitals that they service of the availability of the wage index data files and the process and timeframe for requesting revisions. We also instructed the MACs to advise hospitals that these data were also made available directly through their representative hospital organizations.

If a hospital wished to request a change to its data as shown in the May 16, 2016 wage data files and May 16, 2016 occupational mix data files, the hospital had to submit corrections along with complete, detailed supporting documentation to its MAC by September 2, 2016. Hospitals were notified of this deadline and of all other deadlines and requirements, including the requirement to review and verify their data as posted in the preliminary wage index data files on the Internet, through the letters sent to them by their MACs.

November 4, 2016 was the date by which MACs notified State hospital associations regarding hospitals that failed to respond to issues raised during the desk reviews. The MACs notified the hospitals by mid-January 2017 of any changes to the wage index data as a result of the desk reviews and the resolution of the hospitals’ revision requests. The MACs also submitted the revised data to CMS by January 20, 2017. CMS published the wage index PUFs that included hospitals’ revised wage index data on January 30, 2017. Hospitals had until February 17, 2017, to submit requests to the MACs for reconsideration of adjustments made by the MACs as a result of the desk review, and to correct errors due to CMS’ or the MAC’s mishandling of the wage index data. Hospitals also were required to submit sufficient documentation to support their requests.

After reviewing requested changes submitted by hospitals, MACs were required to transmit to CMS any additional revisions resulting from the hospitals’ reconsideration requests by March 24, 2017. Under our current policy, the deadline for a hospital to request CMS intervention in cases where a hospital disagreed with a MAC’s policy interpretation was April 5, 2017. Beginning next year (that is, April 2018 for wage data revisions for the FY 2019 wage index), we are proposing to require that a hospital that seeks to challenge the MAC’s handling of wage data on any basis (including a policy, factual, or any other dispute) must request CMS to intervene by the date in April that is specified as the deadline for hospitals to appeal MAC determinations and request CMS’ intervention in cases where a hospital disagrees with the MAC’s determination (the wage index timetable would be updated to reflect the specified date). We note that, as we did for the FY 2017 wage index, for the FY 2018 wage index, in accordance with the FY 2018 wage index timeline posted on the CMS Web site at https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Wage-Index-Files-Items/FY2018-Wage-Index-Home-Page.html, the April appeals have to be sent via mail and email. We refer readers to the wage index timeline for complete details.

Hospitals are given the opportunity to examine Table 2, which is listed in section VI. of the Addendum to this proposed rule and available via the Internet on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Wage-Index-Files-Items/FY2018-Wage-Index-Home-Page.html. Table 2 contains each hospital’s proposed adjusted average hourly wage used to construct the wage index values for the past 3 years, including the FY 2014 data used to construct the proposed FY 2018 wage index. We note that the proposed hospital average hourly wages shown in Table 2 only reflect changes made to a hospital’s data that were transmitted to CMS by early February 2017.

We plan to post the final wage index data PUFs in late April 2017 on the Internet at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Wage-Index-Files-Items/FY2018-Wage-Index-Home-Page.html.

If, after reviewing the April 2017 PUF, a hospital disagrees with the final wage index data that resulted from the correction process previously described (revisions submitted to CMS by the MACs by March 24, 2017), after the release of the April 2017 wage index data PUFs, changes to the wage and occupational mix data can only be made in those very limited situations involving an error by the MAC or CMS that the hospital could not have known about before its review of the final wage index data files. Specifically, neither the MAC nor CMS will approve the following types of requests:

- Requests for wage index data corrections that were submitted too late to be included in the data transmitted to CMS by the MACs on or before March 24, 2017.
- Requests for correction of errors that were not, but could have been, identified during the hospital’s review of the January 30, 2017 wage index PUFs.
- Requests to revisit factual determinations or policy interpretations made by the MAC or CMS during the wage index data correction process.

If, after reviewing the April 2017 final wage index data PUFs, a hospital believes that its wage or occupational
mix data were incorrect due to a MAC or CMS error in the entry or tabulation of the final data, the hospital is given the opportunity to notify both its MAC and CMS regarding why the hospital believes an error exists and provide all supporting information, including relevant dates (for example, when it first became aware of the error). The hospital is required to send its request to CMS and to the MAC no later than May 30, 2017. Similar to the April appeals, beginning with the FY 2015 wage index, in accordance with the FY 2018 wage index timeline posted on the CMS Web site at https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Wage-Index-Files-Items/FY2018-Wage-Index-Home-Page.html, the May appeals must be sent via mail and email to CMS and the MACs. We refer readers to the wage index timeline for complete details.

Verified corrections to the wage index data received timely by CMS and the MACs (that is, by May 30, 2017) will be incorporated into the final FY 2018 wage index, which will be effective October 1, 2017.

We created the processes previously described to resolve all substantive wage index data correction disputes before we finalize the wage and occupational mix data for the FY 2018 payment rates. Accordingly, hospitals that do not meet the procedural deadlines set forth above will not be afforded a later opportunity to submit wage index data corrections or to dispute the MAC’s decision with respect to requests for wage index corrections. Specifically, our policy is that hospitals that do not meet the procedural deadlines set forth above (requiring requests to MACs by the specified date in February and, where such requests are unsuccessful, requests for intervention by CMS by the specified date in April) will not be permitted to challenge later, before the PRRB, the failure of CMS to make a requested data revision. We refer readers also to the FY 2000 IPPS final rule (64 FR 41513) for a discussion of the parameters for appeals to the PRRB for wage index data corrections.

Again, we believe the wage index data correction process described earlier provides hospitals with sufficient opportunity to bring errors in their wage and occupational mix data to the MAC’s attention. Moreover, because hospitals have access to the final wage index data PUFs by late April 2017, they have the opportunity to detect any data entry or tabulation errors made by the MAC or CMS before the development and publication of the final FY 2018 wage index by August 2017, and the implementation of the FY 2018 wage index on October 1, 2017. Given these processes, the wage index implemented on October 1 should be accurate.

Nevertheless, in the event that errors are identified by hospitals and brought to our attention after May 30, 2017, we retain the right to make midyear changes to the wage index under very limited circumstances.

Specifically, in accordance with 42 CFR 412.64(k)(1) of our regulations, we make midyear corrections to the wage index for an area only if a hospital can show that: (1) The MAC or CMS made an error in tabulating its data; and (2) the requesting hospital could not have known about the error or did not have an opportunity to correct the error before the beginning of the fiscal year. For purposes of this provision, “before the beginning of the fiscal year” means by the May deadline for making corrections to the wage data for the following fiscal year’s wage index (for example, May 30, 2017 for the FY 2018 wage index). This provision is not available to a hospital seeking to revise another hospital’s data that may be affecting the requesting hospital’s wage index for the labor market area. As indicated earlier, because CMS makes the wage index data available to hospitals on the CMS Web site prior to publishing both the proposed and final IPPS rules, and the MACs notify hospitals directly of any wage index data changes after completing their desk reviews, we do not expect that midyear corrections will be necessary. However, under our current policy, if the correction of a data error changes the wage index value for an area, the revised wage index value will be effective prospectively from the date the correction is made.

In the FY 2006 IPPS final rule (70 FR 47385 through 47387 and 47485), we revised 42 CFR 412.64(k)(2) to specify that, effective on October 1, 2005, that is, beginning with the FY 2006 wage index, a change in the wage index can be made retroactive to the beginning of the Federal fiscal year only when CMS determines all of the following: (1) The MAC or CMS made an error in tabulating data used for the wage index calculation; (2) the hospital knew about the error and requested that the MAC and CMS correct the error using the established process and within the established schedule for requesting corrections to the wage index data, before the beginning of the fiscal year for the applicable IPPS update (that is, by the May 30, 2017 deadline for the FY 2018 wage index); and (3) CMS agreed before October 1 that the MAC or CMS made an error in tabulating the hospital’s wage index data and the wage index should be corrected.

In those circumstances where a hospital requested a correction to its wage index data before CMS calculated the final wage index (that is, by the May 30, 2017 deadline for the FY 2018 wage index), and CMS acknowledges that the error in the hospital’s wage index data was caused by CMS’ or the MAC’s mishandling of the data, we believe that the hospital should not be penalized by our delay in publishing or implementing the correction. As with our current policy, we indicated that the provision is not available to a hospital seeking to revise another hospital’s data. In addition, the provision cannot be used to correct prior years’ wage index data; and it can only be used for the current Federal fiscal year. In situations where our policies would allow midyear corrections other than those specified in 42 CFR 412.64(k)(2)(ii), we continue to believe that it is appropriate to make prospective-only corrections to the wage index.

We note that, as with prospective changes to the wage index, the final retroactive correction will be made irrespective of whether the change increases or decreases a hospital’s payment rate. In addition, we note that the policy of retroactive adjustment will still apply in those instances where a final judicial decision reverses a CMS denial of a hospital’s wage index data revision request.

2. Process for Data Corrections by CMS After the January Public Use File (PUF)

The process set forth with the wage index timeline discussed in section III.M.1. of the preamble of this proposed rule allows hospitals to request corrections to their wage index data within prescribed timeframes. In addition to hospitals’ opportunity to request corrections of wage index data errors or MACs’ mishandling of data, CMS has the authority under section 1886(d)(3)(E) of the Act to make corrections to hospital wage index and occupational mix data in order to ensure the accuracy of the wage index. As we explained in the FY 2016 IPPS/LTC PPS final rule (80 FR 49490 through 49491) and the FY 2017 IPPS/LTC PPS final rule (81 FR 56914), section 1886(d)(3)(E) of the Act requires the Secretary to adjust the proportion of hospitals’ costs attributable to wages and wage-related costs for area differences reflecting the relative hospital wage level in the geographic areas of the hospital compared to the national average hospital wage level. We believe that, under section 1886(d)(3)(E) of the Act, we have discretion to make
corrections to hospitals’ data to help ensure that the costs attributable to wages and wage-related costs in fact accurately reflect the relative hospital wage level in the hospitals’ geographic areas.

We have established a multistep, 15-month process for the review and correction of the hospital wage data that is used to create the IPPS wage index for the upcoming fiscal year. Since the origin of the IPPS, the wage index has been subject to its own annual review process, first by the MACs, and then by CMS. As a standard practice, after each annual desk review, CMS reviews the results of the MACs’ desk reviews and focuses on items flagged during the desk review, requiring that, if necessary, hospitals provide additional documentation, adjustments, or corrections to the data. This ongoing communication with hospitals about their wage data may result in the discovery by CMS of additional items that were reported incorrectly or other data errors, even after the posting of the January PUF, and throughout the remainder of the wage index development process. In addition, the fact that CMS analyzes the data from a regional and even national level, unlike the review performed by the MACs that review a limited subset of hospitals, can facilitate additional editing of the data that may not be readily apparent to the MACs. In these occasional instances, an error may be of sufficient magnitude that the wage index of an entire CBSA is affected. Accordingly, CMS uses its authority to ensure that the wage index accurately reflects the relative hospital wage level in the geographic area of the hospital compared to the national average hospital wage level, by continuing to make corrections to hospital wage data upon discovering incorrect wage data, distinct from instances in which hospitals request data revisions.

We note that CMS corrects errors to hospital wage data as appropriate, regardless of whether that correction will raise or lower a hospital’s average hourly wage. For example, as discussed in section III.D.2. of the preamble of this proposed rule, in the calculation of the proposed FY 2018 wage index, upon discovering that hospitals reported other wage-related costs on Line 18 of Worksheet S–3, despite those other wage-related costs failing to meet the requirement that other wage related costs must exceed 1 percent of total adjusted salaries net of excluded area salaries, CMS made internal edits to remove those other wage-related costs from Line 18. Conversely, if CMS discovers after conclusion of the desk review, for example, that a MAC inadvertently failed to incorporate positive adjustments resulting from a prior year’s wage index appeal to a hospital’s wage related costs such as pension, CMS would correct that data error and the hospital’s average hourly wage would likely increase as a result.

While we maintain CMS’ authority to conduct additional review and make resulting corrections at any time during the wage index development process, we are proposing a process for hospitals to request further review of a correction made by CMS starting with the FY 2019 wage index. In order to allow opportunity for input from hospitals concerning corrections made by CMS after the posting of the January PUF, we are proposing a process similar to the existing process in which hospitals may request corrections to wage index data displayed in the January PUF. Instances where CMS makes a correction to a hospital’s data after the January PUF based on a different understanding than the hospital about certain reported costs, for example, could potentially be resolved using this proposed process before the final wage index is calculated. We believe this proposed process and timeline (as described above) would bring additional transparency to instances where CMS makes data corrections after the January PUF, and would provide opportunities for hospitals to request further review of CMS changes in time for the most accurate data to be reflected in the final wage index calculations.

Effective beginning with the FY 2019 wage index development cycle, we are proposing to use existing appeal deadlines (in place for hospitals to appeal determinations made by the MAC during the desk review process) for hospitals to dispute corrections made by CMS after posting of the January PUF that do not arise from a hospital request for a wage data revision. Starting with the April appeal deadline, hospitals would use the soonest approaching appeal deadline to dispute any adjustments made by CMS. However, if a hospital was notified of an adjustment within 14 days of an appeal deadline, the hospital would have until the next appeal deadline to dispute any adjustments. We believe this would give hospitals sufficient time to prepare an appeal of adjustments made by CMS after the January PUF. Specifically, for any adjustments made by CMS between the date the January PUF is posted and at least 14 calendar days before the April appeals deadline, we are proposing that hospitals would have until the April appeals deadline (which, for example, is April 5 in the FY 2018 Wage Index Timetable) to dispute the adjustments. For any adjustments made by CMS between 13 calendar days before the April appeals deadline and 14 calendar days before the May appeals deadline, we are proposing that hospitals would have until the May appeals deadline (which, for example, is May 30 in the FY 2018 Wage Index Timetable) to dispute the adjustments. In cases where hospitals disagree with CMS adjustments of which they were notified 13 calendar days before the May appeals deadline or later, the hospitals could appeal to the PRRB with no need for further review by CMS before such appeal.

We are using dates from the FY 2018 Wage Index Timetable in the following example (we reiterate that this appeals process would be effective beginning with the FY 2019 wage index cycle, but for illustrative purposes, we are using dates from the FY 2018 Wage Index Timetable, the most recently published wage index timetable): A hospital that is notified by the MAC or CMS of an adjustment to its wage data after the release of the January 30, 2017 PUF could use the April 5, 2017 appeals deadline to dispute the adjustment. If the hospital is notified of an adjustment by CMS or the MAC to its wage data after March 22, 2017 (that is, less than 14 days prior to the April 5 appeals deadline), it could use the May 30, 2017 appeals deadline to dispute the adjustment. If the hospital is first notified about the adjustment after May 16, 2017 (that is, less than 14 days prior to the May 30 appeals deadline), and disagrees with the adjustment, the hospital could appeal directly to the PRRB.

As with the existing process for requesting wage data corrections, we are proposing that a hospital disputing an adjustment made by CMS after the posting of the January PUF would be required to request a correction by the first applicable deadline. For example, if a hospital was notified on March 20 of an adjustment to its data by CMS and does not appeal by April 5, the hospital would not be able to appeal by May 30 or bring the case before the PRRB. That is, hospitals that did not meet the procedural deadlines set forth above would not be afforded a later opportunity to submit wage index data corrections or to dispute CMS’ decision with respect to requested changes. As with the existing process for hospitals to request wage data corrections, our policy is that hospitals that do not meet the procedural deadlines set forth earlier would not be permitted to challenge later, before the PRRB, the failure of CMS to make a requested data revision.
In summary, under the statute, CMS has discretion to make corrections and revisions to hospitals’ wage data throughout the multistep wage index development process, and we are proposing a pathway for hospitals to request additional review of corrections to their wage data made by CMS. Beginning with the development of the FY 2019 wage index, we are proposing a process whereby CMS could continue to correct data after the posting of the January PUF, while allowing hospitals to appeal changes made by CMS using existing deadlines from the process for hospitals to request wage data corrections. As with the existing process, a hospital would be required to appeal by the first applicable deadline, if relevant, to maintain the right to appeal to the PRRB to dispute a correction to its wage data made by CMS.

We are inviting public comments on our proposals.

N. Proposed Labor Market Share for the Proposed FY 2018 Wage Index

Section 1886(d)(3)(E) of the Act directs the Secretary to adjust the proportion of the national prospective payment system base payment rates that are attributable to wages and wage-related costs by a factor that reflects the relative differences in labor costs among geographic areas. It also directs the Secretary to estimate from time to time the proportion of hospital costs that are labor-related and to adjust the proportion (as estimated by the Secretary from time to time) of hospitals’ costs which are attributable to wages and wage-related costs of the DRG prospective payment rates. We refer to the portion of hospital costs attributable to wages and wage-related costs as the labor-related share. The labor-related share of the prospective payment rate is adjusted by an index of relative labor costs, which is referred to as the wage index.

Section 403 of Public Law 108–173 amended section 1886(d)(3)(E) of the Act to provide that the Secretary must employ 62 percent as the labor-related share unless this would result in lower payments to a hospital than would otherwise be made. However, this provision of Public Law 108–173 did not change the legal requirement that the Secretary estimate from time to time the proportion of hospitals’ costs that are attributable to wages and wage-related costs. Thus, hospitals receive payment based on either a 62-percent labor-related share, or the labor-related share estimated from time to time by the Secretary, depending on which labor-related share resulted in a higher payment.

In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50596 through 50667), we rebased and revised the hospital market basket. We established a FY 2010-based IPPS hospital market basket to replace the FY 2006-based IPPS hospital market basket, effective October 1, 2013. In that final rule, we presented our analysis and conclusions regarding the frequency and methodology for updating the labor-related share for FY 2014. Using the FY 2010-based IPPS market basket, we finalized a labor-related share for FY 2014, FY 2015, FY 2016, and FY 2017 of 69.6 percent. In addition, in FY 2014, we implemented this revised and rebased labor-related share in a budget neutral manner (78 FR 51016). However, consistent with section 1886(d)(3)(E) of the Act, we did not take into account the additional payments that would be made as a result of hospitals with a wage index less than or equal to 1.0000 being paid using a labor-related share lower than the labor-related share of hospitals with a wage index greater than 1.0000.

For FY 2018, as described in section IV. of the preamble of this proposed rule, we are proposing to rebase and revise the IPPS market basket reflecting 2014 data. We also are proposing to recalculate the labor-related share for discharges occurring on or after October 1, 2017 using the proposed 2014-based IPPS market basket. As discussed in Appendix A of this proposed rule, we are proposing this revised and rebased labor-related share in a budget neutral manner. However, consistent with section 1886(d)(3)(E) of the Act, we did not take into account the additional payments that would be made as a result of hospitals with a wage index less than or equal to 1.0000 being paid using a labor-related share lower than the labor-related share of hospitals with a wage index greater than 1.0000. The labor-related share is used to determine the proportion of the national IPPS base payment rate to which the area wage index is applied. We include a cost category in the labor-related share if the costs are labor intensive and vary with the local labor market. As described in section IV. of the preamble of this proposed rule, we are proposing to include in the labor-related share the national average proportion of operating costs that are attributable to Wages and Salaries, Employee Benefits, Professional Fees: Labor-Related, Administrative and Facilities Support Services, Installation, Maintenance, and Repair Services, and All Other: Labor-Related Services as measured in the proposed 2014-based IPPS market basket. Therefore, for FY 2018, we are proposing to use a labor-related share of 68.3 percent for discharges occurring on or after October 1, 2017.

Prior to January 1, 2016, Puerto Rico hospitals were paid based on 75 percent of the national standardized amount and 25 percent of the Puerto Rico-specific standardized amount. As a result, we applied the Puerto Rico-specific labor-related share percentage and nonlabor-related share percentage to the Puerto Rico-specific standardized amount. Section 601 of the Consolidated Appropriations Act, 2016 (Pub. L. 114–113) amended section 1886(d)(9)(E) of the Act to specify that the payment calculation with respect to operating costs of inpatient hospital services of a subsection (d) Puerto Rico hospital for inpatient hospital discharges on or after January 1, 2016, shall use 100 percent of the national standardized amount. Because Puerto Rico hospitals are no longer paid with a Puerto Rico-specific standardized amount as of January 1, 2016, under section 1886(d)(9)(E) of the Act as amended by section 601 of the Consolidated Appropriations Act, 2016, there is no longer a need for us to calculate a Puerto Rico-specific labor-related share percentage and nonlabor-related share percentage for application to the Puerto Rico-specific standardized amount. Hospitals in Puerto Rico are now paid 100 percent of the national standardized amount and, therefore, are subject to the national labor-related share and nonlabor-related share percentages that are applied to the national standardized amount.

Accordingly, for FY 2018, we are not proposing a Puerto Rico-specific labor-related share percentage or a nonlabor-related share percentage.

Tables 1A and 1B, which are published in section VI. of the Addendum to this FY 2018 IPPS/LTCH PPS proposed rule and available via the Internet on the CMS Web site, reflect the proposed national labor-related share, which is also applicable to Puerto Rico hospitals. For FY 2018, for all IPPS hospitals (including Puerto Rico hospitals) whose wage indexes are less than or equal to 1.0000, we are proposing to apply the wage index to a labor-related share of 62 percent of the national standardized amount. For all hospitals (including Puerto Rico hospitals) whose wage indexes are greater than 1.0000, for FY 2018, we are proposing to apply the wage index to a proposed labor-related share of 68.3 percent of the national standardized amount.
IV. Proposed Rebasing and Revising of the Hospital Market Baskets for Acute Care Hospitals

A. Background

Effective for cost reporting periods beginning on or after July 1, 1979, we developed and adopted a hospital input price index (that is, the hospital market basket for operating costs). Although “market basket” technically describes the mix of goods and services used in providing hospital care, this term is also commonly used to denote the input price index (that is, cost category weights and price proxies combined) derived from that market basket. Accordingly, the term “market basket” as used in this document refers to the hospital input price index.

The percentage change in the market basket reflects the average change in the price of goods and services hospitals purchase in order to provide inpatient care. We first used the market basket to adjust hospital cost limits by an amount that reflected the average increase in the prices of the goods and services used to provide hospital inpatient care. This approach linked the increase in the cost limits to the efficient utilization of resources.

Since the inception of the IPPS, the projected change in the hospital market basket has been the integral component of the update factor by which the prospective payment rates are updated every year. An explanation of the hospital market basket used to develop the prospective payment rates was published in the Federal Register on September 1, 1983 (48 FR 39764). We also refer readers to the FY 2014 IPPS/LTCH PPS final rule (78 FR 50596), with FY 2010 data used in the hospital market basket. Only when the index is rebased would changes in the quantity and intensity be captured, with those changes being reflected in the cost weights. Therefore, we rebase the market basket periodically so that the cost weights reflect recent changes in the mix of goods and services purchased subsequent to the base period. For example, a hospital hiring more nurses to accommodate the needs of patients would increase the volume of goods and services purchased by the hospital, but would not be factored into the price change measured by a fixed-weight hospital market basket. Only when the index is rebased would the quantities and intensities be captured, with those changes being reflected in the cost weights.

The hospital market basket is a fixed-weight, Laspeyres-type price index. A Laspeyres-type price index measures the change in price, over time, of the same mix of goods and services purchased in the base period. Any changes in the quantity or mix of goods and services (that is, intensity) purchased over time are not measured.

The index itself is constructed in three steps. First, a base period is selected (in this proposed rule, we are proposing to use 2014 as the base period) and total base period expenditures are estimated for a set of mutually exclusive and exhaustive spending categories, with the proportion of total costs that each category represents being calculated. These proportions are called “cost weights” or “expenditure weights.” Second, each expenditure category is matched to an appropriate price or wage variable, referred to as a “price proxy.” In almost every instance, these price proxies are derived from publicly available statistical series that are published on a consistent schedule (preferably at least on a quarterly basis). Finally, the expenditure weight for each cost category is multiplied by the level of its respective price proxy. The sum of these products (that is, the expenditure weights multiplied by their price index levels) for all cost categories yields the composite index level of the market basket in a given period. Repeating this step for other periods produces a series of market basket levels over time. Dividing an index level for a given period by an index level for an earlier period produces a rate of growth in the input price index over that timeframe.

As noted above, the market basket is described as a fixed-weight index because it represents the change in price over time of a constant mix (quantity and intensity) of goods and services needed to provide hospital services. The effects on total expenditures resulting from changes in the mix of goods and services purchased subsequent to the base period are not measured. For example, a hospital hiring more nurses to accommodate the needs of patients would increase the volume of goods and services purchased by the hospital, but would not be factored into the price change measured by a fixed-weight hospital market basket. Only when the index is rebased would changes in the quantities and intensities be captured, with those changes being reflected in the cost weights. Therefore, we rebase the market basket periodically so that the cost weights reflect recent changes in the mix of goods and services purchased subsequent to the base period. For example, a hospital hiring more nurses to accommodate the needs of patients would increase the volume of goods and services purchased by the hospital, but would not be factored into the price change measured by a fixed-weight hospital market basket. Only when the index is rebased would changes in the quantities and intensities be captured, with those changes being reflected in the cost weights.

The index is rebased every 4 years. Therefore, we rebase the hospital market basket cost weights effective for FY 2014 (78 FR 50596), with FY 2010 data used as the base period for the construction of the market basket cost weights. For this FY 2018 IPPS/LTCH PPS proposed rule, we are proposing to rebase the cost structure for the IPPS hospital index from FY 2010 to 2014, as discussed below.

B. Rebasing and Revising the IPPS Market Basket

The terms “rebasing” and “revising,” while often used interchangeably, actually denote different activities. “Rebasing” means moving the base year for the structure of costs of an input price index. For example, under this proposed rule, we are proposing to shift the base year cost structure for the IPPS hospital index from FY 2010 to 2014. We note that we are no longer referring to the market basket as a “FY 2014-based” market basket and instead refer to the proposed market basket as simply “2014-based.” We are proposing this change in naming convention for the market basket because the base year cost weight data for the proposed market basket does not reflect only fiscal year data. For example, the proposed 2014-based IPPS market basket uses Medicare cost report data and other government data that reflect 2014 fiscal year, 2014 calendar year, and 2014 State fiscal year expenses to determine the base year cost weights. Given that it is based on a mix of classifications of 2014 data, we are proposing to refer to the market basket as “2014-based” instead of “FY 2014-based” or “CY 2014-based”.

“Revising” means changing data sources or price proxies used in the input price index. As published in the FY 2006 IPPS final rule (70 FR 47387), in accordance with section 404 of Public Law 108–173, CMS determined a new frequency for rebasing the hospital market basket. We established a rebasing frequency of every 4 years and, therefore, for the FY 2018 IPPS update, we are proposing to rebase and revise the IPPS market basket from FY 2010 to 2014. We are inviting public comments on our proposed methodology.

1. Development of Cost Categories and Weights

a. Use of Medicare Cost Report Data

The major source of expenditure data for developing the proposed rebased and revised hospital market basket cost weights is the 2014 Medicare cost reports. These 2014 Medicare cost reports are for cost reporting periods beginning on and after October 1, 2013 and before October 1, 2014. We note that while these dates appear to reflect fiscal year data, in order to be classified as a “2014 cost report,” a hospital’s cost reporting period must begin between these dates. For example, we found that of the 2014 Medicare cost reports for IPPS hospitals, approximately 40 percent of the reports had a begin date on January 1, 2014, approximately 30 percent had a begin date on July 1, 2014, and approximately 18 percent had a begin date on October 1, 2013. For this reason, we are defining the base year of the market basket as “2014-based” instead of “FY 2014-based.” We are proposing to use 2014 as the base year because we believe that the 2014 Medicare cost reports represent the most recent and reliable cost data available to develop cost weights for IPPS hospitals. As was done...
in previous rebasings, these cost reports are from IPPS hospitals only (hospitals excluded from the IPPS and CAHs are not included) and are based on IPPS Medicare-allowable operating costs. IPPS Medicare-allowable operating costs are costs that are eligible to be paid under the IPPS. For example, the IPPS market basket excludes home health agency (HHA) costs as these costs would be paid under the HHA PPS and, therefore, these costs are not IPPS Medicare-allowable costs.

We are proposing to derive costs for eight major expenditures or cost categories for the 2014-based IPPS market basket from the CMS Medicare cost reports (Form 2552–10, OMB Control Number 0938–0050): Wages and Salaries, Employee Benefits, Contract Labor, Pharmaceuticals, Professional Liability Insurance (Malpractice), Blood and Blood Products, Home Office Contract Labor, and a residual “All Other” category. The residual “All Other” category reflects all remaining costs that are not captured in the other seven cost categories. We are proposing that, for the 2014-based IPPS market basket, we obtain costs for one additional major cost category from the Medicare cost reports compared to the FY 2010-based IPPS market basket—Home Office Contract Labor Costs. We describe below the detailed methodology for obtaining costs for each of the seven cost categories directly determined from the Medicare cost reports.

(1) Wages and Salaries Costs

To derive wages and salaries costs for the Medicare allowable cost centers, we are proposing to first calculate total unadjusted wages and salaries costs as reported on Worksheet S–3, part II. We are then proposing to remove the wages and salaries attributable to non-Medicare allowable cost centers (that is, excluded areas) as well as a portion of overhead wages and salaries attributable to these excluded areas. Specifically, wages and salaries costs are equal to total wages and salaries (as defined earlier) to total wages and salaries (Worksheet S–3, part II, Column 4, Line 1) by total overhead wages and salaries (Worksheet A, Column 1, Lines 4 through 18). A similar methodology was used to derive wages and salaries costs in the FY 2010-based IPPS market basket.

(2) Employee Benefits Costs

We are proposing to derive employee benefits costs using a similar methodology as the wages and salaries costs; that is, reflecting employee benefits costs attributable to the Medicare allowable cost centers. First, we calculate total unadjusted employee benefits costs as the sum of Worksheet S–3, part II, Column 4, Lines 17, 18, 20, and 22. We then exclude those employee benefits attributable to the overhead wages and salaries for the non-Medicare allowable cost centers (that is, excluded areas). Employee benefits attributable to the non-Medicare allowable cost centers are derived by multiplying the ratio of total employee benefits (equal to the sum of Worksheet S–3, part II, Column 4, Lines 17 through 25) to total wages and salaries (Worksheet S–3, part II, Column 4, Line 1) by excluded overhead wages and salaries (as derived above for wages and salaries costs). A similar methodology was used in the FY 2010-based IPPS market basket.

(3) Contract Labor Costs

Contract labor costs are primarily associated with direct patient care services. Contract labor costs for services such as accounting, billing, and legal are estimated using other government data sources as described below. We are proposing to derive contract labor costs for the 2014-based IPPS market basket as the sum of Worksheet S–3, part II, Column 4, Lines 11, 13 and 15. A similar methodology was used in the FY 2010-based IPPS market basket.

(4) Professional Liability Insurance Costs

We are proposing that professional liability insurance (PLI) costs (often referred to as malpractice costs) be equal to premiums, paid losses, and self-insurance costs reported on Worksheet S–2, part I, Columns 1 through 3, Line 118.01. A similar methodology was used for the FY 2010-based IPPS market basket.

(5) Pharmaceuticals Costs

We are proposing to calculate pharmaceuticals costs using nonsalary costs reported for the Pharmacy cost center (Worksheet A, Column 2, Line 15) and Drugs Charged to Patients cost center (Worksheet A, Column 2, Line 73) less estimated employee benefits attributable to these two cost centers. We are proposing to estimate these employee benefits costs by multiplying the ratio of total employee benefits (equal to the sum of Worksheet S–3, part II, Column 4, Lines 17 through 25) to total wages and salaries (Worksheet S–3, part II, Column 4, Line 1) by total wages and salaries costs for the Pharmacy and Drugs Charged to Patients cost centers (equal to the sum of Worksheet A, Column 1, Lines 15 and 73). A similar methodology was used for the FY 2010-based IPPS market basket.

(6) Blood and Blood Products Costs

We are proposing to calculate blood and blood products costs using nonsalary costs reported for the Whole Blood & Packaged Red Blood Cells cost center (Worksheet A, Column 2, Line 62) and the Blood Storing, Processing, & Transfusing cost center (Worksheet A, Column 2, Line 63) less estimated employee benefits attributable to these two cost centers. We estimate these employee benefits costs by multiplying the ratio of total employee benefits (equal to the sum of Worksheet S–3, part II, Column 4, Lines 17 through 25) to total wages and salaries (Worksheet S–3, part II, Column 4, Line 1) by total wages and salaries for the Whole Blood & Packaged Red Blood Cells and Blood Storing, Processing, & Transfusing cost centers (equal to the sum of Worksheet A, Column 4, Lines 62 and 63). A similar methodology was used for the FY 2010-based IPPS market basket.

(7) Home Office Contract Labor Costs

We are proposing to determine home office contract labor costs using data reported on Worksheet S–3, part II, Column 4, line 14. Specifically, we are proposing to determine the Medicare allowable portion of these costs by multiplying them by the ratio of total Medicare allowable operating costs (as defined below in section IV.B.1.b. of the preamble to this proposed rule) to total operating costs (calculated as Worksheet B, part I, Column 26, Line 202, less Worksheet B, part I, Column 0, Lines 1 through 3). Home office contract labor costs in the FY 2010-based IPPS market basket were calculated using the U.S. Census Bureau’s Bureau of Economic Analysis (BEA) Benchmark Input-Output (IO) data, as described below in section IV.B.1.c. of the preamble to this proposed rule.
b. Final Major Cost Category Computation

After we derived costs for the seven major cost categories for each provider using the Medicare cost report data as previously described, we address data outliers using the following steps. First, we divide the costs for each of the seven categories by total Medicare allowable operating costs calculated for the provider to obtain cost weights for each PPS hospital. We are proposing that total Medicare allowable operating costs are equal to noncapital costs (Worksheet B, part I, Column 26 less Worksheet B, part II, Column 26) that are attributable to the Medicare allowable cost centers of the hospital. Medicare allowable cost centers are Lines 30 through 35, 50, 51, 53 through 60, 62 through 76, 90, 91, 92.01 and 93.

We then remove those providers whose derived cost weights fall in the top and bottom five percent of provider-specific cost weights to ensure the removal of outliers. After the outliers have been removed, we sum the costs for each category across all remaining providers. We then divide this by the sum of total Medicare allowable operating costs across all remaining providers to obtain a cost weight for the proposed 2014-based IPPS market basket for the given category. Finally, we calculate the residual “All Other” cost weight that reflects all remaining costs that are not captured in the seven cost categories listed.

Table IV–01 below shows the major cost categories and their respective cost weights as derived from the Medicare cost reports for this proposed rule.

### TABLE IV–01—MAJOR COST CATEGORIES AS DERIVED FROM THE MEDICARE COST REPORTS

<table>
<thead>
<tr>
<th>Major cost categories</th>
<th>FY 2010</th>
<th>Proposed 2014</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wages and Salaries</td>
<td>45.6</td>
<td>42.1</td>
</tr>
<tr>
<td>Employee Benefits</td>
<td>12.7</td>
<td>12.0</td>
</tr>
<tr>
<td>Contract Labor</td>
<td>1.5</td>
<td>1.8</td>
</tr>
<tr>
<td>Professional Liability Insurance (Malpractice)</td>
<td>1.3</td>
<td>1.2</td>
</tr>
<tr>
<td>Pharmaceuticals</td>
<td>5.4</td>
<td>5.9</td>
</tr>
<tr>
<td>Blood and Blood Products</td>
<td>1.1</td>
<td>0.8</td>
</tr>
<tr>
<td>Home Office Contract Labor*</td>
<td>-</td>
<td>4.2</td>
</tr>
<tr>
<td>“All Other” Residual</td>
<td>31.9</td>
<td>32.0</td>
</tr>
</tbody>
</table>

*Home office contract labor costs were included in the “All Other” residual cost weight of the FY 2010-based IPPS market basket.

From FY 2010 to 2014, the Wages and Salaries and Employee Benefits cost weights as calculated directly from the Medicare cost reports decreased by approximately 3.7 and 0.7 percentage points, respectively, while the Contract Labor cost weight was unchanged. The decrease in the Wages and Salaries cost weight occurred among most cost centers and in aggregate for the General Service (overhead), Inpatient Routine Service, Ancillary Service, and Outpatient Service cost centers.

As we did for the FY 2010-based IPPS market basket (78 FR 50597), we are proposing to allocate contract labor costs to the Wages and Salaries and Employee Benefits cost weights based on their relative proportions for employed labor under the assumption that contract labor costs are comprised of both wages and salaries and employee benefits. The contract labor allocation proportion for wages and salaries is equal to the Wages and Salaries cost weight as a percent of the sum of the Wages and Salaries cost weight and the Employee Benefits cost weight. Using the 2014 Medicare cost report data, this percentage is 78 percent. Therefore, we are proposing to allocate approximately 78 percent of the Contract Labor cost weight to the Wages and Salaries cost weight and 22 percent to the Employee Benefits cost weight.

### TABLE IV–02—WAGES AND SALARIES AND EMPLOYEE BENEFITS COST WEIGHTS AFTER CONTRACT LABOR ALLOCATION

<table>
<thead>
<tr>
<th>Major cost categories</th>
<th>FY 2010-based IPPS market basket</th>
<th>Proposed 2014-based IPPS market basket</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wages and Salaries</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Employee Benefits</td>
<td>47.2</td>
<td>43.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>13.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>12.4</td>
</tr>
</tbody>
</table>

c. Derivation of the Detailed Cost Weights

To further divide the “All Other” residual cost weight estimated from the 2014 Medicare cost report data into more detailed cost categories, we are proposing to use the 2007 Benchmark I-O “Use Tables/Before Redefinitions/Purchaser Value” for NAICS 622000, Hospitals, published by the BEA. These data are publicly available at the following Web site: http://www.bea.gov/industry/io_annual.htm. The BEA Benchmark I-O data are generally scheduled for publication every 5 years on a lagged basis, with the most recent data available for 2007. The 2007 Benchmark I-O data are derived from the 2007 Economic Census and are the building blocks for BEA’s economic accounts. Therefore, they represent the most comprehensive and complete set of data on the economic processes or mechanisms by which output is produced and distributed. BEA also produces Annual I-O estimates. However, while based on a similar methodology, these estimates reflect less comprehensive and less detailed data sources and are subject to revision when benchmark data become available. Instead of using the less detailed Annual I-O data, we are proposing to

inflate the detailed 2007 Benchmark I-O data forward to 2014 by applying the annual price changes from the respective price proxies to the appropriate market basket cost categories that are obtained from the 2007 Benchmark I-O data. In our calculations for this proposed rule, we repeated this practice for each year. We then calculated the cost shares that each cost category represents of the 2007 data inflated to 2014. These resulting 2014 cost shares were applied to the “All Other” residual cost weight to obtain the detailed cost weights for the proposed 2014-based IPPS market basket. For example, the cost for Food: Direct Purchases represents 7.3 percent of the sum of the “All Other” 2007 Benchmark I-O Hospital Expenditures inflated to 2014. Therefore, the Food: Direct Purchases cost weight represents 7.3 percent of the proposed 2014-based IPPS market basket’s “All Other” cost category (32.0 percent), yielding a Food: Direct Purchases proposed cost weight of 2.3 percent in the proposed 2014-based IPPS market basket (0.073 × 32.0 percent = 2.3 percent). For the FY 2010-based IPPS market basket (78 FR 50597), we used the same methodology utilizing the 2002 Benchmark I-O data (aged to FY 2010).

Using this methodology, we are proposing to derive 18 detailed cost categories from the proposed 2014-based IPPS market basket residual cost weight (32.0 percent). These categories are: (1) Fuel: Oil and Gas; (2) Electricity; (3) Water and Sewerage; (4) Food: Direct Purchases; (5) Food: Contract Services; (6) Chemicals; (7) Medical Instruments; (8) Rubber and Plastics; (9) Paper and Printing Products; (10) Miscellaneous Products; (11) Professional Fees: Labor-Related; (12) Administrative and Facilities Support Services; (13) Installation, Maintenance, and Repair Services; (14) All Other: Labor-Related Services; (15) Professional Fees: Nonlabor-Related; (16) Financial Services; (17) Telephone Services; and (18) All Other: Nonlabor-Related Services.

Similar to the 2013-based LTCH market basket, the proposed 2014-based IPPS market basket does not include separate cost categories for Apparel, Machinery and Equipment, and Postage. Due to the small weights associated with these detailed categories and relatively stable price growth in the applicable price proxy, we believe that consolidating these smaller cost categories with other cost categories in the proposed market basket that experience similar price increases eliminates unnecessary complexity to the market basket without having a material impact on the total market basket increase. Therefore, we are proposing to include Apparel and Machinery and Equipment in the Miscellaneous Products cost category and Postage in the All-Other: Nonlabor-Related Services cost category. We note that the machinery and equipment expenses are for equipment that is paid for in a given year and not depreciated over the asset’s useful life. Depreciation expenses for movable equipment are reflected in the proposed 2014-based Capital Input Price Index (described in section IV.D. of the preamble of this proposed rule). For the proposed 2014-based IPPS market basket, we also are proposing to include a separate cost category for Installation, Maintenance, and Repair Services in order to proxy these costs by a price index that better reflects the price changes of labor associated with maintenance-related services.

2. Selection of Proposed Price Proxies

After computing the proposed 2014 cost weights for the IPPS market basket, it was necessary to select appropriate wage and price proxies to reflect the rate of price change for each expenditure category. With the exception of the proxy for professional liability insurance (PLI), all the proxies we are proposing are based on Bureau of Labor Statistics (BLS) data and are grouped into one of the following BLS categories:

- Producer Price Indexes—Producer Price Indexes (PPIs) measure price changes for goods sold in markets other than the retail market. PPIs are preferable price proxies for goods and services that hospitals purchase as inputs because PPIs better reflect the actual price changes encountered by hospitals. For example, we are proposing to use a PPI for prescription drugs, rather than the Consumer Price Index (CPI) for prescription drugs, because hospitals generally purchase drugs directly from a wholesaler. The PPIs that we are proposing to use measure price changes at the final stage of production.

- Consumer Price Indexes—Consumer Price Indexes (CPIs) measure change in the prices of final goods and services bought by the typical consumer. Because they may not represent the price faced by a producer, we are proposing to use CPIs only if an appropriate PPI is not available, or if the expenditures are more like those faced by retail consumers in general rather than by purchasers of goods at the wholesale level. For example, the CPI for food purchased away from home is proposed to be used as a proxy for contracted food services.

- Employment Cost Indexes—Employment Cost Indexes (ECIs) measure the rate of change in employee wage rates and employer costs for employee benefits per hour worked. These indexes are fixed-weight indexes and strictly measure the change in wage rates and employee benefits per hour. Appropriately, they are not affected by shifts in employment mix.

We evaluated the price proxies using the criteria of reliability, timeliness, availability, and relevance. Reliability indicates that the index is based on valid statistical methods and has low sampling variability. Timeliness implies that the proxy is published regularly, preferably at least once a quarter. Availability means that the proxy is publicly available. Finally, relevance means that the proxy is applicable and representative of the cost category weight to which it is applied. We believe the proposed PPIs, CPIs, and ECIs selected meet these criteria.

Below we present a detailed explanation of the price proxies that we are proposing for each cost category weight. We note that many of the proxies that we are proposing to use for the 2014-based IPPS market basket are the same as those used for the FY 2010-based IPPS market basket.

(1) Wages and Salaries

We are proposing to use the ECI for Wages and Salaries for All Civilian Workers in Hospitals (BLS series code CIU10262200000000) to measure the price growth of this cost category. This is the same price proxy used in the FY 2010-based IPPS market basket.

(2) Employee Benefits

We are proposing to use the ECI for Total Benefits for All Civilian Workers in Hospitals to measure the price growth of this cost category. This ECI is calculated using the ECI for Total Compensation for All Civilian Workers in Hospitals (BLS series code CIU10162200000000) and the relative importance of wages and salaries within total compensation. This is the same price proxy used in the FY 2010-based IPPS market basket.

(3) Fuel: Oil and Gas

We are proposing to change the proxy used for the Fuel: Oil and Gas cost category. The FY 2010-based IPPS market basket uses the PPI Industry for Petroleum Refineries (BLS series code PCU32411–32441) to proxy these expenses.

For the proposed 2014-based IPPS market basket, we are proposing to use a blend of the PPI Industry for Petroleum Refineries (BLS series code
Input-Output data shows an
price proxy for the Medical Instruments
basket.

Therefore, we are proposing a blended
proxy of 70 percent of the PPI Industry
for Petroleum Refineries (BLS series
code PCU32411–32411–) and 30 percent
of the PPI Commodity for Natural Gas
(BLS series code WPU0531). We believe
that these two price proxies are the most
technically appropriate indices available to measure the price growth of the Fuel: Oil and Gas cost category as obtained from the 2007 Benchmark I–O data. Table IV–03 below shows the proposed weights for each of the four PPIs used to create the blended index compared to those used for the FY 2010-based IPPS market basket.

<table>
<thead>
<tr>
<th>Name</th>
<th>FY 2010-based IPPS weights (%)</th>
<th>Proposed 2014-based IPPS weights (%)</th>
<th>NAICS</th>
</tr>
</thead>
<tbody>
<tr>
<td>PPI for Industrial Gas Manufacturing</td>
<td></td>
<td>35</td>
<td>32</td>
</tr>
<tr>
<td>PPI for Other Basic Inorganic Chemical Manufacturing</td>
<td></td>
<td>25</td>
<td>17</td>
</tr>
<tr>
<td>PPI for Other Basic Organic Chemical Manufacturing</td>
<td></td>
<td>30</td>
<td>45</td>
</tr>
<tr>
<td>PPI for Soap and Cleaning Compound Manufacturing</td>
<td></td>
<td>10</td>
<td>6</td>
</tr>
</tbody>
</table>

(11) Blood and Blood Products

We are proposing to use the PPI Industry for Blood and Organ Banks (BLS series code PCU621991621991) to measure the price growth of this cost category. This is the same price proxy used in the FY 2010-based IPPS market basket.

(12) Medical Instruments

We are proposing to use a blended
price proxy for the Medical Instruments cost category. The 2007 Benchmark Input-Output data shows an approximate 50/50 split between Surgical and Medical Instruments and Medical and Surgical Appliances and Supplies for this cost category, Therefore, we are proposing a blend composed of 50 percent of the PPI Commodity for Surgical and Medical Instruments (BLS series code WPU1562) and 50 percent of the PPI Commodity for Medical and Surgical Appliances and Supplies (BLS series code WPU1563). The FY 2010-based IPPS market basket used the single, higher level PPI Commodity for Medical, Surgical, and Personal Aid Devices (BLS series code WPU156). We believe that the proposed price proxy better reflects the mix of expenses for this cost category as obtained from the 2007 Benchmark I–O data.

(13) Rubber and Plastics

We are proposing to use the PPI Commodity for Rubber and Plastic Products (BLS series code WPU07) to measure the price growth of this cost category. This is the single price proxy used in the FY 2010-based IPPS market basket.
TABLE IV–04—PROPOSED 2014-BASED IPPS MARKET BASKET COST CATEGORIES, COST WEIGHTS, AND PRICE PROXIES COMPARED TO FY 2010-BASED IPPS MARKET BASKET COST WEIGHTS

<table>
<thead>
<tr>
<th>Cost categories</th>
<th>FY 2010-based IPPS market basket cost weights</th>
<th>Proposed 2014-based IPPS market basket cost weights</th>
<th>Proposed 2014-based IPPS market basket price proxies</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Compensation</td>
<td>60.3</td>
<td>55.8</td>
<td>ECI for Wages and Salaries for All Civilian Workers in Hospitals.</td>
</tr>
<tr>
<td>A. Wages and Salaries 1</td>
<td>47.2</td>
<td>43.4</td>
<td>ECI for Total Benefits for All Civilian Workers in Hospitals.</td>
</tr>
<tr>
<td>B. Employee Benefits 1</td>
<td>13.1</td>
<td>12.4</td>
<td>Blend of PPIs for Petroleum Refineries and Natural Gas.</td>
</tr>
<tr>
<td>2. Utilities</td>
<td>2.2</td>
<td>2.5</td>
<td>PPI Commodity for Commercial Electric Power.</td>
</tr>
<tr>
<td>A. Fuel: Oil and Gas</td>
<td>0.4</td>
<td>1.3</td>
<td>CPI for Water and Sewerage Maintenance (All Urban Consumers).</td>
</tr>
<tr>
<td>B. Electricity</td>
<td>1.7</td>
<td>1.0</td>
<td></td>
</tr>
<tr>
<td>C. Water and Sewerage</td>
<td>0.1</td>
<td>0.1</td>
<td></td>
</tr>
<tr>
<td>3. Professional Liability Insurance</td>
<td>1.3</td>
<td>1.2</td>
<td>CMS Hospital Professional Liability Insurance Premium Index.</td>
</tr>
<tr>
<td>4. All Other</td>
<td>36.1</td>
<td>40.5</td>
<td></td>
</tr>
<tr>
<td>A. All Other Products</td>
<td>19.5</td>
<td>17.4</td>
<td>PPI Commodity for Pharmaceuticals for Human Use, Prescription.</td>
</tr>
<tr>
<td>(1.) Pharmaceuticals</td>
<td>5.4</td>
<td>5.9</td>
<td></td>
</tr>
<tr>
<td>(2.) Food: Direct Purchases</td>
<td>4.2</td>
<td>2.3</td>
<td>PPI Commodity for Processed Foods and Feeds.</td>
</tr>
<tr>
<td>(14) Paper and Printing Products</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(15) Miscellaneous Products</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(18) Installation, Maintenance, and</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Repair Services</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(21) Financial Services</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
TABLE IV–04—PROPOSED 2014-BASED IPPS MARKET BASKET COST CATEGORIES, COST WEIGHTS, AND PRICE PROXIES COMPARED TO FY 2010-BASED IPPS MARKET BASKET COST WEIGHTS—Continued

<table>
<thead>
<tr>
<th>Cost categories</th>
<th>FY 2010-based IPPS market basket cost weights</th>
<th>Proposed 2014-based IPPS market basket cost weights</th>
<th>Proposed 2014-based IPPS market basket price proxies</th>
</tr>
</thead>
<tbody>
<tr>
<td>(3) Food: Contract Services ..................</td>
<td>0.6</td>
<td>1.3</td>
<td>CPI for Food Away From Home (All Urban Consumers).</td>
</tr>
<tr>
<td>(4) Chemicals ........................................</td>
<td>1.5</td>
<td>0.9</td>
<td>Blend of Chemical PPIs.</td>
</tr>
<tr>
<td>(5) Blood and Blood Products ..................</td>
<td>1.1</td>
<td>0.8</td>
<td>PPI Industry for Blood and Organ Banks.</td>
</tr>
<tr>
<td>(6) Medical Instruments .........................</td>
<td>2.6</td>
<td>2.9</td>
<td>Blend of PPI for Surgical and Medical Instruments and PPI for Medical and Surgical Appliances and Supplies.</td>
</tr>
<tr>
<td>(7) Rubber and Plastics .........................</td>
<td>1.6</td>
<td>0.8</td>
<td>PPI Commodity for Rubber and Plastic Products.</td>
</tr>
<tr>
<td>(8) Paper and Printing Products ..................</td>
<td>1.5</td>
<td>1.5</td>
<td>PPI Commodity for Converted Paper and Paperboard Products.</td>
</tr>
<tr>
<td>(9) Miscellaneous Products ² ..................</td>
<td>1.0</td>
<td>1.1</td>
<td>PPI Commodity for Finished Goods less Food and Energy.</td>
</tr>
<tr>
<td>B. Labor-Related Services .........................</td>
<td>9.2</td>
<td>12.5</td>
<td>ECI for Total Compensation for Private Industry Workers in Professional and Related.</td>
</tr>
<tr>
<td>(1) Professional Fees: Labor-Related ........</td>
<td>5.5</td>
<td>6.8</td>
<td>ECI for Total Compensation for Private Industry Workers in Professional and Related.</td>
</tr>
<tr>
<td>(2) Administrative and Facilities Support Services</td>
<td>0.6</td>
<td>1.0</td>
<td>ECI for Total Compensation for Private Industry Workers in Office and Administrative Support.</td>
</tr>
<tr>
<td>(3) Installation, Maintenance and Repair Services</td>
<td>2.4</td>
<td>2.4</td>
<td>ECI for Total Compensation for Civilian Workers in Installation, Maintenance, and Repair.</td>
</tr>
<tr>
<td>(4) All Other: Labor-Related Services ........</td>
<td>3.1</td>
<td>2.3</td>
<td>ECI for Total Compensation for Private Industry Workers in Service Occupations.</td>
</tr>
<tr>
<td>C. Nonlabor-Related Services .....................</td>
<td>7.4</td>
<td>10.7</td>
<td>ECI for Total Compensation for Private Industry Workers in Professional and Related.</td>
</tr>
<tr>
<td>(1) Professional Fees: Nonlabor-Related ....</td>
<td>3.7</td>
<td>5.1</td>
<td>ECI for Total Compensation for Private Industry Workers in Professional and Related.</td>
</tr>
<tr>
<td>(2) Financial Services .............................</td>
<td>1.2</td>
<td>3.0</td>
<td>ECI for Total Compensation for Private Industry Workers in Financial Activities.</td>
</tr>
<tr>
<td>(3) Telephone Services ............................</td>
<td>0.6</td>
<td>0.8</td>
<td>CPI for Telephone Services.</td>
</tr>
<tr>
<td>(4) All Other: Nonlabor-Related Services ³ ....</td>
<td>1.9</td>
<td>1.7</td>
<td>CPI for All Items less Food and Energy.</td>
</tr>
<tr>
<td>Total ..................................................</td>
<td>100.0</td>
<td>100.0</td>
<td></td>
</tr>
</tbody>
</table>

Note: The cost weights are calculated using three decimal places. For presentational purposes, we are displaying one decimal and therefore, the detail may not add to the total due to rounding.

¹ Contract labor is distributed to wages and salaries and employee benefits based on the share of total compensation that each category represents.
² The FY 2010-based IPPS market basket Miscellaneous Products cost category also includes Apparel and Machinery and Equipment cost categories. These costs were not broken out separately in the 2014-based IPPS market basket.
³ The FY 2010-based IPPS market basket All Other: Nonlabor-Related Services cost category also includes the Postage cost category. These costs were not broken-out separately in the 2014-based IPPS market basket.

Table IV–05 below compares both the historical and forecasted percent changes in the FY 2010-based IPPS market basket and the proposed 2014-based IPPS market basket. The forecasted growth rates in Table IV–05 are based on IHS Global Insight, Inc.’s (IGI) fourth quarter 2016 forecast with historical data through third quarter 2016.

TABLE IV–05.—FY 2010-BASED AND PROPOSED 2014-BASED IPPS HOSPITAL OPERATING INDEX PERCENT CHANGE, FY 2013 THROUGH FY 2020

<table>
<thead>
<tr>
<th>Fiscal Year (FY)</th>
<th>FY 2010-based IPPS market basket percent change</th>
<th>Proposed 2014-based IPPS market basket percent change</th>
</tr>
</thead>
<tbody>
<tr>
<td>Historical data:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>FY 2013</td>
<td>2.0</td>
<td>2.0</td>
</tr>
<tr>
<td>FY 2014</td>
<td>1.8</td>
<td>1.8</td>
</tr>
<tr>
<td>FY 2015</td>
<td>1.8</td>
<td>1.6</td>
</tr>
<tr>
<td>FY 2016</td>
<td>1.7</td>
<td>1.7</td>
</tr>
<tr>
<td>Average FYs 2013–2016</td>
<td>1.8</td>
<td>1.8</td>
</tr>
<tr>
<td>Forecast:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>FY 2017</td>
<td>2.6</td>
<td>2.7</td>
</tr>
<tr>
<td>FY 2018</td>
<td>2.9</td>
<td>2.9</td>
</tr>
<tr>
<td>FY 2019</td>
<td>3.0</td>
<td>3.0</td>
</tr>
<tr>
<td>FY 2020</td>
<td>3.0</td>
<td>3.0</td>
</tr>
</tbody>
</table>
There is no difference between the average percent change in the FY 2010-based IPPS market basket over the FY 2013 through FY 2016 time period. For FY 2018, the increase is 2.9 percent for both the FY 2010-based and proposed 2014-based IPPS market baskets.

3. Labor-Related Share

Under section 1886(d)(3)(E) of the Act, the Secretary estimates from time to time the proportion of payments that are labor-related. Section 1886(d)(3)(E) of the Act states that the Secretary shall adjust the proportion, (as estimated by the Secretary from time to time) of hospitals’ costs which are attributable to wages and wage-related costs, of the DRG prospective payment rates. We refer to the proportion of hospitals’ costs that are attributable to wages and wage-related costs as the “labor-related share.”

The labor-related share is used to determine the proportion of the national PPS base payment rate to which the area wage index is applied. We include a cost category in the labor-related share if the costs are labor intensive and vary with the local labor market. For the FY 2018 IPPS/LTCH PPS proposed rule, we are proposing to include in the labor-related share the national average proportion of operating costs that are attributable to the following cost categories in the proposed 2014-based IPPS market basket: Wages and Salaries, Employee Benefits, Professional Fees: Labor-Related, Administrative and Facilities Support Services, Installation, Maintenance, and Repair Services, and All Other: Labor-Related Services, as we did in the FY 2014 IPPS/LTCH PPS final rule (78 FR 50594). As noted in section IV.B.1.c. of the preamble of this proposed rule, for the proposed 2014-based IPPS market basket, we are proposing the creation of a separate cost category for Installation, Maintenance, and Repair Services. These expenses were previously included in the All Other: Labor-Related Services cost category in the FY 2010-based IPPS market basket, along with other services, including, but not limited to, janitorial, waste management, security, and dry cleaning/laundry services. Because these services tend to be labor-intensive and are mostly performed at the facility (and, therefore, unlikely to be purchased in the national market), we continue to believe that they meet our definition of labor-related services.

Similar to the FY 2010-based IPPS market basket, we are proposing that the Professional Fees: Labor-Related cost category includes expenses associated with advertising and a proportion of legal services, accounting and auditing, engineering, management consulting, and management of companies and enterprises expenses. As was done in the FY 2010-based IPPS market basket rebasing, we are proposing to determine the proportion of legal, accounting and auditing, engineering, and management consulting services that meet our definition of labor-related services based on a survey of hospitals conducted by CMS in 2008. We notified the public of our intent to conduct this survey on December 9, 2005 (70 FR 73250) and received no comments (71 FR 8588).

A discussion of the composition of the survey and poststratification can be found in the FY 2010 IPPS/LTCH PPS final rule (74 FR 43850 through 43856). Based on the weighted results of the survey, we determined that hospitals purchase, on average, the following portions of contracted professional services outside of their local labor market:

- 34 percent of accounting and auditing services;
- 30 percent of engineering services;
- 33 percent of legal services; and
- 42 percent of management consulting services.

We are proposing to apply each of these percentages to its respective Benchmark I–O cost category underlying the professional fees cost category. This is the methodology that we used to separate the FY 2010-based IPPS market basket professional fees cost category into Professional Fees: Labor-Related and Professional Fees: Nonlabor-Related cost categories. We are proposing to use the same methodology and survey results to separate the professional fees costs for the 2014-based IPPS market basket into Professional Fees: Labor-Related and Professional Fees: Nonlabor-Related cost categories. We believe these survey results are appropriate to use for the 2014-based IPPS market basket as they empirically determine the proportion of contracted professional services purchased by the industry that is attributable to local firms and the proportion that is purchased from national firms.

In the proposed 2014-based IPPS market basket, nonmedical professional fees that were subject to allocation based on these survey results represent 5.9 percent of total operating costs (and are limited to those fees related to Accounting & Auditing, Legal, Engineering, and Management Consulting services). Based on our survey results, we are proposing to apportion 3.1 percentage points of the 4.9 percentage point figure into the Professional Fees: Labor-Related share cost category and Designating the remaining 1.8 percentage point into the Professional Fees: Nonlabor-Related cost category.

In addition to the professional services listed earlier, we also classify a proportion of the home office expenses into the Professional Fees: Labor-Related cost category as was done in the previous rebasing. For the FY 2010-based IPPS market basket, we obtained home office expenses from the Benchmark I–O data for the NAICS 55 industry (Management of Companies and Enterprises). As stated in section IV.B.1.a. of the preamble to this proposed rule, for the 2014-based IPPS market basket, we are proposing to obtain these data from the Medicare cost reports. We believe that many of the home office costs are labor-intensive and vary with the local labor market. However, data indicate that not all IPPS hospitals with home offices have home offices located in their local labor market. Therefore, we are proposing to include in the labor-related share only a proportion of the home office expenses based on the methodology described below.
For the FY 2010-based IPPS market basket, we used data primarily from the Medicare cost reports and a CMS database of Home Office Medicare Records (HOMER) (a database that provides city and state information [addresses] for home offices). We determined the proportion of costs that should be allocated to the labor-related share based on the percent of hospital home office compensation as reported in Worksheet S–3, Part II. Using this methodology, we determined that 62 percent of hospitals’ home office compensation costs were for home offices located in their respective local labor markets (defined as the same Metropolitan Statistical Area (MSA)). Therefore, we classified 62 percent of these costs into the Professional Fees: Labor-Related Services cost category and the remaining 38 percent into the Professional Fees: Nonlabor-Related Services cost category for the FY 2010-based IPPS market basket. For a detailed discussion of this analysis, we refer readers to the FY 2014 IPPS/LTCH PPS final rule (78 FR 50601).

For the proposed 2014-based IPPS market basket, we conducted a similar analysis of home office data. For consistency, we believe that it is important for our analysis on home office data to be conducted on the same IPPS hospitals used to derive the proposed 2014-based IPPS market basket cost weights. The Medicare cost report requires a hospital to report information regarding their home office provider. Approximately 64 percent of IPPS hospitals reported some type of home office information on their Medicare cost report for 2014 (for example, city, State, and zip code). Using the data reported on the Medicare cost report, we compared the location of the hospital with the location of the hospital’s home office. We then determined the proportion of costs that should be allocated to the labor-related share based on the percent of total hospital home office compensation costs for those hospitals that had home offices located in their respective local labor markets—defined as being in the same MSA. We determined a hospital’s and home office’s MSAs using their zip code information from the Medicare cost report.

Similar to the FY 2010-based IPPS market basket, we determined the proportion of costs that should be allocated to the labor-related share based on the percent of hospital home office compensation as reported in Worksheet S–3, Part II. Using this methodology, we determined that 60 percent of hospitals’ home office compensation costs were for home offices located in their respective local labor markets. Therefore, we are proposing to allocate 60 percent of home office expenses to the labor-related share.

### Table IV–06—Comparison of the FY 2010-Based Labor-Related Share and the Proposed 2014-Based Labor-Related Share

<table>
<thead>
<tr>
<th>Service Category</th>
<th>FY 2010-based IPPS market basket cost weights</th>
<th>Proposed 2014-based IPPS market basket cost weights</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wages and Salaries</td>
<td>47.2</td>
<td>43.4</td>
</tr>
<tr>
<td>Employee Benefits</td>
<td>13.1</td>
<td>12.4</td>
</tr>
<tr>
<td>Professional Fees: Labor-Related</td>
<td>5.5</td>
<td>6.8</td>
</tr>
<tr>
<td>Administrative and Facilities Support Services</td>
<td>0.6</td>
<td>1.0</td>
</tr>
<tr>
<td>Installation, Maintenance, and Repair Services</td>
<td></td>
<td>2.4</td>
</tr>
<tr>
<td>All Other: Labor-Related Services</td>
<td>3.1</td>
<td>2.3</td>
</tr>
<tr>
<td>Total Labor-Related Services</td>
<td>69.6</td>
<td>68.3</td>
</tr>
</tbody>
</table>

Note: Detail may not add to total due to rounding.

1. Installation, Maintenance, and Repair Services costs were previously included in the All Other: Labor-Related Services cost category of the FY 2010-based IPPS market basket.

Using the cost category weights from the proposed 2014-based IPPS market basket, we calculated a labor-related share of 68.3 percent, approximately 1.3 percentage points lower than the current labor-related share of 69.6 percent. Therefore, we are proposing to use a labor-related share of 68.3 percent for discharges occurring on or after October 1, 2017. We continue to believe, as we have stated in the past, that these operating cost categories are related to, influenced by, or vary with the local markets. Therefore, our definition of the labor-related share continues to be consistent with section 1886(d)(3) of the Act. We note that section 403 of Pub. L. 108–173 amended sections 1886(d)(3)(E) and 1886(d)(9)(C)(iv) of the Act to provide that the Secretary must employ 62 percent as the labor-related share unless 62 percent would result in lower payments to a hospital than would otherwise be made.

### C. Market Basket for Certain Hospitals Presently Excluded From the IPPS

In the FY 2010 IPPS/RY 2010 LTCH PPS final rule (74 FR 43857), we adopted the use of the FY 2006-based IPPS operating market basket percentage...
increase to update the target amounts for children’s hospitals, PPS-excluded cancer hospitals and religious nonmedical health care institutions (RNHClS). Children’s hospitals and PPS-excluded cancer hospitals and RNHClS are still reimbursed solely under the reasonable cost-based system, subject to the rate-of-increase limits. Under these limits, an annual target amount (expressed in terms of the inpatient operating cost per discharge) is set for each hospital based on the hospital’s own historical cost experience trended forward by the applicable rate-of-increase percentages.

In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50603), under the broad authority in sections 1886(b)(3)(A) and (B), 1886(b)(3)(E), and 1871 of the Act and section 4454 of the BBA, consistent with our use of the IPPS operating market basket percentage increase to update target amounts, we adopted the use of the FY 2010-based IPPS operating market basket percentage increase to update the target amounts for children’s hospitals, PPS-excluded cancer hospitals, and RNHClS that are paid on the basis of reasonable cost subject to the rate-of-increase limits under § 413.40. In addition, as discussed in the FY 2015 IPPS/LTCH PPS final rule (79 FR 50156 through 50157), consistent with §§ 412.23(g), 413.40(a)(2)(ii)(A), and 413.40(c)(3)(viii), we also have used the percentage increase in the FY 2010-based IPPS operating market basket to update the target amounts for short-term acute care hospitals located outside the 50 States, the District of Columbia, and Puerto Rico that are paid on the basis of reasonable cost subject to the rate-of-increase limits under § 413.40.

Due to the small number of children’s and cancer hospitals and RNHClS and hospitals located outside the 50 States, the District of Columbia, and Puerto Rico and because these facilities provide limited Medicare cost report data, we are unable to create a separate market basket specifically for these facilities. Due to the limited cost report data available, we believe that the proposed 2014-based IPPS operating market basket most closely represents the cost structure of children’s hospitals, PPS-excluded cancer hospitals, RNHClS, and hospitals located outside the 50 States, the District of Columbia, and Puerto Rico. We believe this is appropriate as the IPPS operating market basket would reflect the input price growth for providing inpatient hospital services (similar to the services provided by the above excluded facilities) based on the specific mix of goods and services required. Therefore, we are proposing to use the 2014-based IPPS market basket percentage increase to update the target amounts for children’s hospitals, PPS-excluded cancer hospitals, RNHClS, and hospitals located outside the 50 States, the District of Columbia, and Puerto Rico that are paid on the basis of reasonable cost subject to the rate-of-increase limits under § 413.40. We believe it is the best available measure of the average increase in the prices of the goods and services purchased by children’s hospitals, the cancer hospitals, RNHClS, and hospitals located outside the 50 States, the District of Columbia, and Puerto Rico in order to provide care.

D. Rebasings and Revising the Capital Input Price Index (CIPI)

The CIPI was originally described in the FY 1993 IPPS final rule (57 FR 40016). There have been subsequent discussions and CIPI presented in the IPPS proposed and final rules. The FY 2014 IPPS/LTCH PPS final rule (78 FR 50603 through 50607) described the most recent rebasing and revision of the CIPI to a FY 2010 base year, which reflected the capital cost structure of IPPS hospitals available at that time.

For the FY 2018 IPPS update, we are proposing to rebase and revise the CIPI to a 2014 base year to reflect a more current structure of capital costs for IPPS hospitals. This proposed 2014-based CIPI was derived using 2014 cost reports for IPPS hospitals, which includes providers whose cost reporting period began on or after October 1, 2013, and prior to September 30, 2014. While we proposed and finalized the title of the current CIPI in the FY 2014 IPPS/LTCH proposed and final rules as “FY 2010-based CIPI”, for the proposed CIPI, we are now proposing to simply refer to the proposed CIPI as “2014-based CIPI” (dropping the reference to FY). As discussed in section IV.B. of the preamble of this proposed rule, for the 2014-based IPPS operating market basket, we are proposing this change in naming convention for the market basket because the base year cost weight data for the proposed market basket do not reflect only fiscal year data. Similarly, the proposed 2014-based CIPI uses Medicare cost report data and other government data that reflect 2014 fiscal year, 2014 calendar year, and 2014 State fiscal year expenses to determine the base year cost weights and vintage weights. Given that it is based on a mix of cost report data, we are proposing to refer to the CIPI as “2014-based” instead of “FY 2014-based” or “CY 2014-based”. However, the methods and data used to derive each of these CIPI are similar. As with the FY 2010-based index, we are proposing to develop two sets of weights to derive the proposed 2014-based CIPI. The first set of weights identifies the proportion of hospital capital expenditures attributable to each expenditure category, while the second set of weights is a set of relative vintage weights for depreciation and interest. The set of vintage weights is used to identify the proportion of capital expenditures within a cost category that is attributable to each year over the useful life of the capital assets in that category. A more thorough discussion of vintage weights is provided later in this section.

Using 2014 Medicare cost reports, we are able to group capital costs into the following categories: Depreciation, Interest, Lease, and Other. For each of these categories, we are proposing to determine what proportion of total capital costs the category represents using the data reported by IPPS hospitals on Worksheet A–7, which is the same methodology used for the FY 2010-based CIPI. As shown in the left column of Table IV–07, in 2014, depreciation expenses accounted for 66.4 percent of total capital costs, interest expenses accounted for 16.3 percent, leasing expenses accounted for 11.8 percent, and other capital expenses accounted for 5.5 percent.

We also are proposing to allocate lease costs across each of the remaining capital cost categories as was done in the FY 2010-based CIPI. This would result in three primary capital cost categories in the proposed 2014-based CIPI: Depreciation, Interest, and Other. Lease costs are unique in that they are not broken out as a separate cost category in the proposed 2014-based CIPI. Rather, we are proposing to proportionally distribute leasing costs among the cost categories of Depreciation, Interest, and Other, reflecting the assumption that the underlying cost structure of leases is similar to that of capital costs in general. As was done for the FY 2010-based CIPI, we are proposing to assume that 10 percent of the lease costs as a proportion of total capital costs represents overhead and to assign those costs to the Other capital cost category accordingly. Therefore, we are assuming that approximately 1.2 percent (11.8 percent x 0.1) of total capital costs represent lease costs attributable to overhead, and we are proposing to add this 1.2 percent to the 5.5 percent Other cost category weight. We are then proposing to distribute the remaining lease costs...
Finally, we are proposing to further divide the Depreciation and Interest cost categories. We are proposing to separate the Depreciation cost category into the following two categories: (1) Building and Fixed Equipment and (2) Movable Equipment. We also are proposing to separate the Interest cost category into the following two categories: (1) Government/Nonprofit; and (2) For-profit.

To disaggregate the depreciation cost weight, we needed to determine the percent of total depreciation costs for IPPS hospitals (after the allocation of lease costs) that are attributable to building and fixed equipment, which we hereafter refer to as the “fixed percentage.” Based on Worksheet A–7 data from the 2014 IPPS Medicare cost reports, we have determined that depreciation costs for building and fixed equipment account for approximately 49 percent of total depreciation costs, while depreciation costs for movable equipment account for approximately 51 percent of total depreciation costs. As was done for the FY 2010-based CIPI, we are proposing to apply this fixed percentage to the depreciation cost weight (after leasing costs are included) to derive a Depreciation cost weight attributable to Building and Fixed Equipment and a Deprecation cost weight attributable to Movable Equipment.

To disaggregate the interest cost weight, we needed to determine the percent of total interest costs for IPPS hospitals that are attributable to government and nonprofit facilities, which we hereafter refer to as the “nonprofit percentage,” because interest price pressures tend to differ between nonprofit and for-profit facilities. We are proposing to use interest costs data from Worksheet A–7 of the 2014 Medicare cost reports for IPPS hospitals, which is the same methodology used for the FY 2010-based CIPI. The nonprofit percentage determined using this method is 86 percent. Table IV–08 provides a comparison of the FY 2010-based CIPI cost weights and the proposed 2014-based CIPI cost weights.

After the capital cost category weights were computed, it was necessary to select appropriate price proxies to reflect the rate-of-increase for each expenditure category. We are proposing to apply the same price proxies as were used in the FY 2010-based CIPI, which are listed below and provided in Table IV–08. We also are proposing to continue to measure the price growth of capital-related costs, we are proposing to continue to measure the price growth using the average yield on municipal bonds (Bond Buyer 20-bond index) and the average yield on Moody’s Aaa bonds (Federal Reserve), respectively. As stated above, we are proposing two proxies because interest price pressures tend to differ between nonprofit and for-profit facilities. For the Other capital cost category (including insurances, taxes, and other capital-related costs), we are proposing to continue to measure the price growth using the CPI for Rent of Primary Residence (All Urban Consumers) (BLS series code CUUS0000SEHA), which would reflect the price growth of these costs. We believe that these price proxies continue to be the most appropriate proxies for IPPS capital costs that meet our selection criteria of relevance, timeliness, availability, and reliability.

### Table IV–07—Proposed Allocation of Lease Expenses for the Proposed 2014-Based CIPI

<table>
<thead>
<tr>
<th>Cost categories</th>
<th>Proposed cost shares obtained from medicare cost reports (percent of total capital costs)</th>
<th>Proposed cost shares after allocation of lease expenses (percent of total capital costs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depreciation</td>
<td>66.4</td>
<td>74.4</td>
</tr>
<tr>
<td>Interest</td>
<td>16.3</td>
<td>18.2</td>
</tr>
<tr>
<td>Lease</td>
<td>11.8</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>5.5</td>
<td>7.4</td>
</tr>
</tbody>
</table>

Note: The percentage determined using this method is 86 percent. The resulting proposed Other cost weight is 7.4 percent (5.5 percent + 1.2 percent + 0.7 percent). This is the same methodology used for the FY 2010-based CIPI. The resulting cost weights of the proposed allocation of lease expenses are shown in the right column of Table IV–07.
Because capital is acquired and paid for over time, capital expenses in any given year are determined by both past and present purchases of physical and financial capital. The proposed vintage-weighted 2014-based CIPI is intended to capture the long-term consumption of capital, using vintage weights for depreciation (physical capital) and interest (financial capital). These vintage weights reflect the proportion of capital purchases attributable to each year of the expected life of building and fixed equipment, movable equipment, and interest. We are proposing to use vintage weights to compute vintage-weighted price changes associated with depreciation and interest expenses.

Vintage weights are an integral part of the CIPI. Capital costs are inherently complicated and are determined by complex capital purchasing decisions, over time, based on such factors as interest rates and debt financing. In addition, capital is depreciated over time instead of being consumed in the same period it is purchased. By accounting for the vintage nature of capital, we are able to provide an accurate and stable annual measure of price changes. Annual nonvintage price changes for capital are unstable due to the volatility of interest rate changes and, therefore, do not reflect the actual annual price changes for IPPS capital costs. The CIPI reflects the underlying stability of the capital acquisition process.

To calculate the vintage weights for depreciation and interest expenses, we first needed a time series of capital purchases for building and fixed equipment and movable equipment. We found no single source that provides an appropriate time series of capital purchases by hospitals for all of the above categories of capital purchases. The early Medicare cost reports did not have sufficient capital data to meet this need. Data we obtained from the American Hospital Association (AHA) did not include annual capital purchases. However, we were able to obtain data on total expenses back to 1963 from the AHA. Consequently, we are proposing to use data from the AHA Panel Survey and the AHA Annual Survey to obtain a time series of total expenses for hospitals. We then are proposing to use data from the AHA Panel Survey supplemented with the ratio of depreciation to total hospital expenses obtained from the Medicare cost reports to derive a trend of annual depreciation expenses for 1963 through 2014. We are proposing to separate these depreciation expenses into annual amounts of building and fixed equipment depreciation and movable equipment depreciation as determined earlier. From these annual depreciation amounts, we derived annual end-of-year book values for building and fixed equipment and movable equipment using the expected life for each type of asset category. We used the AHA data and similar methodology to derive the FY 2010-based IPPS capital market basket (78 FR 50604).

To continue to calculate the vintage weights for depreciation and interest expenses, we also needed to account for the expected lives for building and fixed equipment, movable equipment, and interest for the proposed 2014-based CIPI. We are proposing to calculate the proposed rule. For the interest vintage weights should represent the average expected life of building and fixed equipment because, based on previous research described in the FY 1997 IPPS final rule (61 FR 46198), the expected life of hospital debt instruments and the expected life of buildings and fixed equipment are similar. We note that the FY 2010-based CIPI was based on an expected average life of building and fixed equipment of 26 years and an expected average life of movable equipment of 12 years.

Multiplying these expected lives by the annual depreciation amounts results in annual year-end asset costs for building and fixed equipment and movable equipment. We then calculated a time series, beginning in 1964, of annual capital purchases by subtracting the previous year’s asset costs from the current year’s asset costs. For the building and fixed equipment and movable equipment vintage weights, we are proposing to use the real annual capital-related purchase amounts for each asset type to capture the actual amount of the physical acquisition, net of the effect of price inflation. These real annual capital-related purchase amounts are produced by deflating the nominal annual purchase amount by the associated price proxy as provided earlier in this proposed rule. For the interest vintage weights, we are proposing to use the total nominal annual capital-related purchase amounts to capture the value of the debt instrument (including, but not limited to, mortgages and bonds). Using these capital purchases time series specific to each asset type, we are proposing to calculate the vintage weights for building and fixed equipment.
equipment, for movable equipment, and for interest.

The vintage weights for each asset type are deemed to represent the average purchase pattern of the asset over its expected life (in the case of building and fixed equipment and interest, 27 years, and in the case of movable equipment, 12 years). For each asset type, we are proposing to use the time series of annual capital purchases amounts available from 2014 back to 1964. These data allow us to derive twenty-five 27-year periods of capital purchases for building and fixed equipment and interest, and forty 12-year periods of capital purchases for movable equipment. For each 27-year period for building and fixed equipment and interest, or 12-year period for movable equipment, we are proposing to calculate annual vintage weights by dividing the capital-related purchase amount in any given year by the total amount of purchases over the entire 27-year or 12-year period. This calculation was done for each year in the 27-year or 12-year period and for each of the periods for which we have data. We then calculated the average vintage weight for a given year of the expected life by taking the average of these vintage weights across the multiple periods of data.

The vintage weights for the proposed 2014-based CIPI and the FY 2010-based CIPI are presented in Table IV–09 below.

### Table IV–09—Proposed 2014-Based CIPI and FY 2010-Based CIPI Vintage Weights

<table>
<thead>
<tr>
<th>Year</th>
<th>Proposed 2014-based 27 years</th>
<th>FY 2010-based 26 years</th>
<th>Proposed 2014-based 12 years</th>
<th>FY 2010-based 12 years</th>
<th>Proposed 2014-based 27 years</th>
<th>FY 2010-based 26 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.024</td>
<td>0.023</td>
<td>0.062</td>
<td>0.064</td>
<td>0.012</td>
<td>0.012</td>
</tr>
<tr>
<td>2</td>
<td>0.025</td>
<td>0.024</td>
<td>0.064</td>
<td>0.066</td>
<td>0.014</td>
<td>0.013</td>
</tr>
<tr>
<td>3</td>
<td>0.027</td>
<td>0.026</td>
<td>0.070</td>
<td>0.071</td>
<td>0.015</td>
<td>0.015</td>
</tr>
<tr>
<td>4</td>
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<td>0.028</td>
<td>0.074</td>
<td>0.073</td>
<td>0.017</td>
<td>0.017</td>
</tr>
<tr>
<td>5</td>
<td>0.030</td>
<td>0.029</td>
<td>0.078</td>
<td>0.076</td>
<td>0.019</td>
<td>0.018</td>
</tr>
<tr>
<td>6</td>
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<td>0.031</td>
<td>0.082</td>
<td>0.078</td>
<td>0.021</td>
<td>0.021</td>
</tr>
<tr>
<td>7</td>
<td>0.033</td>
<td>0.032</td>
<td>0.086</td>
<td>0.084</td>
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<td>0.023</td>
</tr>
<tr>
<td>8</td>
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<td>0.034</td>
<td>0.088</td>
<td>0.088</td>
<td>0.025</td>
<td>0.025</td>
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<tr>
<td>9</td>
<td>0.035</td>
<td>0.036</td>
<td>0.092</td>
<td>0.092</td>
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<td>10</td>
<td>0.036</td>
<td>0.038</td>
<td>0.097</td>
<td>0.098</td>
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<td>0.030</td>
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<tr>
<td>11</td>
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<td>0.103</td>
<td>0.030</td>
<td>0.033</td>
</tr>
<tr>
<td>12</td>
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<td>0.041</td>
<td>0.105</td>
<td>0.106</td>
<td>0.033</td>
<td>0.036</td>
</tr>
<tr>
<td>13</td>
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<td>0.042</td>
<td></td>
<td></td>
<td>0.035</td>
<td>0.038</td>
</tr>
<tr>
<td>14</td>
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<td></td>
<td>0.037</td>
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<tr>
<td>15</td>
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<td>0.043</td>
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<tr>
<td>16</td>
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<tr>
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<td></td>
<td>0.048</td>
<td>0.051</td>
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<tr>
<td>20</td>
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<td>0.044</td>
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<td></td>
<td>0.050</td>
<td>0.052</td>
</tr>
<tr>
<td>21</td>
<td>0.043</td>
<td>0.045</td>
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<td></td>
<td>0.052</td>
<td>0.056</td>
</tr>
<tr>
<td>22</td>
<td>0.043</td>
<td>0.045</td>
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<td></td>
<td>0.054</td>
<td>0.057</td>
</tr>
<tr>
<td>23</td>
<td>0.042</td>
<td>0.045</td>
<td></td>
<td></td>
<td>0.055</td>
<td>0.060</td>
</tr>
<tr>
<td>24</td>
<td>0.042</td>
<td>0.046</td>
<td></td>
<td></td>
<td>0.057</td>
<td>0.062</td>
</tr>
<tr>
<td>25</td>
<td>0.043</td>
<td>0.045</td>
<td></td>
<td></td>
<td>0.059</td>
<td>0.064</td>
</tr>
<tr>
<td>26</td>
<td>0.043</td>
<td>0.045</td>
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<td></td>
<td>0.061</td>
<td>0.066</td>
</tr>
<tr>
<td>27</td>
<td>0.043</td>
<td>0.045</td>
<td></td>
<td></td>
<td>0.062</td>
<td>1.000</td>
</tr>
<tr>
<td>Total</td>
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<td>1.000</td>
<td>1.000</td>
<td>1.000</td>
<td>1.000</td>
<td>1.000</td>
</tr>
</tbody>
</table>

**Note:** Numbers may not add to total due to rounding.

1 Vintage weight in the last year (for example, year 27 for the proposed 2014-based CIPI) is applied to the most recent data point and prior vintage weights are applied going back in time. For example, year 27 vintage weight would be applied to the 2018q3 fixed price proxy level, year 26 vintage weight would be applied to the 2017q3 fixed price proxy level, etc.

The process of creating vintage-weighted price proxies requires applying the vintage weights to the price proxy index where the last applied vintage weight in Table IV–09 is applied to the most recent data point. We have provided on the CMS Web site an example of how the vintage weighting price proxies are calculated, using example vintage weights and example price indices. The example can be found under the following CMS Web site link: [http://www.cms.gov/Research-Statistics-Data-and-Systems/Statistics-Trends-and-Reports/MedicareProgramRatesStats/MarketBasketResearch.html](http://www.cms.gov/Research-Statistics-Data-and-Systems/Statistics-Trends-and-Reports/MedicareProgramRatesStats/MarketBasketResearch.html) in the zip file titled “Weight Calculations as described in the IPPS FY 2010 Proposed Rule.”

Table IV–10 below compares both the historical and forecasted percent changes in the FY 2010-based CIPI and the proposed 2014-based CIPI.
IHS Global Insight, Inc. forecasts a 1.2 percent increase in the proposed 2014-based CIPI for FY 2018, as shown in Table IV–10. The underlying vintage-weighted price increases for depreciation (including building and fixed equipment and movable equipment) and interest (including government/nonprofit and for-profit) are included on the proposed 2014-based CIPI.

TABLE IV–11—PROPOSED 2014-BASED CAPITAL INPUT PRICE INDEX PERCENT CHANGES, TOTAL AND DEPRECIATION AND INTEREST COMPONENTS—FYs 2013 THROUGH 2020

<table>
<thead>
<tr>
<th>Fiscal year</th>
<th>Total</th>
<th>Depreciation</th>
<th>Interest</th>
</tr>
</thead>
<tbody>
<tr>
<td>FY 2013</td>
<td>1.0</td>
<td>1.7</td>
<td>-2.5</td>
</tr>
<tr>
<td>FY 2014</td>
<td>1.2</td>
<td>1.8</td>
<td>-1.8</td>
</tr>
<tr>
<td>FY 2015</td>
<td>1.1</td>
<td>1.8</td>
<td>-2.7</td>
</tr>
<tr>
<td>FY 2016</td>
<td>1.0</td>
<td>1.7</td>
<td>-3.0</td>
</tr>
<tr>
<td>FY 2017</td>
<td>1.0</td>
<td>1.6</td>
<td>-2.7</td>
</tr>
<tr>
<td>FY 2018</td>
<td>1.2</td>
<td>1.6</td>
<td>-1.6</td>
</tr>
<tr>
<td>FY 2019</td>
<td>1.4</td>
<td>1.6</td>
<td>-0.6</td>
</tr>
<tr>
<td>FY 2020</td>
<td>1.5</td>
<td>1.6</td>
<td>0.1</td>
</tr>
</tbody>
</table>


Rebasing the CIPI from FY 2010 to 2014 decreased the percent change in the forecasted update for FY 2018 by 0.1 percentage point, from 1.3 percent to 1.2 percent, as shown in Table IV–10. The lower FY 2018 update is primarily due to a change in the vintage weights for the proposed 2014-based CIPI, which includes updating the asset purchase data through 2014 and changing the building and fixed equipment and interest asset lives from 26 years to 27 years. This lower update is only partially offset by the change in the base year weights, which produce a faster increase due to more weight being given to the Depreciation cost category and less weight being given to the Interest cost category. As shown in Table IV–11, for FY 2018, vintage-weighted price growth is projected to be positive for the Depreciation cost category and negative for Interest cost category.

V. Other Decisions and Proposed Changes to the IPPS for Operating System

A. Proposed Changes to MS–DRGs Subject to the Postacute Care Transfer and MS–DRG Special Payment Policies (§ 412.4)

1. Background

Existing regulations at 42 CFR 412.4(a) define discharges under the IPPS as situations in which a patient is formally released from an acute care hospital or dies in the hospital. Section 412.4(b) defines acute care transfers, and §412.4(c) defines postacute care transfers. Our policy set forth in §412.4(f) provides that when a patient is transferred and his or her length of stay is less than the geometric mean length of stay for the MS–DRG to which the case is assigned, the transferring hospital is generally paid based on a graduated per diem rate for each day of stay, not to exceed the full MS–DRG payment that would have been made if the patient had been discharged without being transferred.

The per diem rate paid to a transferring hospital is calculated by dividing the full MS–DRG payment by the geometric mean length of stay for the MS–DRG. Based on an analysis that showed that the first day of hospitalization is the most expensive (60 FR 45804), our policy generally provides for payment that is twice the per diem amount for the first day, with each subsequent day paid at the per diem amount up to the full MS–DRG payment (§ 412.4(f)(1)). Transfer cases also are eligible for outlier payments. In general, the outlier threshold for transfer cases, as described in §412.80(b), is equal to the fixed-loss outlier threshold for nontransfer cases (adjusted for geographic variations in costs), divided by the geometric mean length of stay for the MS–DRG, and multiplied by the length of stay for the case, plus 1 day.
We established the criteria set forth in §412.4(d) for determining which DRGs qualify for postacute care transfer payments in the FY 2006 IPPS final rule (70 FR 47419 through 47420). The determination of whether a DRG is subject to the postacute care transfer policy was initially based on the Medicare Version 23.0 GROUPER (FY 2006) and data from the FY 2004 MedPAR file. However, if a DRG did not exist in Version 23.0 or a DRG included in Version 23.0 is revised, we use the current version of the Medicare GROUPER and the most recent complete year of MedPAR data to determine if the DRG is subject to the postacute care transfer policy. Specifically, if the MS–DRG’s total number of discharges to postacute care equals or exceeds the 55th percentile for all MS–DRGs and the proportion of short-stay discharges to postacute care to total discharges in the MS–DRG exceeds the 55th percentile for all MS–DRGs, CMS will apply the postacute care transfer policy to that MS–DRG and to any other MS–DRG that shares the same base MS–DRG. The statute directs us to identify MS–DRGs based on a high volume of discharges to postacute care facilities and a disproportionate use of postacute care services. As discussed in the FY 2006 IPPS final rule (70 FR 47416), we determined that the 55th percentile is an appropriate level at which to establish these thresholds. In that same final rule (70 FR 47419), we stated that we will not revise the list of DRGs subject to the postacute care transfer policy annually unless we are making a change to a specific MS–DRG.

To account for MS–DRGs subject to the postacute care policy that exhibit exceptionally higher shares of costs very early in the hospital stay, §412.4(f) also includes a special payment methodology. For these MS–DRGs, hospitals receive 50 percent of the full MS–DRG payment, plus the single per diem payment, for the first day of the stay, as well as a per diem payment for subsequent days (up to the full MS–DRG payment [§412.4(f)(6)]. For an MS–DRG to qualify for the special payment methodology, the geometric mean length of stay must be greater than 4 days, and the average charges of 1-day discharge cases in the MS–DRG must be at least 50 percent of the average charges for all cases within the MS–DRG. MS–DRGs that are part of an MS–DRG severity level group will qualify under the MS–DRG special payment methodology policy if any one of the MS–DRGs that share that same base MS–DRG qualifies (§412.4(f)(6)).

2. Proposed Changes for FY 2018

Based on our annual review of MS–DRGs, we have identified three MS–DRGs that we are proposing to be included on the list of MS–DRGs subject to the special payment transfer policy. As we discuss in section II.F. of the preamble of this proposed rule, in response to public comments and based on our analysis of FY 2016 MedPAR claims data, we are proposing to make changes to MS–DRGs, effective for FY 2018. As discussed in section II.F.14.b. of the preamble of this proposed rule, we are proposing to delete MS–DRGs 984, 985, and 986 (Prostatic O.R. Procedure Unrelated to Principal Diagnosis with MCC, with CC and without CC/MCC, respectively) and reassign the procedure codes currently assigned to these three MS–DRGs to MS–DRGs 987, 988, and 989 (Non-Extensive O.R. Procedure Unrelated to Principal Diagnosis with MCC, with CC and without CC/MCC, respectively).

In light of these proposed changes to the MS–DRGs for FY 2018, according to the regulations under §412.4(d), we evaluated proposed revised MS–DRGs 987, 988, and 989 (which would contain the proposed reassigned procedures from MS–DRGs 984, 985, and 986) against the general postacute care transfer policy criteria using the FY 2016 MedPAR data. If an MS–DRG qualified for the postacute care transfer policy, we also evaluated that MS–DRG under the special payment methodology criteria according to regulations at §412.4(f)(6). We continue to believe it is appropriate to reassess MS–DRGs when proposing realignment or procedure or diagnosis codes that would result in material changes to an MS–DRG. MS–DRGs 987, 988, and 989 are currently subject to the postacute care transfer policy. As a result of our review, the proposed revised MS–DRGs 987, 988, and 989 continue to qualify to be included on the list of MS–DRGs that are subject to the postacute care transfer policy. We are not proposing to change the postacute care transfer policy status for MS–DRGs 987, 988, and 989.

### LIST OF PROPOSED REvised MS–DRGs SUBJECT TO REVIEW OF POSTACUTE CARE TRANSFER POLICY STATUS FOR FY 2018

<table>
<thead>
<tr>
<th>Proposed revised MS–DRG</th>
<th>MS–DRG title</th>
<th>Total cases</th>
<th>Postacute care transfers (55th percentile: 1,419)</th>
<th>Short-stay postacute care transfers</th>
<th>Percent of short-stay postacute care transfers to all cases (55th percentile: 8.01068%)</th>
<th>Postacute care transfer policy status</th>
</tr>
</thead>
<tbody>
<tr>
<td>987</td>
<td>Non-Extensive O.R. Procedure Unrelated to Principal Diagnosis with MCC.</td>
<td>8,131</td>
<td>4,210</td>
<td>1,355</td>
<td>16.66462</td>
<td>YES.</td>
</tr>
<tr>
<td>988</td>
<td>Non-Extensive O.R. Procedure Unrelated to Principal Diagnosis with CC.</td>
<td>8,239</td>
<td>3,416</td>
<td>706</td>
<td>8.56900</td>
<td>YES.</td>
</tr>
<tr>
<td>989</td>
<td>Non-Extensive O.R. Procedure Unrelated to Principal Diagnosis without MCC/CC.</td>
<td>2,216</td>
<td>*499</td>
<td>47</td>
<td>*2.12094</td>
<td>**YES.</td>
</tr>
</tbody>
</table>

* Indicates a current postacute care transfer policy criterion that the MS–DRG did not meet.

** As described in the policy at 42 CFR 412.4(d)(3)(ii)(D), MS–DRGs that share the same base MS–DRG will all qualify under the postacute care transfer policy if any one of the MS–DRGs that share that same base MS–DRG qualifies.

We also have determined that proposed revised MS–DRGs 987, 988, and 989 would meet the criteria for the MS–DRG special payment methodology. MS–DRGs 987, 988, and 989 are not currently listed as being subject to the special payment policy. Therefore, we are proposing that these three proposed revised MS–DRGs would be subject to...
the MS–DRG special payment methodology, effective FY 2018.

**LIST OF PROPOSED REVISED MS–DRGs SUBJECT TO REVIEW OF SPECIAL PAYMENT POLICY STATUS FOR FY 2018**

<table>
<thead>
<tr>
<th>Proposed revised MS–DRG</th>
<th>MS–DRG title</th>
<th>Geometric mean length of stay</th>
<th>Average charges of 1-day discharges</th>
<th>50 Percent of average charges for all cases within MS–DRG</th>
<th>Special payment policy status</th>
</tr>
</thead>
<tbody>
<tr>
<td>987</td>
<td>Non-Extensive O.R. Procedure Unrelated to Principal Diagnosis with MCC.</td>
<td>8.1</td>
<td>$36,526</td>
<td>$53,449</td>
<td>*YES.</td>
</tr>
<tr>
<td>988</td>
<td>Non-Extensive O.R. Procedure Unrelated to Principal Diagnosis with CC.</td>
<td>8.6</td>
<td>35,629</td>
<td>29,119</td>
<td>YES.</td>
</tr>
<tr>
<td>989</td>
<td>Non-Extensive O.R. Procedure Unrelated to Principal Diagnosis without MCC/CC.</td>
<td>2.2</td>
<td>0</td>
<td>0</td>
<td>*YES.</td>
</tr>
</tbody>
</table>

*As described in the policy at 42 CFR 412.4(d)(6)(iv), MS–DRGs that share the same base MS–DRG will all qualify under the MS–DRG special payment policy if any one of the MS–DRGs that share that same base MS–DRG qualifies.

The proposed postacute care transfer policy status and special payment policy status of these MS–DRGs are reflected in Table 5 associated with this proposed rule, which is listed in section VI. of the Addendum to this proposed rule and available via the Internet on the CMS Web site.

**B. Proposed Changes in the Inpatient Hospital Update for FY 2018**

1. Proposed FY 2018 Inpatient Hospital Update

In accordance with section 1886(b)(3)(B)(i) of the Act, each year we update the national standardized amount for inpatient hospital operating costs by a factor called the “applicable percentage increase.” For FY 2018, we are setting the applicable percentage increase by applying the adjustments listed in this section in the same sequence as we did for FY 2017. Specifically, consistent with section 1886(b)(3)(B) of the Act, as amended by sections 3401(a) and 10319(a) of the Affordable Care Act, we are setting the applicable percentage increase by applying the following adjustments in the following sequence. The applicable percentage increase under the IPPS is equal to the rate-of-increase in the hospital market basket for IPPS hospitals in all areas, subject to—

(a) A reduction of one-quarter of the applicable percentage increase (prior to the application of other statutory adjustments; also referred to as the market basket update or rate-of-increase (with no adjustments)) for hospitals not considered to be meaningful EHR users in accordance with section 1886(b)(3)(B)(ix) of the Act;

(b) A reduction of three-quarters of the applicable percentage increase (prior to the application of other statutory adjustments; also referred to as the market basket update or rate-of-increase (with no adjustments)) for hospitals that fail to submit quality information under rules established by the Secretary in accordance with section 1886(b)(3)(B)(viii) of the Act (hereafter referred to as a hospital that submits quality data) and is a meaningful EHR user under section 1886(b)(3)(B)(ix) of the Act (hereafter referred to as a hospital that is a meaningful EHR user), there are four possible applicable percentage increases that can be applied to the standardized amount as specified in the table that appears later in this section.

We note that, in compliance with section 404 of the MMA, in this proposed rule, we are proposing to replace the FY 2010-based IPPS operating and capital market baskets with the revised and rebased 2014-based IPPS operating and capital market baskets for FY 2018. We are proposing to base the proposed FY 2018 market basket update used to determine the applicable percentage increase for the IPPS on IHS Global Insight, Inc.’s ([IGI’s](https://www.bls.gov/mfp)) fourth quarter 2016 forecast of the proposed 2014-based IPPS market basket rate-of-increase with historical data through third quarter 2016, which is estimated to be 2.9 percent. We are proposing that if more recent data subsequently become available (for example, a more recent estimate of the market basket and the MFP adjustment), we would use such data, if appropriate, to determine the FY 2018 market basket update and the MFP adjustment in the final rule.

For FY 2018, depending on whether a hospital submits quality data under the rules established in accordance with section 1886(b)(3)(B)(viii) of the Act (hereafter referred to as a hospital that submits quality data) and is a meaningful EHR user under section 1886(b)(3)(B)(ix) of the Act (hereafter referred to as a hospital that is a meaningful EHR user), there are four possible applicable percentage increases that can be applied to the standardized amount as specified in the table that appears later in this section.

In the FY 2012 IPPS/LTCH PPS final rule (76 FR 51689 through 51692), we finalized our methodology for calculating and applying the MFP adjustment. As we explained in that rule, section 1886(b)(3)(B)(xii) of the Act, as added by section 3401(a) of the Affordable Care Act, state that application of the MFP adjustment and the additional FY 2018 adjustment of 0.75 percentage point may result in the applicable percentage increase being less than zero.

We note that, in compliance with section 404 of the MMA, in this proposed rule, we are proposing to replace the FY 2010-based IPPS operating and capital market baskets with the revised and rebased 2014-based IPPS operating and capital market baskets for FY 2018. We are proposing to base the proposed FY 2018 market basket update used to determine the applicable percentage increase for the IPPS on IHS Global Insight, Inc.’s ([IGI’s](https://www.bls.gov/mfp)) fourth quarter 2016 forecast of the proposed 2014-based IPPS market basket rate-of-increase with historical data through third quarter 2016, which is estimated to be 2.9 percent. We are proposing that if more recent data subsequently become available (for example, a more recent estimate of the market basket and the MFP adjustment), we would use such data, if appropriate, to determine the FY 2018 market basket update and the MFP adjustment in the final rule.

For FY 2018, depending on whether a hospital submits quality data under the rules established in accordance with section 1886(b)(3)(B)(viii) of the Act (hereafter referred to as a hospital that submits quality data) and is a meaningful EHR user under section 1886(b)(3)(B)(ix) of the Act (hereafter referred to as a hospital that is a meaningful EHR user), there are four possible applicable percentage increases that can be applied to the standardized amount as specified in the table that appears later in this section.

In the FY 2012 IPPS/LTCH PPS final rule (76 FR 51689 through 51692), we finalized our methodology for calculating and applying the MFP adjustment. As we explained in that rule, section 1886(b)(3)(B)(xii) of the Act, as added by section 3401(a) of the Affordable Care Act, state that application of the MFP adjustment and the additional FY 2018 adjustment of 0.75 percentage point may result in the applicable percentage increase being less than zero.

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For FY 2018, depending on whether a hospital submits quality data under the rules established in accordance with section 1886(b)(3)(B)(viii) of the Act (hereafter referred to as a hospital that submits quality data) and is a meaningful EHR user under section 1886(b)(3)(B)(ix) of the Act (hereafter referred to as a hospital that is a meaningful EHR user), there are four possible applicable percentage increases that can be applied to the standardized amount as specified in the table that appears later in this section.

In the FY 2012 IPPS/LTCH PPS final rule (76 FR 51689 through 51692), we finalized our methodology for calculating and applying the MFP adjustment. As we explained in that rule, section 1886(b)(3)(B)(xii) of the Act, as added by section 3401(a) of the Affordable Care Act, state that application of the MFP adjustment and the additional FY 2018 adjustment of 0.75 percentage point may result in the applicable percentage increase being less than zero.

We note that, in compliance with section 404 of the MMA, in this proposed rule, we are proposing to replace the FY 2010-based IPPS operating and capital market baskets with the revised and rebased 2014-based IPPS operating and capital market baskets for FY 2018. We are proposing to base the proposed FY 2018 market basket update used to determine the applicable percentage increase for the IPPS on IHS Global Insight, Inc.’s ([IGI’s](https://www.bls.gov/mfp)) fourth quarter 2016 forecast of the proposed 2014-based IPPS market basket rate-of-increase with historical data through third quarter 2016, which is estimated to be 2.9 percent. We are proposing that if more recent data subsequently become available (for example, a more recent estimate of the market basket and the MFP adjustment), we would use such data, if appropriate, to determine the FY 2018 market basket update and the MFP adjustment in the final rule.

For FY 2018, depending on whether a hospital submits quality data under the rules established in accordance with section 1886(b)(3)(B)(viii) of the Act (hereafter referred to as a hospital that submits quality data) and is a meaningful EHR user under section 1886(b)(3)(B)(ix) of the Act (hereafter referred to as a hospital that is a meaningful EHR user), there are four possible applicable percentage increases that can be applied to the standardized amount as specified in the table that appears later in this section.

In the FY 2012 IPPS/LTCH PPS final rule (76 FR 51689 through 51692), we finalized our methodology for calculating and applying the MFP adjustment. As we explained in that rule, section 1886(b)(3)(B)(xii) of the Act, as added by section 3401(a) of the Affordable Care Act, state that application of the MFP adjustment and the additional FY 2018 adjustment of 0.75 percentage point may result in the applicable percentage increase being less than zero.
We are proposing to revise the existing regulations at 42 CFR 412.64(d) to reflect the current law for the FY 2018 update. Specifically, in accordance with section 1886(b)(3)(B) of the Act, we are proposing to revise paragraph (vii) of § 412.64(d)(1) to include the applicable percentage increase to the FY 2018 operating standardized amount as the percentage increase in the market basket index, subject to the reductions specified under § 412.64(d)(2) for a hospital that does not submit quality data and § 412.64(d)(3) for a hospital that is not a meaningful EHR user, less an MFP adjustment and less an additional reduction of 0.75 percentage point.

Section 1886(b)(3)(B)(iv) of the Act provides that the applicable percentage increase to the hospital-specific rates for SCHs equals the applicable percentage increase set forth in section 1886(b)(3)(B)(i) of the Act (that is, the same update factor as for all other hospitals subject to the IPPS). Therefore, the update to the hospital-specific rates for SCHs also is subject to section 1886(b)(3)(B)(i) of the Act, as amended by sections 3401(a) and 10319(a) of the Affordable Care Act.

As discussed in section V.H. of the preamble of this proposed rule, section 205 of the Medicare Access and CHIP Reauthorization Act of 2015 (MACRA) (Pub. L. 114–10, enacted on April 16, 2015) extended the MDH program (which, under previous law, was to be in effect for discharges on or before March 31, 2015 only) for discharges occurring on or after April 1, 2015, through FY 2017 (that is, for discharges occurring on or before September 30, 2017). Therefore, under current law, the MDH program will expire at the end of FY 2017.

For FY 2018, we are proposing the following updates to the hospital-specific rates applicable to SCHs: A proposed update of 1.75 percent for a hospital that submits quality data and is a meaningful EHR user; a proposed update of 1.025 percent for a hospital that fails to submit quality data and is a meaningful EHR user; a proposed update of −0.425 percent for a hospital that submits quality data and is not a meaningful EHR user; and a proposed update of −1.15 percent for a hospital that fails to submit quality data and is not a meaningful EHR user. As mentioned previously, for this FY 2018 proposed rule, we are using ICI’s fourth quarter 2016 forecast of the proposed 2014-based IPPS market basket update with historical data through third quarter 2016. Similarly, we are using ICI’s fourth quarter 2016 forecast of the MFP adjustment. We are proposing that if more recent data subsequently become available, we would use such data, if appropriate, to determine the update in the final rule.

Based on these data, for this proposed rule, we have determined four proposed applicable percentage increases to the standardized amount for FY 2018, as specified in the following table:

<table>
<thead>
<tr>
<th>Proposed Market Basket Rate-of-Increase</th>
<th>Hospital submitted quality data and is a meaningful EHR user</th>
<th>Hospital submitted quality data and is NOT a meaningful EHR user</th>
<th>Hospital did NOT submit quality data and is a meaningful EHR user</th>
<th>Hospital did NOT submit quality data and is NOT a meaningful EHR user</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proposed Adjustment for Failure to Submit Quality Data under Section 1886(b)(3)(B)(viii) of the Act</td>
<td>2.9</td>
<td>2.9</td>
<td>2.9</td>
<td>2.9</td>
</tr>
<tr>
<td>Proposed Adjustment for Failure to be a Meaningful EHR User under Section 1886(b)(3)(B)(x) of the Act</td>
<td>0.0</td>
<td>0.0</td>
<td>−0.725</td>
<td>−0.725</td>
</tr>
<tr>
<td>Proposed MFP Adjustment under Section 1886(b)(3)(B)(xi) of the Act</td>
<td>−0.4</td>
<td>−0.4</td>
<td>−0.4</td>
<td>−0.4</td>
</tr>
<tr>
<td>Statutory Adjustment under Section 1886(b)(3)(B)(xii) of the Act</td>
<td>−0.75</td>
<td>−0.75</td>
<td>−0.75</td>
<td>−0.75</td>
</tr>
<tr>
<td>Proposed Applicable Percentage Increase Applied to Standardized Amount</td>
<td>1.75</td>
<td>−0.425</td>
<td>1.025</td>
<td>−1.15</td>
</tr>
</tbody>
</table>

2. Proposed FY 2018 Puerto Rico Hospital Update

As discussed in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56937 through 56938), prior to January 1, 2016, Puerto Rico hospitals were paid based on 75 percent of the national standardized amount and 25 percent of the Puerto Rico-specific standardized amount. Section 601 of Public Law 114–113 amended section 1886(d)(9)(E) of the Act to specify that the payment calculation with respect to operating costs of inpatient hospital services of a subsection (d) Puerto Rico hospital for inpatient hospital discharges on or after January 1, 2016, shall use 100 percent of the national standardized amount. Because Puerto Rico hospitals are no longer paid with a Puerto Rico-specific standardized amount under the amendments to section 1886(d)(9)(E) of the Act, there is no longer a need for us to propose an update to the Puerto Rico standardized amount. Hospitals in Puerto Rico are now paid 100 percent of the national standardized amount and, therefore, are subject to the same update to the national standardized amount discussed under section V.B.1. of the preamble of this proposed rule. Accordingly, for FY 2018, we are proposing an applicable percentage increase of 1.75 to the standardized amount for hospitals located in Puerto Rico.
applicable percentage increase for “subsection (d)” hospitals that do not submit quality data under the rules established by the Secretary, is not applicable to hospitals located in Puerto Rico.

In addition, section 602 of Public Law 114–113 amended section 1886(n)(6)(B) of the Act to specify that Puerto Rico hospitals are eligible for incentive payments for the meaningful use of certified EHR technology, effective beginning FY 2016, and also to apply the adjustments to the applicable percentage increase under section 1886(b)(3)(B)(ix) of the Act to Puerto Rico hospitals that are not meaningful EHR users, effective FY 2022. Accordingly, because the provisions of section 1886(b)(3)(B)(ix) of the Act are not applicable to hospitals located in Puerto Rico until FY 2022, the adjustments under this provision are not applicable for FY 2018.

C. Proposed Change to Volume Decrease Adjustment for Sole Community Hospitals (SCHs) and Medicare-Dependent, Small Rural Hospitals (MDHs) (§ 412.92)

1. Background

Sections 1886(d)(5) and (d)(5)(G) of the Act provide special payment protections under the IPPS to sole community hospitals (SCHs) and Medicare-dependent, small rural hospitals (MDHs), respectively. Section 1886(d)(5)(D)(ii) and section 1886(d)(5)(G)(iii) of the Act require that the Secretary adjust the payments made to an SCH and MDH, respectively, as may be necessary to fully compensate the hospital for the fixed costs it incurs in providing inpatient hospital services, including the reasonable cost of maintaining necessary core staff and services, when it experiences a decrease of more than 5 percent in its total number of inpatient discharges due to circumstances beyond its control. These adjustments are known as “volume decrease adjustments.”

The regulations governing volume decrease adjustments are found at § 412.92(e) for SCHs and § 412.108(d) for MDHs. As noted earlier, the MDH program is set to expire as of October 1, 2017. As such, we are not proposing specific amendments to the regulations at § 412.108(d) for MDHs. However, we are proposing that if the MDH program ends up being extended by law, similar to how it was extended by section 205 of the MACRA (Pub. L. 114–10) and prior legislation, the following proposed changes to the volume decrease adjustment methodology and the proposed amendment to § 412.92(e)(3) for SCHs would also be made to the parallel requirements for MDHs under § 412.108(d)(3).

To qualify for a volume decrease adjustment, the SCH must: (a) Submit documentation demonstrating the size of the decrease in discharges and the resulting effect on per discharge costs; and (b) show that the decrease is due to circumstances beyond the hospital’s control. If an SCH demonstrates to the MAC’s satisfaction that it has suffered a qualifying decrease in total inpatient discharges, the MAC determines the appropriate amount, if any, due to the SCH as an adjustment.

As we have noted in the PRM and in adjudications rendered by the PRRB and the CMS Administrator, under the current methodology, the MAC determines a volume decrease adjustment amount not to exceed the cap calculated as the difference between the lesser of (1) the hospital’s current year’s Medicare inpatient operating costs or (2) its prior year’s Medicare inpatient operating costs multiplied by the appropriate IPPS update factor and the hospital’s total MS–DRG revenue for inpatient operating costs (including outlier payments, DSH payments, and IME payments). In determining the volume decrease adjustment amount, the MAC considers the individual hospital’s needs and circumstances, including the reasonable cost of maintaining necessary core staff and services in view of minimum staffing requirements imposed by State agencies; the hospital’s fixed costs (including whether any semi-fixed costs are to be considered fixed) other than those costs paid on a reasonable cost basis; and the length of time the hospital has experienced a decrease in utilization.

We have set forth interpretive guidance regarding volume decrease adjustments in the preamble to various rules and in Section 2810.1 of the Provider Reimbursement Manual, Part 1 (PRM–1). The adjustment also has been the subject of a series of adjudications, rendered by the PRRB and the CMS Administrator. For example, we refer readers to Greenwood County Hospital v. Blue Cross Blue Shield Association/Blue Cross Blue Shield of Kansas, 2006 WL 3050893 (PRRB, August 29, 2006); Unity Healthcare Muscle Shoals v. Blue Cross Blue Shield Association/ Wisconsin Physicians Service, 2014 WL 5450066 (CMS Administrator September 4, 2014); Lakes Regional Healthcare Spirit Lake, Iowa v. Blue Cross Blue Shield Association/Wisconsin Physicians Service, 2014 WL 5450078 (CMS Administrator September 4, 2014); and Fairbanks Memorial Hospital v. Wisconsin Physician Services/ BlueCross BlueShield Association, 2015 WL 5852432 (CMS Administrator, August 5, 2015). In those adjudications, the PRRB and the CMS Administrator have recognized that: (1) The volume decrease adjustment is intended to compensate qualifying SCHs for their fixed costs only, and that variable costs are to be excluded from the adjustment; and (2) an SCH’s volume decrease adjustment should be reduced to reflect the compensation of fixed costs that has already been made through MS–DRG payments.

However, some hospitals have recently expressed concerns regarding the exact calculations that the MACs use when determining the volume decrease...
adjustment. The issue also has been addressed in some recent decisions of the PRRB. Under the current calculation methodology, the MACs calculate the volume decrease adjustment by subtracting the hospital’s total MS–DRG revenue for inpatient operating costs, including outlier payments and IME and DSH payments in the cost reporting period in which the volume decrease occurred, from fixed costs in the cost reporting period in which the volume decrease occurred, minus any adjustment for excess staff. If the result of that calculation is greater than zero and less than the cap, the hospital receives that amount in a lump-sum payment. If the result of that calculation is zero or less than zero, the hospital does not receive a volume decrease payment adjustment.

Under the IPPS, MS–DRG payments are not based on an individual hospital’s actual costs in a given cost reporting period. However, the main issue raised by the PRRB and individual hospitals is that, under the current calculation methodology, if the hospital’s total MS–DRG revenue for treating Medicare beneficiaries for which it incurs inpatient operating costs (consisting of fixed, semi-fixed, and variable costs) exceeds the hospital’s fixed costs, the calculation by the MACs results in no volume decrease adjustment for the hospital. In some recent decisions, the PRRB has indicated that it believes it would be more appropriate for the MACs to adjust the hospital’s total MS–DRG revenue from Medicare by looking at the ratio of a hospital’s fixed costs to its total costs (as determined by the MAC) and applying that ratio as a proxy for the share of the hospital’s MS–DRG payments that it assumes are attributable (or allocable) to fixed costs, and then comparing that estimate of the fixed portion of MS–DRG payments to the hospital’s fixed costs. In this way, the calculation would compare estimated Medicare revenue for fixed costs to the hospital’s fixed costs when determining the volume decrease adjustment.

We continue to believe that our current approach in calculating volume decrease adjustments is reasonable and consistent with the statute. Nevertheless, we understand why hospitals might take the view that CMS should make an effort, in some way, to ascertain whether a portion of MS–DRG payments can be allocated or attributed to fixed costs in order to fulfill the statutory mandate to “fully compensate” a qualifying SCH for its fixed costs.

Accordingly, after considering these views, we are proposing to prospectively change how the MACs calculate the volume decrease adjustments and require that the MACs compare estimated Medicare revenue for fixed costs to the hospital’s fixed costs to remove any conceivable possibility that a hospital that qualifies for the volume decrease adjustment could ever be less than fully compensated for fixed costs as a result of the application of the adjustment. We are proposing that, in order to estimate the fixed portion of the Medicare revenue, the MACs would apply the ratio of the hospital’s fixed costs to total costs in the cost reporting period when it experienced the volume decrease to the hospital’s total Medicare revenue in that same cost reporting period. We are proposing to revise the regulations at 42 CFR 412.92(e)(3) to reflect our proposed change in the MAC’s calculation of the volume decrease adjustment that would apply prospectively to cost reporting periods beginning on or after October 1, 2017, and to reflect that the language requiring that the volume decrease adjustment amount not exceed the difference between the hospital’s Medicare inpatient operating costs and the hospital’s total DRG revenue for inpatient operating costs would only apply to cost reporting periods beginning before October 1, 2017, but not to subsequent cost reporting periods. Under the proposed methodology, if a hospital’s total MS–DRG payment is less than its total Medicare inpatient operating costs, the sum of any resulting volume decrease adjustment payment and its MS–DRG payment would never exceed its total Medicare inpatient operating costs due to the fact that the fixed cost percentage is applied to the MS–DRG payment in calculating the volume decrease adjustment amount. By taking the ratio derived from the subset of fixed costs to total costs and applying that same ratio to the MS–DRG payment, we ensure that the sum of a hospital’s IPPS payment and its volume decrease adjustment payment would never exceed its total Medicare inpatient operating costs, thus negating the need for a cap calculation. Thus, the proposed methodology renders the current volume decrease adjustment cap calculation obsolete. Conversely, if a hospital’s total MS–DRG payment is greater than its total Medicare inpatient operating costs, calculating a volume decrease adjustment using the proposed methodology would result in a negative payment amount, which would yield a volume decrease adjustment payment of zero. Finally, if a hospital’s total MS–DRG payment is equal to its total Medicare inpatient operating costs, calculating a volume decrease adjustment using the proposed methodology would also yield a volume decrease adjustment payment of zero. Furthermore, we believe that because a hospital could not foresee a decrease in its volume from one year to the next and would therefore not plan for a volume decrease adjustment, the volume decrease adjustment payment should therefore not be limited to a cap that is based on the previous year’s costs. For these reasons, we are proposing to remove the cap calculation from the volume decrease adjustment calculation methodology in future periods.

We are proposing that these proposed changes in the MAC’s calculation of the volume decrease adjustment would be prospective, effective for cost reporting periods beginning on or after October 1, 2017. If these proposed changes are adopted, we also intend to update Section 2810.1 of the PRM–1 to reflect the changes in the calculation of the volume decrease adjustment by the MAC. For volume decrease adjustments for earlier cost reporting periods, the current calculation methodology will continue. In addition, we are not proposing to change any part of the methodology, criteria, rules, or presumptions we consider and apply in determining whether to classify a given cost as fixed, semi-fixed, or variable for purposes of the volume decrease adjustment.

The following example illustrates the calculation of the volume decrease adjustment by the MAC under our proposed change.

Example: In its cost reporting period beginning October 1, 2017, Hospital A has total Medicare inpatient operating costs equaling $1,600,000 and total MS–DRG revenue (including outlier payments, IME and DSH) of $1,400,000. The MAC determines that the hospital qualifies for a volume decrease adjustment for this cost reporting period. The MAC classifies $1,360,000 of Hospital A’s costs as fixed and $240,000 as variable. Hospital A’s fixed cost ratio is therefore .85 = $1,360,000/ $1,600,000. The MAC applies this ratio to the total MS–DRG revenue of $1,400,000 to estimate the hospital’s fixed MS–DRG revenue to be $1,190,000. The volume decrease adjustment payment is then calculated by comparing the fixed MS–DRG revenue of $1,190,000 to the fixed costs of $1,360,000, resulting in a volume decrease adjustment payment of $170,000 ($1,360,000 minus $1,190,000).
Under the current methodology used by the MACs, Hospital A would receive no volume decrease adjustment payment because its total MS–DRG revenue from Medicare of $1,400,000 exceeded the hospital’s fixed costs of $1,360,000. Furthermore, under the current methodology, but not under our proposed methodology, it is possible that a hospital would still receive no volume decrease adjustment payment even if its fixed costs exceeded its total MS–DRG revenue if those fixed costs exceeded the previous year’s costs updated for inflation.

We also are proposing changes to an adjustment that might be made to a hospital’s staffing costs in calculating the volume decrease adjustment. The statute and regulations and the PRM imply, and we have expressly indicated in prior rulemaking, most recently in the FY 2006 rulemaking, our belief that not all staff costs can necessarily be considered fixed costs (71 FR 48056 through 48060). Therefore, we currently require a hospital, when applying for a volume decrease adjustment, to demonstrate that it appropriately adjusted the number of staff in inpatient areas of the hospital based on the decrease in the number of inpatient days but not beyond minimum levels as required by State or local laws. If a hospital does not appropriately adjust its number of staff, the cost of maintaining those staff members is deducted from the total volume decrease adjustment payment. In reviewing the volume decrease adjustment calculation, we have also weighed the administrative burden on the hospital of making this demonstration to CMS, as compared to an assumption that it is likely that a hospital would, in its normal course of business, adjust its staffing levels as revenue declines. In the absence of evidence to contrary, we believe that a hospital would adjust its staffing levels as revenue declines rather than maintain those staffing levels for the sole purpose of potentially having those staffing costs eventually reflected in a Medicare volume decrease adjustment payment that the hospital may or may not qualify for when it files its cost report. Therefore, we are proposing to modify the volume decrease adjustment process to no longer require that a hospital explicitly demonstrate that it appropriately adjusted the number of staff in inpatient areas of the hospital based on the decrease in the number of inpatient days and to no longer require the MAC to adjust the volume decrease adjustment payment amount for excess staffing. We are proposing that these changes be effective for cost reporting periods beginning on or after October 1, 2017.

In summary, we are proposing to prospectively modify the volume decrease adjustment process to no longer require that a hospital explicitly demonstrate that it appropriately adjusted the number of staff in inpatient areas of the hospital based on the decrease in the number of inpatient days and to no longer require the MAC to adjust the volume decrease adjustment calculation methodology. We are proposing to revise the regulations at § 412.92(e)(3) to reflect our proposed changes. We also are proposing to prospectively modify the volume decrease adjustment process to no longer require that a hospital explicitly demonstrate that it appropriately adjusted the number of staff in inpatient areas of the hospital based on the decrease in the number of inpatient days and to no longer require the MAC to adjust the volume decrease adjustment payment amount for excess staffing. We are proposing that these changes be effective for cost reporting periods beginning on or after October 1, 2017. As we noted earlier, we are proposing that if the MDH program ends up being extended by law, similar to how it was extended by section 205 of the MACRA (Pub. L. 114–10) and prior legislation, these proposed changes to the volume decrease adjustment methodology and the proposed amendment to § 412.92(e)(3) for SCHs would also be made to the parallel requirements for MDHs under § 412.108(d)(3).

D. Rural Referral Centers (RRCs):

Proposed Annual Updates to Case-Mix Index and Discharge Criteria (§ 412.98)

Under the authority of section 1886(d)(5)[C][i] of the Act, the regulations at § 412.96 set forth the criteria that a hospital must meet in order to qualify under the IPPS as a rural referral center (RRC). RRCs receive some special treatment under both the DSH payment adjustment and the criteria for geographic reclassification. Section 402 of Public Law 108–173 raised the DSH payment adjustment for RRCs such that they are not subject to the 12-percent cap on DSH payments that is applicable to other rural hospitals. RRCs also are not subject to the proximity criteria when applying for geographic reclassification. In addition, they do not have to meet the requirement that a hospital’s average hourly wage must exceed, by a certain percentage, the average hourly wage of the labor market area in which the hospital is located.

Section 4202(b) of Public Law 105–33 states, in part, that any hospital classified as an RRC by the Secretary for FY 1991 shall be classified as such an RRC for FY 1998 and each subsequent fiscal year. In the August 29, 1997 IPPS final rule with comment period (62 FR 45999), we reinstated RRC status for all hospitals that lost that status due to triennial review or MGCRB reclassification. However, we did not reinstate the status of hospitals that lost RRC status because they were now urban for all purposes because of the OMB designation of their geographic area as urban. Subsequently, in the August 1, 2000 IPPS final rule (65 FR 47089), we indicated that we were revisiting that decision. Specifically, we stated that we would permit hospitals that previously qualified as an RRC and lost their status due to OMB redesignation of the county in which they are located from rural to urban, to be reinstated as an RRC. Otherwise, a hospital seeking RRC status must satisfy all of the other applicable criteria. We use the definitions of “urban” and “rural” specified in subpart D of 42 CFR part 412. One of the criteria under which a hospital may qualify as an RRC is to have 275 or more beds available for use (§ 412.96(b)(1)(iii)). A rural hospital that does not meet the bed size requirement can qualify as an RRC if the hospital meets two mandatory prerequisites (a minimum case-mix index (CMI) and a minimum number of discharges), and at least one of three optional criteria (relating to specialty composition of medical staff, source of inpatients, or referral volume). We refer readers to § 412.96(c)(1) through (c)(5) and the September 30, 1988 Federal Register (53 FR 38513) for additional discussion.) With respect to the two mandatory prerequisites, a hospital may be classified as an RRC if—

- The hospital’s CMI is at least equal to the lower of the median CMI for urban hospitals in its census region, excluding hospitals with approved teaching programs, or the median CMI for all urban hospitals nationally; and

- The hospital’s number of discharges is at least 5,000 per year, or, if fewer, the median number of discharges for urban hospitals in the census region in which the hospital is located. The number of discharges criterion for an osteopathic hospital is at least 3,000 discharges per year, as specified in section 1886(d)(5)[C][i] of the Act.

1. Case-Mix Index (CMI)

Section 412.96(c)(1) provides that CMS establish updated national and regional CMI values in each year’s annual notice of prospective payment
rates for purposes of determining RRC status. The methodology we used to determine the national and regional CMI values is set forth in the regulations at § 412.96(c)(1)(ii). The proposed national median CMI value for FY 2018 is based on the CMI values of all urban hospitals nationwide, and the proposed regional median CMI values for FY 2018 are based on the CMI values of all urban hospitals within each census region, excluding those hospitals with approved teaching programs (that is, those hospitals that train residents in an approved GME program as provided in § 413.75). These proposed values are based on discharges occurring during FY 2016 (October 1, 2015 through September 30, 2016), and include bills posted to CMS’ records through December 2016.

In this proposed rule, we are proposing that, in addition to meeting other criteria, if rural hospitals with fewer than 275 beds are to qualify for initial RRC status for cost reporting periods beginning on or after October 1, 2017, they must have a CMI value for FY 2016 that is at least—

- 1.6635 (national—all urban); or
- The median CMI value (not transfer-adjusted) for urban hospitals (excluding hospitals with approved teaching programs as identified in § 413.75) calculated by CMS for the census region in which the hospital is located.

The proposed median CMI values by region are set forth in the following table.

<table>
<thead>
<tr>
<th>Region</th>
<th>Case-mix index value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. New England (CT, ME, MA, NH, RI, VT)</td>
<td>1.4186</td>
</tr>
<tr>
<td>2. Middle Atlantic (PA, NJ, NY)</td>
<td>1.5126</td>
</tr>
<tr>
<td>3. South Atlantic (DE, DC, FL, GA, MD, NC, SC, VA, WV)</td>
<td>1.5393</td>
</tr>
<tr>
<td>4. East North Central (IL, IN, MI, OH, WI)</td>
<td>1.5921</td>
</tr>
<tr>
<td>5. East South Central (AL, KY, MS, TN)</td>
<td>1.5179</td>
</tr>
<tr>
<td>6. West North Central (IA, KS, MN, MO, NE, ND, SD)</td>
<td>1.6346</td>
</tr>
<tr>
<td>7. West South Central (AR, LA, OK, TX)</td>
<td>1.6949</td>
</tr>
<tr>
<td>8. Mountain (AZ, CO, ID, MT, NV, NM, UT, WY)</td>
<td>1.7614</td>
</tr>
<tr>
<td>9. Pacific (AK, CA, HI, OR, WA)</td>
<td>1.8466</td>
</tr>
</tbody>
</table>

We intend to update these proposed CMI values in the FY 2018 final rule to reflect the updated FY 2016 MedPAR file, which will contain data from additional bills received through March 2017.

A hospital seeking to qualify as an RRC should obtain its hospital-specific CMI value (not transfer-adjusted) from its MAC. Data are available on the Provider Statistical and Reimbursement (PS&R) System. In keeping with our policy on discharges, the CMI values are computed based on all Medicare patient discharges subject to the IPPS MS–DRG-based payment.

2. Discharges

Section 412.96(c)(2)(i) provides that CMS set forth the national and regional numbers of discharges criteria in each year’s annual notice of prospective payment rates for purposes of determining RRC status. As specified in section 1886(d)(5)(C)(ii) of the Act, the national standard is set at 5,000 discharges. In this proposed rule, for FY 2018, we are proposing to update the regional standards based on discharges for urban hospitals’ cost reporting periods that began during FY 2015 (that is, October 1, 2014 through September 30, 2015), which are the latest cost report data available at the time this proposed rule was developed.

Therefore, we are proposing that, in addition to meeting other criteria, a hospital, if it is to qualify for initial RRC status for cost reporting periods beginning on or after October 1, 2017, must have, as the number of discharges for its cost reporting period that began during FY 2015, at least—

- 5,000 (3,000 for an osteopathic hospital); or
- The median number of discharges for urban hospitals in the census region in which the hospital is located as reflected in the following table.

<table>
<thead>
<tr>
<th>Region</th>
<th>Number of discharges</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. New England (CT, ME, MA, NH, RI, VT)</td>
<td>7,991</td>
</tr>
<tr>
<td>2. Middle Atlantic (PA, NJ, NY)</td>
<td>10,268</td>
</tr>
<tr>
<td>3. South Atlantic (DE, DC, FL, GA, MD, NC, SC, VA, WV)</td>
<td>10,503</td>
</tr>
<tr>
<td>4. East North Central (IL, IN, MI, OH, WI)</td>
<td>8,802</td>
</tr>
<tr>
<td>5. East South Central (AL, KY, MS, TN)</td>
<td>8,697</td>
</tr>
<tr>
<td>6. West North Central (IA, KS, MN, MO, NE, ND, SD)</td>
<td>7,532</td>
</tr>
<tr>
<td>7. West South Central (AR, LA, OK, TX)</td>
<td>5,189</td>
</tr>
<tr>
<td>8. Mountain (AZ, CO, ID, MT, NV, NM, UT, WY)</td>
<td>8,887</td>
</tr>
<tr>
<td>9. Pacific (AK, CA, HI, OR, WA)</td>
<td>8,856</td>
</tr>
</tbody>
</table>

We intend to update these numbers in the FY 2018 final rule based on the latest available cost report data.

We note that the median number of discharges for hospitals in each census region is greater than the national standard of 5,000 discharges. Therefore, under this proposed rule, 5,000 discharges is the minimum criterion for all hospitals, except for osteopathic hospitals for which the minimum criterion is 3,000 discharges.

E. Proposed Payment Adjustment for Low-Volume Hospitals (§ 412.101)

1. Expiration of Temporary Changes to Low-Volume Hospital Payment Policy

Under section 1886(d)(12) of the Act, as amended, most recently by section 204 of the Medicare Access and CHIP Reauthorization Act of 2015 (MACRA), Public Law 114–10, the temporary changes in the low-volume hospital payment policy originally provided by the Affordable Care Act and extended through subsequent legislation are effective through FY 2017. Beginning with FY 2018, the preexisting low-volume hospital payment adjustment and qualifying criteria, as implemented
in FY 2005 and discussed later in this section, will resume. We discuss the proposed payment policies for FY 2018 in section V.E.3. of the preamble of this proposed rule.

2. Background

Section 1886(d)(12) of the Act, as added by section 406(a) of Public Law 108–173, provides for a payment adjustment to account for the higher costs per discharge for low-volume hospital–provider, the IPPS, effective beginning FY 2005. Sections 3125 and 10314 of the Affordable Care Act amended section 1886(d)(12) of the Act by modifying the definition of a low-volume hospital and the methodology for calculating the payment adjustment for low-volume hospitals, effective only for discharges occurring during FYs 2011 and 2012. Specifically, the provisions of the Affordable Care Act amended the qualifying criteria for low-volume hospitals to specify, for FYs 2011 and 2012, that a hospital qualifies as a low-volume hospital if it is more than 15 road miles from another subsection (d) hospital and has less than 1,600 discharges of individuals entitled to, or enrolled for, benefits under Medicare Part A during the fiscal year. In addition, the statute, as amended by the Affordable Care Act, provides that the low-volume hospital payment adjustment (that is, the percentage increase) is determined using a continuous linear sliding scale ranging from 25 percent for low-volume hospitals with 200 or fewer discharges of individuals entitled to, or enrolled for, benefits under Medicare Part A in the fiscal year to 0 percent for low-volume hospitals with greater than 1,600 discharges of such individuals in the fiscal year. The temporary changes to the low-volume hospital qualifying criteria and the payment adjustment originally provided by the Affordable Care Act were extended by subsequent legislation, most recently through FY 2017 by section 204 of the MAGRA. (We refer readers to the FY 2017 IPPS/LTC PPS final rule (FR 50941 through 59943) for a detailed summary of the applicable legislation.) Under current law, beginning with FY 2018, the preexisting low-volume hospital qualifying criteria and payment adjustment, as implemented in FY 2005 and described in this section, will resume. The regulations implementing the low-volume hospital adjustment provided by section 1886(d)(12) of the Act are located at 42 CFR 412.101.

The additional payment adjustment to a low-volume hospital for under section 1886(d)(12) of the Act is in addition to any payment calculated under this section. Therefore, the additional payment adjustment is based on the per discharge amount paid to the qualifying hospital under section 1886 of the Act. In other words, the low-volume add-on payment amount is based on total per discharge payments made under section 1886 of the Act, including capital, DSH, IME, and outliers. For hospitals paid based on the hospital-specific rate, the low-volume add-on payment amount is based on either the Federal rate or the hospital-specific rate, whichever results in a greater operating IPPS payment. Section 1886(d)(12)(C)(i) of the Act defines a low-volume hospital, for fiscal years other than FYs 2011 through 2017, as a subsection (d) hospital (as defined in paragraph (1)(B)) that the Secretary determines is located more than 25 road miles from another subsection (d) hospital and that has less than 800 discharges during the fiscal year. Section 1886(d)(12)(C)(ii) of the Act further stipulates that the term “discharge” means an inpatient acute care discharge of an individual regardless of whether the individual is entitled to benefits under Medicare Part A. Therefore, for fiscal years other than FYs 2011 through 2017, the term “discharge” refers to total discharges, regardless of payer (that is, not only Medicare discharges). Furthermore, section 1886(d)(12)(B) of the Act requires, for discharges occurring in FYs 2005 through 2010 and FY 2018 and subsequent years, that the Secretary determine an applicable percentage increase for these low-volume hospitals based on the “empirical relationship” between the standardized cost-per-case for such hospitals and the total number of discharges of such hospitals and the amount of the additional incremental costs (if any) that are associated with such number of discharges. The statute thus mandates that the Secretary develop an empirically justifiable adjustment based on the relationship between costs and discharges for these low-volume hospitals. Section 1886(d)(12)(B)(iii) of the Act limits the applicable percentage increase adjustment to no more than 25 percent.

Based on an analysis we conducted for the FY 2005 IPPS final rule (69 FR 49099 through 49102), a 25-percent low-volume adjustment to all qualifying hospitals with less than 200 discharges was found to be most consistent with the statutory requirement to provide relief to low-volume hospitals where there is empirical evidence that higher incremental costs are associated with low numbers of total discharges. In the FY 2006 IPPS final rule (70 FR 47432 through 47434), we stated that multivariate analyses supported the existing low-volume adjustment implemented in FY 2005.

3. Proposed Payment Adjustment for FY 2018 and Subsequent Fiscal Years

In accordance with section 1886(d)(12) of the Act, beginning with FY 2018, the low-volume hospital definition and payment adjustment methodology will revert back to the statutory requirements that were in effect prior to the amendment made by the Affordable Care Act and extended by subsequent legislation. Therefore, effective for FY 2018 and subsequent years, in order to qualify as a low-volume hospital, a subsection (d) hospital must be more than 25 road miles from another subsection (d) hospital and have less than 200 discharges (that is, less than 800 discharges total, including both Medicare and non-Medicare discharges) during the fiscal year. As discussed earlier, the statute specifies that a low-volume hospital must have less than 800 discharges during the fiscal year. However, as required by section 1886(d)(12)(B)(i) of the Act and as discussed earlier, the Secretary has developed an empirically justifiable payment adjustment based on the relationship, for IPPS hospitals with less than 800 discharges, between the additional incremental costs (if any) that are associated with a particular number of discharges. Based on an analysis we conducted for the FY 2005 IPPS final rule (69 FR 49099 through 49102), a 25-percent low-volume adjustment to all qualifying hospitals with less than 200 discharges was found to be most consistent with the statutory requirement to provide relief for low-volume hospitals where there is empirical evidence that higher incremental costs are associated with low numbers of total discharges. Under the policy we established in that same final rule, hospitals with between 200 and 799 discharges do not receive a low-volume hospital adjustment.

As described earlier, for FYs 2005 through 2010 and FY 2018 and subsequent years, the discharge determination is made based on the hospital’s number of total discharges, that is, Medicare and non-Medicare discharges. The hospital’s most recently submitted cost report is used to determine if the hospital meets the discharge criterion to receive the low-volume payment adjustment in the current year (§ 412.101(b)(2)(i)). We use cost report data to determine if a hospital meets the discharge criterion because this is the best available data source that includes information on
both Medicare and non-Medicare discharges. We note that, for FYs 2011 through 2017, we used the most recently available MedPAR data to determine the hospital’s Medicare discharges because only Medicare discharges were used to determine if a hospital met the discharge criterion for those years.

For FY 2018 and for subsequent fiscal years, in addition to a discharge criterion, the eligibility for the low-volume payment adjustment is also dependent upon the hospital meeting the mileage criterion specified at § 412.101(b)(2)(ii). Specifically, to meet the mileage criterion to qualify for the low-volume payment adjustment for FY 2018 and subsequent fiscal years, a hospital must be located more than 25 road miles from the nearest subsection (d) hospital. We define, at § 412.101(a), the term “road miles” to mean “miles” as defined at § 412.92(c)(1)(75 FR 50238 through 50275 and 50414).

In the FY 2011 IPPS/LTCH PPS final rule (75 FR 50238 through 50275 and 50414) and subsequent rulemaking, most recently in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56942 through 56943), we discussed the process for requesting and obtaining the low-volume hospital payment adjustment. In order to qualify for the low-volume hospital payment adjustment, a hospital must provide to its MAC sufficient evidence to document that it meets the discharge and distance requirements. The MAC will determine, based on the most recent data available, if the hospital qualifies as a low-volume hospital, so that the hospital will know in advance whether or not it will receive a payment adjustment. The MAC and CMS may review available data, in addition to the data the hospital submits with its request for low-volume hospital status, in order to determine whether or not the hospital meets the qualifying criteria.

In order to receive a low-volume hospital payment adjustment under § 412.101, a hospital must notify and provide documentation to its MAC that it meets the mileage criterion. The use of a Web-based mapping tool as part of documenting that the hospital meets the mileage criterion for low-volume hospitals is acceptable. The MAC will determine if the information submitted by the hospital, such as the name and street address of the nearest hospitals, location on a map, and distance (in road miles, as defined in the regulations at § 412.101(a)) from the hospital requesting low-volume hospital status, is sufficient to document that it meets the mileage criterion. If not, the MAC will follow up with the hospital to obtain additional necessary information to determine whether or not the hospital meets the low-volume mileage criterion. In addition, the MAC will refer to the hospital’s most recently submitted cost report to determine whether or not the hospital meets the discharge criterion. A hospital should refer to its most recently submitted cost report for total discharges (Medicare and non-Medicare) in order to decide whether or not to apply for low-volume hospital status for a particular fiscal year. A hospital must continue to meet the qualifying criteria at § 412.101(b)(2)(ii) as a low-volume hospital (that is, the discharge criterion and the mileage criterion) in order to receive the payment adjustment in that year; that is, low-volume hospital status is not based on a “one-time” qualification (75 FR 50238 through 50275).

In order to be a low-volume hospital in FY 2018 and subsequent fiscal years, in accordance with our previously established procedure, a hospital must make a written request for low-volume hospital status that is received by its MAC by September 1 immediately preceding the start of the Federal fiscal year for which the hospital is applying for low-volume hospital status in order for the 25-percent, low-volume, add-on payment adjustment to be applied to payments for its discharges for the fiscal year beginning on or after October 1 immediately following the request (that is, the start of the Federal fiscal year). For a hospital whose request for low-volume hospital status is received after September 1, if the MAC determines the hospital meets the criteria to qualify as a low-volume hospital, the MAC will apply the 25-percent, low-volume, add-on payment adjustment to determine payment for the hospital’s discharges for the fiscal year, effective prospectively within 30 days of the date of the MAC’s low-volume status determination. Specifically, for FY 2018, a hospital must make a written request for low-volume hospital status that is received by its MAC no later than September 1, 2017, in order for the 25-percent, low-volume, add-on payment adjustment to be applied to payments for its discharges beginning on or after October 1, 2017 (through September 30, 2018). Under this procedure, a hospital that qualified for the low-volume hospital payment adjustment for FY 2017 may continue to receive a low-volume hospital payment adjustment for FY 2018 without reapplying if it meets both the discharge criterion and the mileage criterion applicable for FY 2018. As in previous years, we are proposing that such a hospital must send written notification that is received by its MAC no later than September 1, 2017, stating that it meets the mileage criterion applicable for FY 2018. For FY 2018, we are further proposing that this written verification must also state, based upon the most recently submitted cost report, that the hospital meets the discharge criterion applicable for FY 2018 (that is, less than 200 discharges total, including both Medicare and non-Medicare discharges). If a hospital’s request for low-volume hospital status for FY 2018 is received after September 1, 2017, and if the MAC determines the hospital meets the criteria to qualify as a low-volume hospital, the MAC will apply the 25-percent, low-volume, add-on payment adjustment to determine the payment for the hospital’s FY 2018 discharges, effective prospectively within 30 days of the date of the MAC’s low-volume hospital status determination. We note that this process mirrors our established application process but is updated to ensure that providers currently receiving the low-volume hospital payment adjustment verify that they meet both the mileage criterion and the discharge criterion applicable for FY 2018 to continue receiving the adjustment for FY 2018. For additional information on our established application process for the low-volume hospital payment adjustment, we refer readers to the FY 2017 IPPS/LTCH PPS final rule (81 FR 56942 through 56943).

In the FY 2016 IPPS interim final rule with comment period (80 FR 49594 through 49597 and 49767), we made conforming changes to the regulations at 42 CFR 412.101 to reflect the extension of the changes to the qualifying criteria and the payment adjustment methodology for low-volume hospitals through FY 2017 in accordance with section 204 of the MACRA. Under these revisions, beginning with FY 2018, consistent with current law, the low-volume hospital qualifying criteria and payment adjustment methodology will return to the criteria and methodology that were in effect prior to the amendments made by the Affordable Care Act (that is, the low-volume hospital policy in effect for FYs 2005 through 2010). Therefore, no further revisions to the policy or to the regulations at § 412.101 are required to conform them to the statutory requirement that the low-volume hospital policy in effect prior to the Affordable Care Act will again be in effect for FY 2018 and subsequent years. For this reason, we are not proposing specific amendments to the regulations at § 412.101 to reflect the expiration of the temporary changes to the low-volume hospital payment adjustment policy originally provided for by the
Affordable Care Act. However, we are proposing that if these temporary changes to the low-volume hospital payment policy were to be extended by law, similar to extensions provided through FY 2013, by the American Taxpayer Relief Act of 2012 (ATRA), Public Law 112–240; through March 31, 2014, by the Pathway for SGR Reform Act of 2013, Public Law 113–167; through March 31, 2015, by the Protecting Access to Medicare Act of 2014 (PAMA), Public Law 113–93; and most recently through FY 2017, by the Medicare Access and CHIP Reauthorization Act of 2015 (MACRA), Public Law 114–10, we would make conforming changes to the regulations at § 412.101(b) through (d), as appropriate, to reflect any such extension.

These conforming changes would only be made if the temporary changes to the low-volume hospital payment adjustment policy were to be extended by statute beyond the current expiration date of September 30, 2017. If these temporary changes were to be extended by statute, for FY 2018, consistent with our historical policy and our implementation of the prior extensions, qualifying low-volume hospitals and their payment adjustment would be determined using the most recently available Medicare discharge data available at the time of the final rule, which we expect would be from the March 2017 update of the FY 2016 MedPAR file. Consistent with past practice, if these temporary changes were to be extended for FY 2018 before the development of the final rule, we would list the subsection (d) hospitals with fewer than 1,600 Medicare discharges based on the claims data from the March 2017 update of the FY 2016 MedPAR file and their potential low-volume hospital payment adjustment for FY 2018 in Table 14 listed in the Addendum of the final rule. In such an event, hospitals would still submit requests or verification to the MAC, as outlined earlier, but updated as needed to reflect the applicable discharge and mileage criteria in accordance with any such extension for FY 2018.

4. Proposed Parallel Low-Volume Hospital Payment Adjustment Regarding Hospitals Operated by the Indian Health Services (IHS) or a Tribe

As previously stated, section 1886(d)(12)(C) of the Act and our regulations at 42 CFR 412.101(b)(2) require that, in order to qualify for the low-volume hospital payment adjustment, a hospital must be located more than a specified number of miles from the nearest subsection (d) hospital (referred to as the mileage criterion). Section 1886(d)(1)(B) of the Act defines a “subsection (d) hospital” as a hospital located in one of the 50 States or District of Columbia, other than the specified excluded types of hospitals. As stated in prior rulemaking (for example, 79 FR 50153), CMS considers IHS and Tribal hospitals to be subsection (d) hospitals. However, given the unique nature of IHS and Tribal hospitals and the populations they serve, as discussed below, we believe it would be appropriate to provide additional flexibility in determining eligibility for the low-volume hospital payment adjustment for IHS and non-IHS hospitals and Tribal hospitals that are located less than the specified mileage from one another. Specifically, we are proposing that, for an IHS or Tribal hospital, only its proximity to other IHS or Tribal hospitals would be used to determine if the mileage criterion is met. Similarly, for a non-IHS hospital, only its proximity to other non-IHS hospitals would be used to determine if the mileage criterion is met.

Except for emergencies and a few other limited special cases, those individuals who are not members of a Federally recognized Tribe are not eligible for treatment at IHS or Tribal hospitals. Therefore, such a hospital is not a valid option for the general Medicare population, including local residents who are not members of a Federally recognized Tribe or not otherwise eligible for IHS services. Therefore, we believe it would be appropriate to not consider IHS and Tribal hospitals when evaluating whether a non-IHS hospital meets the mileage criterion.

Likewise, we believe it would be appropriate to not consider non-IHS hospitals when evaluating whether an IHS or Tribal hospital meets the mileage criterion. The principal mission of the IHS is the provision of health care to American Indians and Alaska Natives throughout the United States. In carrying out that mission, IHS operates under two primary authorizing statutes. The first statute, the Snyder Act, authorizes IHS to expend such moneys as Congress may determine from time to time appropriate for the conservation of the health of American Indians or Alaska Natives. We refer readers to 25 U.S.C. 13 (providing that the Bureau of Indian Affairs (BIA) will expend funds as appropriated for, among other things, the conservation of health of American Indians and Alaska Natives); and 42 U.S.C. 2001(a) (transferring the responsibility for American Indian and Alaska Native health care from BIA to HHS). The second statute, the Indian Health Care Improvement Act (HCIA), established IHS as an agency within the Public Health Service of HHS and provides authority for numerous programs to address particular health initiatives for American Indians and Alaska Natives, such as alcohol and substance abuse and diabetes (25 U.S.C. 1601 et seq.).

IHS and Tribal hospitals are charged with addressing the health of American Indians and Alaska Natives and are uniquely situated to provide services to this population. For this reason, we believe it would be appropriate to not consider the non-IHS hospitals when evaluating whether an IHS or Tribal hospital meets the mileage criterion.

Because IHS and Tribal hospitals are subsection (d) hospitals, we are proposing to use our authority under section 1886(d)(5)(I)(i) of the Act to provide an adjustment equal to the applicable low-volume adjustment provided for under section 1886(d)(12) of the Act for an IHS or Tribal hospital whose sole disqualifier for the low-volume hospital adjustment is its proximity to a non-IHS hospital, and for a non-IHS hospital whose sole disqualifier is its proximity to an IHS or Tribal hospital. Such an adjustment would provide that, practically speaking, an IHS or Tribal hospital would be able to receive a low-volume hospital adjustment based on its distance to the nearest IHS or Tribal hospital, and a non-IHS hospital would be able to qualify to receive a low-volume hospital adjustment based on its distance to the nearest non-IHS hospital. We believe it is appropriate to apply this authority here, given the unique characteristics of IHS and Tribal hospitals, as discussed above. To implement this proposed adjustment, we are proposing to revise 42 CFR 412.101 by adding paragraph (e) to provide that, for discharges occurring in FY 2018 and subsequent years, only the distance between IHS or Tribal hospitals would be considered when assessing whether an IHS or Tribal hospital meets the mileage criterion under § 412.101(b)(2). Similarly, only the distance between non-IHS hospitals would be considered when assessing whether a non-IHS hospital meets the mileage criterion under § 412.101(b)(2).

F. Indirect Medical Education (IME) Payment Adjustment Factor for FY 2018 (§ 412.105)

Under the IPPS, an additional payment amount is made to hospitals with residents in an approved graduate medical education (GME) program in order to reflect the higher indirect patient care costs of teaching hospitals.

As previously stated, section 1886(d)(12)(C) of the Act and our regulations at 42 CFR 412.101(b)(2) require that, in order to qualify for the low-volume hospital payment adjustment, a hospital must be located more than a specified number of miles from the nearest subsection (d) hospital (referred to as the mileage criterion).
benefits by the hospital’s total number of patient days furnished to patients entitled to benefits under Medicare Part A. The Medicaid fraction is computed by dividing the hospital’s number of inpatient days furnished to patients who, for such days, were eligible for Medicaid, but were not entitled to benefits under Medicare Part A, by the hospital’s total number of inpatient days in the same period.

Because the DSH payment adjustment is part of the IPPS, the statutory references to “days” in section 1886(d)(5)(F)(i) of the Act have been interpreted to apply only to hospital acute care inpatient days. Regulations located at §412.106 govern the Medicare DSH payment adjustment and specify how the DPP is calculated as well as how beds and patient days are counted in determining the Medicare payment adjustment. Under §412.106(a)(1)(i), the number of beds for the Medicare DSH payment adjustment is determined in accordance with bed counting rules for the IPPS in effect for FY 2018. The remaining amount, equal to an estimate of 75 percent of what otherwise would have been paid as Medicare DSH payments, reduced to reflect changes in the percentage of individuals who are uninsured, is available to make additional payments to each hospital that qualifies for Medicare DSH payments and that has uncompensated care. The payments to each hospital for a fiscal year are based on the hospital’s amount of uncompensated care for a given time period, divided by the total amount of uncompensated care for that same time period reported by all hospitals that receive Medicare DSH payments for the fiscal year.

As provided by section 3133 of the Affordable Care Act, subsection (d) of the Act requires that, for FY 2014 and each subsequent fiscal year, a subsection (d) hospital that would otherwise receive DSH payments made under section 1886(d)(5)(F) of the Act receive two separately calculated payments. Specifically, section 1886(r)(1) of the Act provides that the Secretary shall pay to such subsection (d) hospital (including a Pickle hospital) 25 percent of the amount the hospital would have received under section 1886(d)(5)(F) of the Act for DSH payments, which represents the empirically justified amount for such payment, as determined by the Medicare DSH payment adjustment. In addition to this empirically justified Medicare DSH payment, section 1886(r)(2) of the Act provides that, for FY 2014 and each subsequent fiscal year, the Secretary shall pay to such subsection (d) hospital an additional amount equal to the product of three factors. The first factor is the difference between the aggregate amount of payments that would be made to subsection (d) hospitals under section 1886(d)(5)(F) of the Act if subsection (r) did not apply and the aggregate amount of payments that are made to subsection (d) hospitals under section 1886(r)(1) of the Act for each fiscal year. Therefore, this factor amounts to 75 percent of the payments that would otherwise be made under section 1886(d)(5)(F) of the Act.
most recent estimate of the Congressional Budget Office before the final vote on the Health Care and Education Reconciliation Act of 2010, which is contained in a March 20, 2010 letter from the Director of the Congressional Budget Office to the Speaker of the House. (The March 20, 2010 letter is available for viewing on the following Web site: http://www.cbo.gov/sites/default/files/cbofiles/fjd/docs/113xx/doc11379/amendreconprop.pdf.)

For FY 2018 and subsequent fiscal years, the second factor is 1 minus the percent change in the percent of individuals who are uninsured, as determined by comparing the percent of individuals who were uninsured in 2013 (as estimated by the Secretary, based on data from the Census Bureau or other sources the Secretary determines appropriate, and certified by the Chief Actuary of CMS), and the percent of individuals who were uninsured in the most recent period for which data are available (as so estimated and certified), minus 0.2 percentage point for FYs 2018 and 2019.

The third factor is a percent that, for each subsection (d) hospital, represents the quotient of the amount of uncompensated care for such hospital for a period selected by the Secretary (as estimated by the Secretary, based on appropriate data), including the use of alternative data where the Secretary determines that alternative data are available which are a better proxy for the costs of subsection (d) hospitals for treating the uninsured, and the aggregate amount of uncompensated care for all subsection (d) hospitals that receive a payment under section 1886(r) of the Act. Therefore, this third factor represents a hospital’s uncompensated care amount for a given time period relative to the uncompensated care amount for that same time period for all hospitals that receive Medicare DSH payments in the applicable fiscal year, expressed as a percent.

For each hospital, the product of these three factors represents its additional payment for uncompensated care for the applicable fiscal year. We refer to the additional payment determined by these factors as the ‘‘uncompensated care payment.’’

Section 1886(r) of the Act applies to FY 2014 and each subsequent fiscal year. In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50620 through 50647) and the FY 2014 IPPS interim final rule with comment period (78 FR 61191 through 61197), we set forth our policies for implementing the required changes to the Medicare DSH payment methodology by section 3133 of the Affordable Care Act for FY 2014. In those rules, we noted that, because section 1886(r) of the Act modifies the payment required under section 1886(d)(5)(F) of the Act, it affects only the DSH payment under the operating IPPS. It does not revise or replace the capital IPPS DSH payment provided under the regulations at 42 CFR part 412, subpart M, which were established through the exercise of the Secretary’s discretion in implementing the capital IPPS under section 1886(g)(1)(A) of the Act.

Finally, section 1886(r)(3) of the Act provides that there shall be no administrative or judicial review under section 1869, section 1878, or otherwise of any estimate of the Secretary for purposes of determining the factors described in section 1886(r)(2) of the Act or of any period selected by the Secretary for the purpose of determining those factors. Therefore, there is no administrative or judicial review of the estimates developed for purposes of applying the three factors used to determine uncompensated care payments, or the periods selected in order to develop such estimates.

2. Eligibility for Empirically Justified Medicare DSH Payments and Uncompensated Care Payments

As indicated earlier, the payment methodology under section 3133 of the Affordable Care Act applies to ‘‘subsection (d) hospitals’’ that would otherwise receive a DSH payment made under section 1886(d)(5)(F) of the Act. Therefore, hospitals must receive empirically justified Medicare DSH payments in a fiscal year in order to receive an additional Medicare uncompensated care payment for that year. Specifically, section 1886(r)(2) of the Act states that, in addition to the payment made to a subsection (d) hospital under section 1886(r)(1) of the Act, the Secretary shall pay to such subsection (d) hospitals an additional amount. Because section 1886(r)(1) of the Act refers to empirically justified Medicare DSH payments, the additional payment under section 1886(r)(2) of the Act is limited to hospitals that receive empirically justified Medicare DSH payments in accordance with section 1886(r)(1) of the Act for the applicable fiscal year.

In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50622) and the FY 2014 IPPS interim final rule with comment period (78 FR 61193), we provided that hospitals that are not eligible to receive empirically justified Medicare DSH payments will not receive uncompensated care payments for that year. We also specified that we would make a determination concerning eligibility for interim uncompensated care payments based on each hospital’s estimated DSH status for the applicable fiscal year (using the most recent data that are available). We indicated that our final determination on the hospital’s eligibility for uncompensated care payments will be based on the hospital’s actual DSH status at cost report settlement for that payment year.

In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50622) and the FY 2015 IPPS/LTCH PPS final rule (79 FR 50006), we specified our policies for several specific classes of hospitals within the scope of section 1886(r) of the Act. We refer readers to those two final rules for a detailed discussion of our policies. In summary, we specified the following:

- **Subsection (d) Puerto Rico hospitals** that are eligible for DSH payments also are eligible to receive empirically justified Medicare DSH payments and uncompensated care payments under the new payment methodology (78 FR 50623 and 79 FR 50006).
- **Maryland hospitals** are not eligible to receive empirically justified Medicare DSH payments and uncompensated care payments under the payment methodology of section 1886(r) of the Act because they are not paid under the IPPS. As discussed in the FY 2015 IPPS/LTCH PPS final rule (79 FR 50007), effective January 1, 2014, the State of Maryland elected to no longer have Medicare pay Maryland hospitals in accordance with section 1814(b)(3) of the Act and entered into an agreement with CMS that Maryland hospitals will be paid under the Maryland All-Payer Model. However, under the Maryland All-Payer Model, Maryland hospitals still are not paid under the IPPS. Therefore, they remain ineligible to receive empirically justified Medicare DSH payments or uncompensated care payments under section 1886(r) of the Act.
- **SCHs that are paid under their hospital-specific rate** are not eligible for Medicare DSH payments. SCHs that are paid under the IPPS Federal rate receive interim payments based on what we estimate and project their DSH status to be prior to the beginning of the Federal fiscal year (based on the best available data at that time) subject to settlement through the cost report, and if they receive interim empirically justified Medicare DSH payments in a fiscal year, they also will receive interim uncompensated care payments for that fiscal year on a per discharge basis, subject to settlement through the cost report. Final eligibility determinations will be made at the end.
of the cost reporting period at settlement, and both interim empirically justified Medicare DSH payments and uncompensated care payments will be adjusted accordingly (78 FR 50624 and 79 FR 50007).

- **MDHs** are paid based on the IPPS Federal rate or, if higher, the IPPS Federal rate plus 75 percent of the amount by which the Federal rate is exceeded by the updated hospital-specific rate from certain specified base years (76 FR 51684). The IPPS Federal rate used in the MDH payment methodology is the same IPPS Federal rate that is used in the SCH payment methodology. Section 205 of the Medicare Access and CHIP Reauthorization Act of 2015 (MACRA), Public Law 114–10, enacted April 16, 2015, extended the MDH program for discharges on or after April 1, 2015, through September 30, 2017. Because MDHs are paid based on the IPPS Federal rate, for FY 2017, MDHs continue to be eligible to receive empirically justified Medicare DSH payments and uncompensated care payments if their DPP is at least 15 percent. We apply the same process to determine MDHs’ eligibility for empirically justified Medicare DSH and uncompensated care payments, as we do for all other IPPS hospitals, through September 30, 2017. We note that there has not been legislation at the time of development of this proposed rule that would extend the MDH program beyond September 30, 2017. However, if the MDH program were to be extended beyond its current expiration date, similar to how it was extended under MACRA, MDHs would continue to be paid based on the IPPS Federal rate or, if higher, the IPPS Federal rate plus 75 percent of the amount by which the Federal rate is exceeded by the updated hospital-specific rate from certain specified base years. Accordingly, if the MDH program were to be extended beyond its current expiration date of September 30, 2017, we would continue to make a determination concerning eligibility for interim uncompensated care payments based on each hospital’s estimated DSH status for the applicable fiscal year (using the most recent data that are available). Our final determination on the hospital’s eligibility for uncompensated care payments would be based on the hospital’s actual DSH status at cost report settlement for that payment year. In addition, as we do for all IPPS hospitals, we would calculate the numerator for Factor 3 for all MDHs, regardless of whether they are projected to be eligible for Medicare DSH payments during the fiscal year, but the denominator for Factor 3 would be based on the uncompensated care data from the hospitals that we have projected to be eligible for Medicare DSH payments during the fiscal year. These policies for MDHs would only apply in FY 2018 if the MDH program is extended by statute, beyond its current expiration date of September 30, 2017.

- **IPPS hospitals that have elected to participate in the Bundled Payments for Care Improvement initiative and IPPS hospitals that are participating in the mandatory Comprehensive Care for Joint Replacement Model, the Episode Payment Models, or the Cardiac Rehabilitation Incentive Payment Model** continue to be paid under the IPPS (77 FR 53342) and, therefore, are eligible to receive empirically justified Medicare DSH payments and uncompensated care payments (78 FR 50625 and 79 FR 50008).

- **Hospitals Participating in the Rural Community Hospital Demonstration Program** are not eligible to receive empirically justified Medicare DSH payments and uncompensated care payments under section 1886(r) of the Act because they are not paid under the IPPS (78 FR 50625 and 79 FR 50008). The Rural Community Hospital Demonstration Program was originally authorized for a 5-year period by section 410A of the Medicare Prescription Drug, Improvement, and Modernization Act of 2003 (MMA) (Pub. L. 108–173), and extended for another 5-year period by sections 3123 and 10313 of the Affordable Care Act (Pub. L. 114–255). The period of performance for this 5-year extension period ended December 31, 2016. Section 15003 of the 21st Century Cures Act (Pub. L. 114–255), enacted December 13, 2016, again amended section 410A of Public Law 108–173 to require a 10-year extension period (in place of the 5-year extension required by the Affordable Care Act), to begin on the date immediately following the last day of the initial 5-year period. Section 15003 also requires that, no later than 120 days after enactment of Public Law 114–255, the Secretary issue a solicitation to select additional hospitals to participate in the demonstration program for the second 5 years of the 10-year extension period so long as the maximum number of 30 hospitals stipulated by the Affordable Care Act is not exceeded. (We refer readers to section V.L. of the preamble of this proposed rule for a full discussion of the provisions of section 15003 of Public Law 114–255 and our proposals for implementation.) As of the time of development of this proposed rule, the entire set of hospitals that will participate in the second 5 years of the extension period is unknown. However, we intend to apply a similar payment methodology during the remainder of the extension period. As a result, we expect that hospitals participating in the demonstration will not receive empirically justified DSH payments, and that they will be excluded from receiving interim and final uncompensated care payments for FY 2018 and subsequent fiscal years for the duration of the second 5 years of the extension period.

3. **Empirically Justified Medicare DSH Payments**

As we have discussed earlier, section 1886(r)(1) of the Act requires the Secretary to pay 25 percent of the amount of the Medicare DSH payment that would otherwise be made under section 1886(d)(5)(F) of the Act to a subsection (d) hospital. Because section 1886(r)(1) of the Act merely requires the program to pay a designated percentage of these payments, without revising the criteria governing eligibility for DSH payments or the underlying payment methodology, we stated in the FY 2014 IPPS/LTC PPS final rule that we did not believe that it was necessary to develop any new operational mechanisms for making such payments. Therefore, in the FY 2014 IPPS/LTC PPS final rule (78 FR 50626), we implemented this provision by advising MACs to simply adjust the interim claim payments to the requisite 25 percent of what would have otherwise been paid. We also made corresponding changes to the hospital cost report so that these empirically justified Medicare DSH payments can be settled at the appropriate level at the time of cost report settlement. We provided more detailed operational instructions and cost report instructions following issuance of the FY 2014 IPPS/LTC PPS final rule that are available on the CMS Web site: http://www.cms.gov/Regulations-and-Guidance/Guidance/Transmittals/2014-Transmittals-Items/R5P240.html.

4. **Uncompensated Care Payments**

As we discussed earlier, section 1886(r)(2) of the Act provides that, for each eligible hospital in FY 2014 and subsequent years, the uncompensated care payment is the product of three factors. These three factors represent our estimate of 75 percent of the amount of Medicare DSH payments that would otherwise have been paid, an adjustment to this amount based on the percent change in the national rate of uninsurance compared to the rate of
uninsurance in 2013, and each eligible hospital’s estimated uncompensated care amount relative to the estimated uncompensated care amount for all eligible hospitals. Below we discuss the data sources and methodologies for computing each of these factors, our final policies for FYs 2014 through 2017, and our proposed policies for FY 2018.

a. Proposed Calculation of Factor 1 for FY 2018

Section 1886(r)(2)(A) of the Act establishes Factor 1 in the calculation of the uncompensated care payment. Section 1886(r)(2)(A) of the Act states that this factor is equal to the difference between (1) the aggregate amount of payments that would be made to subsection (d) hospitals under section 1886(d)(5)(F) of the Act if section 1886(r) of the Act did not apply for such fiscal year (as estimated by the Secretary); and (2) the aggregate amount of payments that are made to subsection (d) hospitals under section 1886(r)(1) of the Act for such fiscal year (as so estimated). Therefore, section 1886(r)(2)(A)(i) of the Act represents the estimated Medicare DSH payments that would have been made under section 1886(d)(5)(F) of the Act if section 1886(r) of the Act did not apply for such fiscal year. Under a prospective payment system, we would not know the precise aggregate Medicare DSH payment amount that would be paid for a Federal fiscal year until cost report settlement for all IPPS hospitals is completed, which occurs several years after the end of the Federal fiscal year. Therefore, section 1886(r)(2)(A)(i) of the Act provides authority to estimate this amount, by specifying that, for each fiscal year to which the provision applies, such amount is to be estimated by the Secretary. Similarly, section 1886(r)(2)(A)(ii) of the Act represents the estimated empirically justified Medicare DSH payments to be made in a fiscal year, as prescribed under section 1886(r)(1) of the Act. Again, section 1886(r)(2)(A)(ii) of the Act provides authority to estimate this amount.

Therefore, Factor 1 is the difference between our estimates of: (1) The amount that would have been paid in Medicare DSH payments for the fiscal year, in the absence of the new payment provision; and (2) the amount of empirically justified Medicare DSH payments that are made for the fiscal year, which takes into account the requirement to pay 25 percent of what would have otherwise been paid under section 1886(r) of the Act. In other words, this factor represents our estimate of 75 percent (100 percent minus 25 percent) of our estimate of Medicare DSH payments that would otherwise be made, in the absence of section 1886(r) of the Act, for the fiscal year.

As we did for FY 2017, in this FY 2018 IPPS/LTCH PPS proposed rule, in order to determine Factor 1 in the uncompensated care payment formula for FY 2018, we are proposing to continue the policy established in the FY 2014 IPPS/LTCH PPS final rule (78 FR 50628 through 50630) and in the FY 2014 IPPS interim final rule with comment period (78 FR 61194) of determining Factor 1 by developing estimates of both the aggregate amount of Medicare DSH payments that would be made in the absence of section 1886(r)(1) of the Act and the aggregate amount of empirically justified Medicare DSH payments to hospitals under 1886(r)(1) of the Act. These estimates will not be revised or updated after we know the final Medicare DSH payments for FY 2018.

Therefore, to determine the two elements of proposed Factor 1 for FY 2018 (Medicare DSH payments prior to the application of section 1886(r)(1) of the Act, and empirically justified Medicare DSH payments after application of section 1886(r)(1) of the Act), for this proposed rule, we used the most recently available projections of Medicare DSH payments for the fiscal year, as calculated by CMS’ Office of the Actuary using the most recently filed Medicare hospital cost report with Medicare DSH payment information and the most recent Medicare DSH patient percentages and Medicare DSH payment adjustments provided in the IPPS Impact File.

For purposes of calculating proposed Factor 1 and modeling the impact of this FY 2018 IPPS/LTCH PPS proposed rule, we used the Office of the Actuary’s January 2017 Medicare DSH estimates, which are based on data from the December 2016 update of the Medicare Hospital Cost Report Information System (HCRIS) and the FY 2017 IPPS/LTCH PPS final rule IPPS Impact file, published in conjunction with the publication of the FY 2017 IPPS/LTCH PPS final rule. Because SCHs that are projected to be paid under their hospital-specific rate are excluded from the application of section 1886(r) of the Act, these hospitals also were excluded from the January 2017 Medicare DSH estimates. Furthermore, because section 1886(r) of the Act specifies that the uncompensated care payment is in addition to the empirically justified Medicare DSH payments, 75 percent of DSH payments that would be made without regard to section 1886(r) of the Act, Maryland hospitals participating in the Maryland All-Payer Model that do not receive DSH payments were also excluded from the Office of the Actuary’s January 2017 Medicare DSH estimates. Hospitals that had been participating in the Rural Community Hospital Demonstration Program through December 31, 2016 were included in these estimates. (As discussed earlier, the Affordable Care Act authorized a 5-year extension period for the demonstration, which ended December 31, 2016.) The demonstration was extended for an additional 5 years by section 15003 of Public Law 114–255. Although the hospitals that will participate in the second 5 years of the extension period had not been determined at the time of development of this proposed rule, we intend to apply a similar payment methodology during the second 5 years of the extension period as for the earlier periods of the demonstration. Therefore, hospitals participating in the demonstration would not be eligible to receive DSH payments. If the hospitals participating in the second 5 years of the extension period are known prior to the development of the Medicare DSH estimates for the FY 2018 final rule, these hospitals would be excluded from the Office of the Actuary’s final Medicare DSH estimates for FY 2018.

For this proposed rule, using the data sources discussed above, the Office of the Actuary used the most recently submitted Medicare cost report data to identify Medicare DSH payments and the most recent Medicare DSH payment adjustments and applied inflation updates and assumptions for future changes in utilization and case-mix to estimate Medicare DSH payments for the upcoming fiscal year. The January 2017 Office of the Actuary estimate for Medicare DSH payments for FY 2018, without regard to the application of section 1886(r)(1) of the Act, is approximately $16.003 billion. This estimate excludes Maryland hospitals participating in the Maryland All-Payer Model and SCHs paid under their hospital-specific payment rate. Therefore, based on the January 2017 estimate, the estimate for empirically justified Medicare DSH payments for FY 2018, with the application of section 1886(r)(1) of the Act, is approximately $4.001 billion (or 25 percent of the total amount of estimated Medicare DSH payments for FY 2018). Under §412.106(g)(1)(i) of the regulations, Factor 1 is the difference between these two estimates of the Office of the Actuary. Therefore, in this proposed rule, we are proposing that Factor 1 for
FY 2018 is $12,001,915,095.04, which is equal to 75 percent of the total amount of estimated Medicare DSH payments for FY 2018 ($16,002,553,460.05 minus $4,000,638,365.01).

The Office of the Actuary’s estimates for FY 2018 for this proposed rule began with a baseline of $12,405 billion in Medicare DSH expenditures for FY 2014. The following table shows the factors applied to update this baseline through the current estimate for FY 2018:

**FACTORS APPLIED FOR FY 2015 THROUGH FY 2018 TO ESTIMATE MEDICARE DSH EXPENDITURES USING 2014 BASELINE**

<table>
<thead>
<tr>
<th>FY</th>
<th>Update</th>
<th>Discharge</th>
<th>Case-mix</th>
<th>Other</th>
<th>Total</th>
<th>Estimated DSH payment (in billions)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>2015</td>
<td>1.014</td>
<td>1.0068</td>
<td>1.005</td>
<td>1.0493</td>
<td>1.076581</td>
<td>$13.355</td>
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<td>2016</td>
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<td>0.9757</td>
<td>1.027</td>
<td>1.0689</td>
<td>1.080724</td>
<td>14.433</td>
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<td>2017</td>
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<td>1.005</td>
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<td>1.005</td>
<td>0.9934</td>
<td>1.039603</td>
<td>16.003</td>
</tr>
</tbody>
</table>

*Rounded.

In this table, the discharge column shows the increase in the number of Medicare fee-for-service (FFS) inpatient hospital discharges. The figures for FY 2015 are based on Medicare claims data that have been adjusted by a completion factor. The discharge figure for FY 2016 is based on preliminary data for 2016. The discharge figures for FYs 2017 and 2018 are assumptions based on recent trends recovering back to the long-term trend and assumptions related to how many beneficiaries will be enrolled in Medicare Advantage (MA) plans. The case-mix column shows the increase in case-mix for IPPS hospitals. The case-mix figures for FY 2015 are based on actual data adjusted by a completion factor. The FY 2016 increase is based on preliminary data adjusted by a completion factor. The FYs 2017 and 2018 increases are based on the recommendation of the 2010–2011 Medicare Technical Review Panel. The “Other” column shows the increase in other factors that contribute to the Medicare DSH estimates. These factors include the difference between the total inpatient hospital discharges and the IPPS discharges, and various adjustments to the payment rates that have been included over the years but are not reflected in the other columns (such as the change in rates for the 2-midnight stay policy). In addition, the “Other” column includes a factor for the Medicaid expansion due to the Affordable Care Act. In the past, commenters have contended that the “Other” column understates the effect of the Medicaid expansion. The factor for Medicaid expansion was developed using public information and statements for each State regarding its intent to implement the expansion. Based on this information, it is assumed that 50 percent of all individual who were potentially newly eligible Medicaid enrollees in 2016 resided in States that had elected to expand Medicaid eligibility and, for 2017 and thereafter, that 55 percent of such individuals would reside in expansion States. In the future, these assumptions may change based on actual participation by States. For a discussion of general issues regarding Medicaid projections, we refer readers to the 2016 Actuarial Report on the Financial Outlook for Medicaid (https://www.cms.gov/Research-Statistics-Data-and-Systems/Research/ActuarialStudies/Downloads/MedicaidReport2016.pdf). We note that, in developing their estimates of the effect of Medicaid expansion on Medicare DSH expenditures, our actuaries have assumed that the new Medicaid enrollees are healthier than the average Medicaid recipient and, therefore, use fewer hospital services.

The table below shows the factors that are included in the “Update” column of the above table:

<table>
<thead>
<tr>
<th>FY</th>
<th>Market basket percentage</th>
<th>Affordable Care Act payment reductions</th>
<th>Multifactor productivity adjustment</th>
<th>Documentation and coding</th>
<th>Total update percentage</th>
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<td></td>
<td></td>
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<td>– 0.5</td>
<td>– 0.8</td>
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<tr>
<td>2016</td>
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<td></td>
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<td>– 0.3</td>
<td>– 1.5</td>
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<td></td>
<td></td>
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<td>– 0.4</td>
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</table>

*Note: All numbers are based on FY 2018 President’s Budget projections.

We are inviting public comments on our proposed calculation of Factor 1 for FY 2018.

b. Proposed Calculation of Factor 2 for FY 2018

(1) Background

Section 1886(r)[2][B] of the Act establishes Factor 2 in the calculation of the uncompensated care payment. Specifically, section 1886(r)[2][B][i] of the Act provides that, for each of FYs 2014, 2015, 2016, and 2017, a factor equal to 1 minus the percent change in the percent of individuals under the age of 65 who are uninsured, as determined by comparing the percent of such individuals (1) who were uninsured in 2013, the last year before coverage expansion under the Affordable Care Act (as calculated by the Secretary based on the most recent estimates available from the Director of the Congressional Budget Office before a vote in either House on the Health Care and Education Reconciliation Act of 2010 that, if determined in the affirmative, would clear such Act for enrollment); and (2) who are uninsured in the most recent period for which data are available (as so calculated), minus 0.1 percentage point for FY 2014 and minus 0.2 percentage point for each of FYs 2015, 2016, and 2017.

Section 1886(r)[2][B][ii] of the Act further indicates that the percent of
individuals under 65 without insurance in 2013 must be the percent of such individuals who were uninsured in 2013, the last year before coverage expansion under the Affordable Care Act (as calculated by the Secretary based on the most recent estimates available from the Director of the Congressional Budget Office before a vote in either House on the Health Care and Education Reconciliation Act of 2010). The Health Care and Education Reconciliation Act (Pub. L. 111–152) was enacted on March 30, 2010. It was passed in the House of Representatives on March 21, 2010, and by the Senate on March 25, 2010. Because the House of Representatives was the first House to vote on the Health Care and Education Reconciliation Act of 2010 on March 21, 2010, we have determined that the most recent estimate available from the Director of the Congressional Budget Office “before a vote in either House on the Health Care and Education Reconciliation Act of 2010.” (emphasis added) appeared in a March 20, 2010 letter from the director of the CBO to the Speaker of the House. Therefore, we believe that only the estimates in this March 20, 2010 letter meet the statutory requirement under section 1886(r)(2)(B)(i) of the Act. (To view the March 20, 2010 letter, we refer readers to the Web site at: https://www.cbo.gov/sites/default/files/111th-congress-2009-2010/costestimate/amendreconprop.pdf.)

In its March 20, 2010 letter to the Speaker of the House of Representatives, the CBO provided two estimates of the “post-policy uninsured population.” The first estimate is of the “Insured Share of the Nonelderly Population Including All Residents” (82 percent) and the second estimate is of the “Insured Share of the Nonelderly Population Excluding Unauthorized Immigrants” (83 percent). In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50634), we used the first estimate that includes all residents, including unauthorized immigrants. We stated that we believe this estimate is most consistent with the statute, which requires us to measure “the percent of individuals under the age of 65 who are uninsured” and provides no exclusions except for individuals over the age of 65. In addition, we stated that we believe this estimate more fully reflects the levels of uninsurance in the United States than the estimate for uncompensated care for hospitals than the estimate that reflects only legal residents. The March 20, 2010 CBO letter reports these figures as the estimated percentage of individuals with insurance. However, because section 1886(r)(2)(B)(i) of the Act requires that we compare the percent of individuals who are uninsured in the most recent period for which data are available with the percent of individuals who were uninsured in 2013, in the FY 2014 IPPS/LTCH PPS final rule, we used the CBO insurance rate figure and subtracted that amount from 100 percent (that is, the total population without regard to insurance status) to estimate the 2013 baseline percent of individuals without insurance. Therefore, for FYs 2014 through 2017, our estimate of the uninsurance percentage for 2013 was 18 percent. Section 1886(r)(2)(B)(i) of the Act requires that we compare the baseline uninsurance rate to the percent of such individuals who are uninsured in the most recent period for which data are available (as so calculated). In the FY 2014, FY 2015, FY 2016, and FY 2017 IPPS/LTCH PPS final rules (78 FR 50634, 79 FR 50014, 80 FR 49522, and 81 FR 56952, respectively), we used the same data source, CBO estimates, to calculate this percent of individuals without insurance. In response to public comments, we also agreed that we should normalize the CBO estimates, which are based on the calendar year, for the Federal fiscal years for which each calculation of Factor 2 is made (78 FR 50633). Therefore, for the FY 2017 IPPS/LTCH PPS final rule (81 FR 56952), we used the most recently available estimate of the uninsurance rate, which was based on the CBO’s March 2016 estimates of the effects of the Affordable Care Act on health insurance coverage (which are available at https://www.cbo.gov/sites/default/files/114th-congress-2015-2016/reports/53185-HealthInsuranceBaseline.pdf). The CBO’s March 2016 estimate of individuals under the age of 65 with insurance in CY 2016 was 90 percent. Therefore, the CBO’s most recent estimate of the rate of uninsurance in CY 2016 was 10 percent (that is, 100 percent minus 90 percent). The CBO’s March 2016 estimate of individuals under the age of 65 with insurance in CY 2017 was also 90 percent. Therefore, the CBO’s most recent estimate of the rate of uninsurance in CY 2017 available for the FY 2017 final rule was also 10 percent (that is, 100 percent minus 90 percent). The calculation of the final Factor 2 for FY 2017, employing a weighted average of the CBO projections for CY 2016 and CY 2017, was as follows:

1
1–((0.10–0.18)/0.18)
1–0.0444 = 0.5555
1
0.5555
= 0.5536

Therefore, the final Factor 2 for FY 2017 was 55.36 percent.

The FY 2017 final uncompensated care amount was: $10,797,476,782.62 × 0.5536 = $5,977,483,146.86.

FY 2017 uncompensated care total available ...... $5,977,483,146.86

(2) Proposed Methodology for Calculation of Factor 2 for FY 2018

Section 1886(r)(2)(B)(ii) of the Act permits the use of a data source other than the CBO estimates to determine the percent change in the rate of uninsurance beginning in FY 2018. In addition, for FY 2018 and subsequent years, the statute does not require that the estimate of the percent of individuals who are uninsured be limited to individuals who are under 65. Specifically, the statute states that, for FY 2018 and subsequent fiscal years, the second factor is 1 minus the percent change in the percent of individuals who are uninsured, as determined by comparing the percent of individuals who were uninsured in 2013 (as estimated by the Secretary, based on data from the Census Bureau or other sources the Secretary determines appropriate, and certified by the Chief Actuary of CMS) and the percent of individuals who were uninsured in the most recent period for which data are available (as so estimated and certified), minus 0.2 percentage point for FYs 2018 and 2019. In the FY 2017 IPPS/LTCH PPS final rule (81 FR 56952), we indicated that we planned to address changes to the methodology for determining Factor 2 and the viability of potential alternative data sources in the FY 2018 IPPS/LTCH PPS proposed rule. In our analysis of a potential data source for the rate of uninsurance for purposes of computing Factor 2 in FY 2018, we considered the following: (a)
The extent to which the source accounted for the full U.S. population; (b) the extent to which the source comprehensively accounted for both public and private health insurance coverage in deriving its estimates of the number of uninsured; (c) the extent to which the source utilized data from the Census Bureau; (d) the timeliness of the estimates; (e) the continuity of the estimates over time; (f) the accuracy of the estimates; and (g) the availability of projections (including the availability of projections using an established estimation methodology that would allow for calculation of the rate of uninsurance for the applicable Federal fiscal year). These considerations are consistent with the statutory requirement that this estimate be based on data from the Census Bureau or other sources the Secretary determines appropriate and help to ensure the data source will provide reasonable estimates for the rate of uninsurance that are available in conjunction with the IPPS rulemaking cycle.

We have determined that the source that, on balance, best meets all of these considerations is the uninsured estimates produced by CMS’ Office of the Actuary (OACT) as part of the development of the National Health Expenditure Accounts (NHEA). The NHEA represents the government’s official estimates of economic activity (spending) within the health sector. The information contained in the NHEA has been used to study numerous topics related to the health care sector, including, but not limited to, changes in the amount and cost of health services purchased and the payers or programs that provide or purchase these services; the economic causal factors at work in the health sector; the impact of policy changes, including major health reform; and comparisons to other countries’ health spending. Of relevance to the determination of Factor 2 is that the comprehensive and integrated structure of the NHEA creates an ideal tool for evaluating changes to the health care system, such as the mix of the insured and uninsured because this mix is integral to the well-established NHEA methodology. Below we describe some aspects of the methodology used to develop the NHEA that we believe are particularly relevant in estimating the percent change in the rate of uninsurance for FY 2018. A full description of the methodology used to develop the NHEA is available on the CMS Web site at: https://www.cms.gov/Research-Statistics-Data-and-Systems/Statistics-Trends-and-Reports/NationalHealthExpendData/Downloads/DSM-15.pdf.

The NHEA estimates of U.S. population reflect the Census Bureau’s definition of the resident-based population, which includes all people who usually reside in the 50 States or the District of Columbia, but excludes residents living in Puerto Rico and areas under U.S. sovereignty, members of the U.S. Armed Forces overseas, and U.S. citizens whose usual place of residence is outside of the United States, plus a small (typically less than 0.2 percent of population) adjustment to reflect Census undercounts. In past years, the estimates for Factor 2 were made using the CBO’s uninsured population estimates for the under 65 population. For FY 2018 and subsequent years, the statute does not restrict the estimate to the measurement of the percent of individuals under the age of 65 who are uninsured. Accordingly, we believe it is appropriate to use an estimate that reflects the rate of uninsurance in the United States across all age groups. In addition, we continue to believe that a resident-based population estimate more fully reflects the levels of uninsurance in the United States that influence uncompensated care for hospitals than an estimate that reflects only legal residents. The NHEA estimates of uninsurance are for the total U.S. population (all ages) and not by specific age cohort, such as the population under the age of 65.

The NHEA includes comprehensive enrollment estimates for total private health insurance (PHI) (including direct and employer-sponsored plans), Medicare, Medicaid, the Children’s Health Insurance Program (CHIP), and other public programs, and estimates of the number of individuals who are uninsured. Estimates of total PHI enrollment are available for 1960 through 2015, estimates of Medicaid, Medicare, and CHIP enrollment are available for the length of the respective programs, and all other estimates (including the more detailed estimates of direct-purchased and employer-sponsored insurance) are available for 1987 through 2015. The NHEA data are publicly available on the CMS Web site at: https://www.cms.gov/Research-Statistics-Data-and-Systems/Statistics-Trends-and-Reports/NationalHealthExpendData/index.html. In order to compute Factor 2, the first metric that is needed is the proportion of the total U.S. population that was uninsured in 2013. In developing the estimates for the NHEA, OACT’s methodology includes estimating the number of uninsured individuals for 1987 through 2009 based on the enhanced Current Population Survey (CPS) from the State Health Access Data Assistance Center (SHADAC). The CPS, sponsored jointly by the U.S. Census Bureau and the U.S. Bureau of Labor Statistics (BLS), is the primary source of labor force statistics for the population of the United States. (We refer readers to the Web site at: http://www.census.gov/programs-surveys/cps.html.) The enhanced CPS, available from SHADAC (available at http://datacenter.shadac.org) accounts for changes in the CPS methodology over time. OACT further adjusts the enhanced CPS for an estimated undercount of Medicaid enrollees (a population that is often not fully captured in surveys that include Medicaid enrollees due to a perceived stigma associated with being enrolled in the Medicaid program or confusion about the source of their health insurance).

To estimate the number of uninsured individuals for 2010 through 2014, OACT extrapolates from the 2009 CPS data using data from the National Health Interview Survey (NHIS). The NHIS is one of the major data collection programs of the National Center for Health Statistics (NCHS), which is part of the Centers for Disease Control and Prevention (CDC). The U.S. Census Bureau is the data collection agent for the NHIS. The NHIS results have been instrumental over the years in providing data to track health status, health care access, and progress toward achieving national health objectives. For further information regarding the NHIS, we refer readers to the CDC Web site at: https://www.cdc.gov/nchs/nhis/index.htm. For 2015, the estimate of the rate of uninsurance in the NHEA matches with the estimate from the NHIS. The next metrics needed to compute Factor 2 are projections of the rate of uninsurance in both calendar years 2017 and 2018. On an annual basis, the OACT projects enrollment and spending trends for the coming 10-year period. Those projections (currently for years 2016 through 2025) use the latest NHEA historical data, which presently run through 2015. The NHEA projection methodology accounts for expected changes in enrollment across all of the categories of insurance coverage previously listed. The sources for projected growth rates in enrollment for Medicare, Medicaid, and CHIP include the latest Medicare Trustees Report, the Medicaid Actuarial Report, or other updated estimates as produced by the OACT. Projected rates of growth in enrollment for private health insurance and the uninsured are based largely on

The use of data from the NHEA to estimate the rate of uninsurance is consistent with the statute and meets the criteria we have identified for determining the appropriate data source. Section 1886(f)(2)(B)(ii) of the Act instructs the Secretary to estimate the rate of uninsurance for purposes of Factor 2 based on data from the Census Bureau or other sources the Secretary determines appropriate. The NHEA utilizes data from the Census Bureau; the estimates are available in time for the IPPS rulemaking cycle; the estimates are produced by OACT on an annual basis and are expected to continue to be produced for the foreseeable future; and projections are available for calendar year time periods that span the upcoming fiscal year. Timeliness and continuity are important considerations because of our need to be able to update this estimate annually. Accuracy is also a very important consideration and, all things being equal, we would choose the most accurate data source that sufficiently meets our other criteria.

As with the CBO estimates on which we based Factor 2 in prior fiscal years, the NHEA estimates are for a calendar year. In the rulemaking for FY 2014, many commenters noted that the uncompensated care payments are made on fiscal year and not a calendar year basis and requested that CMS normalize the CBO estimate to reflect a fiscal year basis. Specifically, commenters requested that CMS calculate a weighted average of the CBO estimate for October through December 2013 and the CBO estimate for January through September 2014 when determining Factor 2 for FY 2014. We agreed with the commenters that normalizing the estimate to cover FY 2014 would more accurately reflect the rate of uninsurance that hospitals would experience during the FY 2014 payment year. Accordingly, we estimated the rate of uninsurance for FY 2014 by calculating a weighted average of the CBO estimates for CY 2013 and CY 2014 (78 FR 50633). We have continued this weighted average approach in each fiscal year since FY 2014.

We continue to believe that, in order to estimate the rate of uninsurance during a fiscal year more accurately, Factor 2 should reflect the estimated rate of uninsurance that hospitals will experience during the fiscal year, rather than the rate of uninsurance during only one of the calendar years that the fiscal year spans. However, we have concerns about the future potential for the uninsured rate to vary nonuniformly in the 2 calendar years that the fiscal year spans (for example, due to changes in the economy or changes in legislation). Nevertheless, for FY 2018, because OACT’s current estimates of the percent of individuals without insurance in CY 2017 and CY 2018 are relatively close, we do not believe this is a significant policy issue and are proposing to continue with the weighted average approach used in past fiscal years in order to estimate the rate of uninsurance for FY 2018.

The calculation of the proposed Factor 2 for FY 2018 using a weighted average of OACT’s projections for CY 2017 and CY 2018 is as follows:

1. Percent of individuals without insurance for CY 2013: 14 percent.
2. Percent of individuals without insurance for CY 2017: 8.3 percent.
3. Percent of individuals without insurance for CY 2018: 8.1 percent.

Factor 2 for FY 2018 is 58.01 percent.

\[ 0.5801 = \text{Factor 2} \]

Therefore, the proposed Factor 2 for FY 2018 is 58.01 percent.

The proposed FY 2018 uncompensated care amount for all hospitals estimated to receive Medicare DSH payments is:

\[ \text{Proposed FY 2018 uncompensated care amount} = 0.5801 \times 6,962,310,946.63 = \$6,962,310,946.63 \]

We are inviting public comments on our proposed methodology for calculation of Factor 2 for FY 2018.

(1) Background

Section 1886(r)(2)(C) of the Act defines Factor 3 in the calculation of the uncompensated care payment. As we have discussed earlier, section 1886(r)(2)(C) of the Act states that Factor 3 is equal to the percent, for each subsection (d) hospital, that represents the quotient of (1) the amount of uncompensated care for such hospital for a period selected by the Secretary (as estimated by the Secretary, based on appropriate data (including, in the case where the Secretary determines alternative data are available that are a better proxy for the costs of subsection (d) hospitals for treating the uninsured, the use of such alternative data)); and (2) the aggregate amount of uncompensated care for all subsection (d) hospitals that receive a payment under section 1886(r) of the Act for such period (as so estimated, based on such data).

Therefore, Factor 3 is a hospital-specific value that expresses the proportion of the estimated uncompensated care amount for each subsection (d) hospital and each subsection (d) Puerto Rico hospital with the potential to receive Medicare DSH payments relative to the estimated uncompensated care amount for all hospitals estimated to receive Medicare DSH payments in the fiscal year for which the uncompensated care payment is to be made. Factor 3 is applied to the product of Factor 1 and Factor 2 to determine the amount of the uncompensated care payment that each eligible hospital will receive for FY 2014 and subsequent fiscal years. In order to implement the statutory requirements for this factor of the uncompensated care payment formula, it was necessary to determine: (1) The definition of uncompensated care or, in other words, the specific items that are to be included in the numerator (that is, the estimated uncompensated care amount for an individual hospital) and the denominator (that is, the estimated uncompensated care amount for all hospitals estimated to receive Medicare DSH payments in the applicable fiscal year); (2) the data source(s) for the estimated uncompensated care amount; and (3) the timing and manner of computing the quotient for each hospital estimated to receive Medicare DSH payments. The statute instructs the Secretary to estimate the amounts of uncompensated care for a period based on appropriate data. In addition, we note that the statute permits the Secretary to use alternative data in the

| Proposed FY 2018 uncompensated care total available | \$6,962,310,946.63 |

We are inviting public comments on our proposed methodology for calculation of Factor 2 for FY 2018.
case where the Secretary determines that such alternative data are available that are a better proxy for the costs of subsection (d) hospitals for treating individuals who are uninsured.

In the course of considering how to determine Factor 3 during the rulemaking process for FY 2014, we considered defining the amount of uncompensated care for a hospital as the uncompensated care costs of each hospital and determined that Worksheet S–10 of the Medicare cost report potentially provides the most complete data regarding uncompensated care costs for Medicare hospitals. However, because of concerns regarding variations in the data reported on Worksheet S–10 and the completeness of these data, we did not propose to use data from Worksheet S–10 to determine Factor 3 for FY 2014, the first year this provision was in effect, or for FY 2015, 2016, or 2017. When we first discussed using Worksheet S–10 to allocate hospitals’ shares of uncompensated care costs in the FY 2014 IPPS/LTCH PPS final rule (78 FR 50638), we explained why we believed that it was premature to use uncompensated care costs reported on Worksheet S–10 for FY 2014. Specifically, at that time, the most recent available cost reports would have been from FY’s 2010 and 2011, which were submitted on or after May 1, 2010, when the new Worksheet S–10 went into effect. We believed that concerns about the standardization and completeness of the Worksheet S–10 data could be more acute for data collected in the first year of the Worksheet’s use (78 FR 50635). In addition, we believed that it would be most appropriate to use data elements that have been historically publicly available, subject to audit, and used for payment purposes (or that the public understands will be used for payment purposes) to determine the amount of uncompensated care for purposes of Factor 3 (78 FR 50635). At the time we issued the FY 2014 IPPS/LTCH PPS final rule, we did not believe that the available data regarding uncompensated care from Worksheet S–10 met these criteria and, therefore, we believed they were not reliable enough to use for determining FY 2014 uncompensated care payments. Accordingly, for FY 2014, we concluded that utilization of insured low-income patients would be a better proxy for the costs of hospitals in treating the uninsured. For FY’s 2015, 2016, and 2017, the cost reports used for calculating uncompensated care payments (Worksheet S–10, 2011, 2012, and 2013) were also submitted prior to the time that hospitals were on notice that Worksheet S–10 could be the data source for calculating uncompensated care payments. Therefore, we believed it was also appropriate to use proxy data to calculate Factor 3 for these years.

We stated in the preamble of the FY 2017 IPPS/LTCH PPS proposed rule that we believed that, for FY 2018, many of the above concerns would no longer be relevant. That is, hospitals were on notice as of FY 2014 that Worksheet S–10 could eventually become the data source for CMS to calculate uncompensated care payments. Furthermore, hospitals’ cost reports from FY 2014 had been publicly available for some time, and CMS had analyses of Worksheet S–10 conducted both internally and by stakeholders demonstrating that Worksheet S–10 accuracy had improved over time. Specifically, as discussed in the FY 2017 IPPS/LTCH PPS proposed rule (81 FR 25090), MedPAC has provided analyses that found that current Worksheet S–10 data are a better proxy for predicting audited uncompensated care costs than Medicaid/Medicare SSI days, and that the data on Worksheet S–10 would improve over time as the data are actually used to make payments. CMS has also undertaken an extensive analysis of the Worksheet S–10 data, benchmarking it against the data on uncompensated care costs reported to the Internal Revenue Service (IRS) on Form 990 by not-for-profit hospitals. (This analysis, performed by Dobson DaVanzo & Associates, LLC, under contract to CMS, was included in a report entitled “Improvements to Medicare Disproportionate Share Hospital (DSH) Payments Report: Benchmarking S–10 Data Using IRS Form 990 Data and Worksheet S–10 Trend Analyses,” which is available on the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/dsh.html under the Downloads section.) The analysis determined a strong and converging correlation between the amounts for Factor 3 derived using the IRS Form 990 and Worksheet S–10 data suggesting that Worksheet S–10 uncompensated care data are becoming more stable over time. Given these results and in light of the fact that hospitals have been on notice since the FY 2014 rulemaking that CMS intended eventually to use Worksheet S–10 as the data source for calculating uncompensated care payments, we believed it would be appropriate to propose to begin incorporating the Worksheet S–10 data for purposes of calculating Factor 3 starting in FY 2018. In section IV.F.4.d. of the preamble of the FY 2017 IPPS/LTCH PPS proposed rule (81 FR 25090 through 25094), we proposed a methodology and timeline for incorporating Worksheet S–10 data in the calculation of Factor 3 beginning in FY 2018 and invited public comments on that proposal.

While some commenters, including MedPAC, were supportive of the proposal, many other commenters expressed concerns about a perceived lack of clarity in the Worksheet S–10 instructions and their belief in the necessity of a strict audit mechanism to capture aberrant uncompensated care costs reported on Worksheet S–10. Many commenters also cited the report from Dobson DaVanzo, which concluded that hospitals are doing a better job of reporting their uncompensated care data on Worksheet S–10 than they did a few years ago.

However, these commenters disagreed with CMS about the significance of this observation. One commenter stated that even if it is true in the aggregate that hospitals are reporting data more accurately on Worksheet S–10, the zero-sum nature of the calculation of uncompensated care payments is such that the remaining inaccuracy and lack of uniformity in the data reported can have a very large impact on hospitals. The commenter asserted that if hospitals, for whatever reason, over-report their uncompensated care, they benefit financially from doing so, while those that do not aggressively report suffer financial harm. The commenter concluded that, for this reason, the possibility that some hospitals are generally “doing better” with reporting data is not good enough. All hospitals must do better, and until they do, the commenter believed that data from Worksheet S–10 are not accurate enough for public policymaking purposes. Other commenters asserted that the Dobson/ DaVanzo study did not illustrate or even evaluate whether data from Worksheet S–10 are a reasonable proxy for the costs hospitals incur in providing care to the uninsured. These commenters pointed to their own analyses, which indicated that the most notable aberrations in Worksheet S–10 data reporting occur among public hospitals, which do not file a Form 990 and are therefore missing from the Dobson/DaVanzo analysis.

On balance, after considering all of the comments, we elected not to finalize our proposal to begin to incorporate Worksheet S–10 into the calculation of Factor 3 for FY 2018 in the FY 2017 IPPS/LTCH PPS rule. We stated that we were postponing the decision regarding when to begin incorporating
We found further evidence for this tipping point when we examined changes to the FY 2014 Worksheet S–10 data submitted by hospitals since the publication of the FY 2017 IPPS/LTCH PPS final rule. In the FY 2017 IPPS/LTCH PPS final rule, as part of our ongoing quality control and data improvement measures for the Worksheet S–10, we referred readers to Change Request 9648, Transmittal 1681, titled "The Supplemental Security Income (SSI)/Medicare Beneficiary Data for Fiscal Year 2014 for Inpatient Prospective Payment System (IPPS) Hospitals, Inpatient Rehabilitation Facilities (IRFs), and Long Term Care Hospitals (LTCH),” issued on July 15, 2016 (available at: https://www.cms.gov/ Regulations-and-Guidance/Guidance/ Transmittals/Downloads/ RI6810TN.pdf). In this transmittal, as part of the process for ensuring complete submission of Worksheet S–10 by all eligible DSH hospitals, we instructed MACs to accept amended Worksheets S–10 for FY 2014 cost reports submitted by hospitals (or initial submissions of Worksheet S–10 if none had been submitted previously) and to upload them to the Health Care Provider Cost Report Information System (HCRIS) in a timely manner. The transmittal stated that, for revisions to be considered, hospitals were required to submit their amended FY 2014 cost report containing the revised Worksheet S–10 (or a completed Worksheet S–10 if no data were included on the previously submitted cost report) to the MAC no later than September 30, 2016.

We have examined hospitals’ FY 2014 cost reports to see if the Worksheet S–10 data on those cost reports have changed as a result of the opportunity for hospitals to submit revised Worksheet S–10 data for FY 2014. Specifically, we compared hospitals’ FY 2014 Worksheet S–10 data as they existed in the first quarter of CY 2016 with data from the fourth quarter of CY 2016. We found that the FY 2014 Worksheet S–10 data had changed over that time period for approximately one quarter of hospitals that receive uncompensated care payments. The fact that the Worksheet S–10 data changed for such a significant number of hospitals following a review of the cost report data they originally submitted and that the revised Worksheet S–10 information is available to be used in determining uncompensated care costs contributes to our belief that we can no longer conclude that alternative data are available that are a better proxy than the Worksheet S–10 data for the costs of...
subsection (d) hospitals for treating individuals who are uninsured.

Commenters have also provided equity arguments with respect to the relationship between uncompensated care payments and the expansion of Medicaid in certain States under the authority provided by the Affordable Care Act. The commenters have made a twofold argument. First, they have argued that hospitals in States that did not expand Medicaid treat a higher number of uninsured patients compared to hospitals in States that did expand Medicaid and, as a result, provide more uncompensated care. However, since the implementation of the new DSH payment methodology under section 3133 of the Affordable Care Act in FY 2014, these hospitals have experienced reductions in the payments for uncompensated care due to the national decline in the uninsured rate driven in large part by Medicaid expansions in other States. Second, they have argued that hospitals in non-expansion States will be penalized a second time when Medicaid utilization is used as part of the basis for determining Factor 3 because their Medicaid utilization has not grown as much relative to hospitals in expansion States. We note that, while CMS has not yet used data affected by Medicaid expansion when determining Factor 3, commenters are concerned that they will be penalized in future calculations when these data are used.

In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50639), we recognized that, in using Medicaid days as part of the proxy for uncompensated care, it would be possible for hospitals in States that choose to expand Medicaid to receive higher uncompensated care payments because they may have more Medicaid patient days than hospitals in a State that does not choose to expand Medicaid. Because the earliest Medicaid expansions under the Affordable Care Act began in 2014, the 2011, 2012, and 2013 Medicaid days data used to determine Factor 3 for FY 2017 are the most recent available data on Medicaid utilization that do not reflect the effects of these Medicaid expansions. Accordingly, if we were to use only low-income insured days to estimate uncompensated care in FY 2018, we would need to hold the time period of these data constant and use data on Medicaid days from 2011, 2012, and 2013 in order to avoid the risk of any redistributive effects arising from the decision to expand Medicaid in certain States. As a result, we would be using older data that may provide a less accurate proxy for the level of uncompensated care being furnished by hospitals in FY 2018, contributing to our growing concerns regarding the continued use of low-income insured days as a proxy for uncompensated care costs in FY 2018.

We also note that when weighing the new information that has become available to us regarding the Worksheet S–10 and the low-income days proxy since the FY 2018 rulemaking, we are not considering these developments in isolation, but rather in the context of the information that we previously considered as part our discussions of the Worksheet S–10 data in prior rulemaking. Part of this background is provided by the 2007 MedPAC analysis of data from the Government Accountability Office (GAO) and the American Hospital Association (AHA), which suggests that Medicaid days and low income Medicare days are not a good proxy for uncompensated care costs (80 FR 49525). Additional analyses performed by MedPAC showed that the correlation between audited uncompensated care data from 2009 and the data from the FY 2011 Worksheet S–10 was over 0.50, as compared to a correlation of approximately 0.50 between the audited uncompensated care data and 2011 Medicare SSI and Medicaid days. Based on this analysis, MedPAC concluded that use of Worksheet S–10 data was already better than using Medicare SSI and Medicaid days as a proxy for uncompensated care costs, and that the data on Worksheet S–10 would improve over time as the data are actually used to make payments (81 FR 25090). Furthermore, MedPAC in the past has commented about the low-income days proxy we have used historically because it is an inpatient measure and much of the uncompensated care provided by certain hospitals, including rural hospitals, occurs in the emergency room or other outpatient areas. In its comments on the FY 2017 proposed rule, MedPAC again recommended we start using the Worksheet S–10 data with a phase-in (81 FR 56962). In summary, when weighing the new information that has become available to us since the FY 2017 rulemaking in conjunction with the information regarding Worksheet S–10 data and the low-income days proxy that we have analyzed as part of our consideration of this issue in prior rulemaking, we can no longer conclude that alternative data to the Worksheet S–10 are available for FY 2014 that are a better proxy for the costs of subsection (d) hospitals for treating individuals who are uninsured. We discuss below our proposed methodology and how we would begin to incorporate Worksheet S–10 data for FY 2014 into the calculation of Factor 3 of the uncompensated care payment methodology.

(3) Proposed Time Period for Calculating Factor 3 for FY 2018, Including Methodology for Incorporating Worksheet S–10 Data

Section 1886(r)(2)(C) of the Act not only governs the selection of the data to be used in calculating Factor 3, but also allows the Secretary the discretion to determine the time periods from which we will derive the data to estimate the numerator and denominator of the Factor 3 quotient. Specifically, section 1886(r)(2)(C)(i) of the Act defines the numerator of the quotient as the amount of uncompensated care for such hospital for a period selected by the Secretary. Section 1886(r)(2)(C)(ii) of the Act defines the denominator as the aggregate amount of uncompensated care for all subsection (d) hospitals that receive a payment under section 1886(r) of the Act for such period. In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50639), we adopted a process of making interim payments with final cost report settlement for both the empirically justified Medicare DSH payments and the uncompensated care payments required by section 3133 of the Affordable Care Act. Consistent with that process, we also determined the time period from which to calculate the numerator and denominator of the Factor 3 quotient in a way that would be consistent with making interim and final payments. Specifically, we must have Factor 3 values available for hospitals that we estimate will qualify for Medicare DSH payments and for those hospitals that we do not estimate will qualify for Medicare DSH payments but that may ultimately qualify for Medicare DSH payments at the time of cost report settlement.

In the FY 2017 IPPS/LTCH PPS final rule, in order to mitigate undue fluctuations in the amount of uncompensated care payments to hospitals from year to year and smooth over anomalies between cost reporting periods, we finalized a policy of calculating a hospital’s share of uncompensated care based on data derived from three cost reporting periods instead of one cost reporting period. As explained in the preamble to the FY 2017 IPPS/LTCH PPS final rule (81 FR 56957 through 56959), instead of determining Factor 3 using Medicaid days from a single cost reporting period and the most recent available data on Medicare SSI utilization, as we did in FY 2014, FY 2015, and FY 2016, we used selected Medicaid days from three cost reporting periods (FYs 2011, 2012, and 2013) and SSI days from the three most...
recent available years of SSI utilization data (FYs 2012, 2013, and 2014) to compute Factor 3 for FY 2017. We continued to extract Medicaid days data from the most recent update of HCRIS, which for FY 2017 was the March 2016 update. Furthermore, instead of determining a single Factor 3 as we have done since the first year of the uncompensated care payment in FY 2014, we calculated an individual Factor 3 for each of the three cost reporting periods, which we then averaged by the number of cost reporting years with data to compute the final Factor 3 for a hospital. Under this policy, if a hospital had merged, we would combine data from both hospitals for the cost reporting periods in which the merger was not reflected in the surviving hospital’s cost report data to compute Factor 3 for the surviving hospital. Moreover, to further reduce undue fluctuations in a hospital’s uncompensated care payments, if a hospital filed multiple cost reports beginning in the same fiscal year, we combined data from the multiple cost reports so that a hospital could have a Factor 3 calculated using more than one cost report within a cost reporting period. We codified these changes for FY 2017 by amending the regulations at § 412.106(g)(1)(ii)(C).

For FY 2018, we are proposing to continue to use the methodology finalized in FY 2017 and to compute Factor 3 using an average of data from three cost reporting periods instead of one cost reporting period. Consistent with the methodology used to calculate Factor 3 for FY 2017, we are proposing to advance the time period of the data used in the calculation of Factor 3 forward by one year and using data from FY 2012, FY 2013, and FY 2014 cost reports. For the reasons we described earlier, we believe it would not be appropriate to use Worksheet S–10 data for periods prior to FY 2014. Rather, for cost reporting periods prior to FY 2014, we believe it would be appropriate to continue to use low-income insured days. Accordingly, with a time period that includes three cost reporting years consisting of FY 2014, FY 2013, and FY 2012, we are proposing to use Worksheet S–10 data for the FY 2014 cost reporting period and the low-income insured day proxy data for the two earlier cost reporting periods. In order to perform this calculation, we will draw three sets of data (2 years of Medicaid utilization data and 1 year of Worksheet S–10 data) from the most recent available HCRIS extract, which for FY 2018 is the December 2016 update of HCRIS for the proposed rule and the March 2017 update of HCRIS for the final rule. Accordingly, for FY 2018, in addition to the Worksheet S–10 data for FY 2014, we are proposing to use Medicaid days from FY 2012 and FY 2013 cost reports and FY 2014 and FY 2015 SSI ratios. We also would continue to use FY 2012 cost report data submitted to CMS by IHS and Tribal hospitals to determine Medicaid days for those hospitals. (We note that cost report data from IHS and Tribal hospitals are included in HCRIS beginning in FY 2013 and are no longer submitted separately.) We also are proposing to continue the policies that were finalized in the FY 2015 IPPS/LTCH PPS final rule (79 FR 50020) to address several specific issues concerning the process and data to be employed in determining Factor 3 in the case of hospital mergers as well as the policies finalized in the FY 2017 IPPS/LTCH PPS final rule concerning multiple cost reports beginning in the same fiscal year (81 FR 56957).

We believe this approach, if we were to propose to use FY 2019 and FY 2020, would have the effect of transitioning the incorporation of data from Worksheet S–10 into the calculation of Factor 3. Starting with one year of Worksheet S–10 data in FY 2018, an additional year of Worksheet S–10 data would be incorporated into the calculation of Factor 3 in FY 2019, and the use of low-income insured days would be phased out by FY 2020. In addition, we acknowledge the concerns regarding IHS/Tribal hospitals and subsection (d) Puerto Rico hospitals that some commenters expressed in response to the FY 2017 proposal to begin using Worksheet S–10 data to determine Factor 3 in FY 2018. According to some of these commenters, the use of data from Worksheet S–10 to calculate uncompensated care may jeopardize all of the IHS/Tribal hospitals’ uncompensated care payments due to their the unique funding structure. With respect to Puerto Rico, other commenters asserted that the use of Worksheet S–10 data may not be appropriate given the historical treatment of subsection (d) Puerto Rico hospitals under the statutory provisions governing payments under Medicaid and Medicare Part A and its impact on the reporting of uncompensated care payments by these hospitals. After consideration of the concerns, we believe that the uncompensated care data reported by Puerto Rico and IHS/Tribal hospitals needs to be further examined and should not be used for FY 2018. For the reasons described earlier related to the impact of the Medicaid expansion beginning in FY 2014, we also do not believe it would be appropriate to calculate a Factor 3 for these hospitals using FY 2014 low-income insured days. Because we do not believe it is appropriate to use the FY 2014 uncompensated care data for these hospitals and we also do not believe it is appropriate to use the FY 2014 low-income insured days, we believe that the best proxy for the costs of Puerto Rico and IHS/Tribal hospitals for treating the uninsured is the low-income-insured days data for FY 2012 and FY 2013. Accordingly, we are proposing for these hospitals that when we compute the individual Factor 3s for each of the three cost reporting periods that are used to determine Factor 3, rather than computing a Factor 3 using Worksheet S–10 data from the hospital’s FY 2014 cost report, we would substitute the Factor 3 calculated using the hospital’s FY 2013 low-income insured days. That is, in order to determine the Factor 3 for FY 2018, we would calculate an average of three individual Factor 3s using the Factor 3 calculated using FY 2013 cost report data twice and the Factor 3 calculated using FY 2012 cost report data once. We believe it is appropriate to double-weight the Factor 3 calculated using FY 2013 data as it reflects the most recent available information regarding the hospital’s low-income insured days before any expansion of Medicaid. We note that as we are not making any proposals with respect to the calculation of Factor 3 for FY 2019 at this time, we will reexamine the use of the Worksheet S–10 data for Puerto Rico and IHS/Tribal hospitals as part of the FY 2019 rulemaking. In addition, we are proposing to continue to use a proxy for SSI days consisting of 14 percent of a hospital’s Medicaid days for Puerto Rico hospitals, as finalized in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56953 through 56956).

Therefore, for FY 2018, we are proposing to compute Factor 3 for each hospital by—

- **Step 1:** Calculating Factor 3 using the low-income insured days proxy based on FY 2012 cost report data and the FY 2014 SSI ratio;
- **Step 2:** Calculating Factor 3 using the insured low-income days proxy based on FY 2013 cost report data and the FY 2015 SSI ratio;
- **Step 3:** Calculating Factor 3 based on the FY 2014 Worksheet S–10 data (or using the Factor 3 calculated in Step 2 for Puerto Rico and IHS/Tribal hospitals); and
- **Step 4:** Averaging the Factor 3 values from Steps 1, 2, and 3; that is, adding the Factor 3 values from FY 2012, FY 2013, and FY 2014 for each
hospital, and dividing that amount by the number of cost reporting periods with data to compute an average Factor 3.

We are inviting public comments on our proposed methodology for calculating Factor 3 for FY 2018.

We note that if this proposed methodology is adopted for FY 2018, we would expect to propose to use a similar methodology for calculating Factor 3 for subsequent years, meaning that for FY 2019 we would expect to incorporate data from the FY 2015 Worksheet S–10 into the methodology and drop the FY 2012 low-income insured day proxy data. However, we are not making any proposals with respect to the calculation of Factor 3 for FY 2019 at this time.

For new hospitals that do not have data for any of the three cost reporting periods used in the proposed Factor 3 calculation, we will continue to apply the new hospital policy finalized in the FY 2014 IPPS/LTCH PPS final rule (78 FR 50643). That is, the hospital will not receive either interim empirically justified Medicare DSH payments or interim uncompensated care payments. However, if the hospital is later determined to be eligible to receive empirically justified Medicare DSH payments based on its FY 2018 cost report, the hospital will also receive an uncompensated care payment calculated using a Factor 3, where the numerator is the uncompensated care costs reported on Worksheet S–10 of the hospital’s FY 2018 cost report, and the denominator is the sum of uncompensated care costs reported on Worksheet S–10 of all DSH eligible hospitals’ FY 2014 cost reports as prospectively determined during rulemaking. We note that, given the selected time period of the data used to calculate Factor 3, any hospitals with a CCN established after October 1, 2014 would be considered new and subject to this policy.

As we have done for every proposed and final rule beginning in FY 2014, in conjunction with both the FY 2018 IPPS/LTCH PPS proposed rule and final rule, we will publish on the CMS Web site a table listing Factor 3 for all hospitals that we estimate would receive empirically justified Medicare DSH payments in FY 2018 (that is, those hospitals that would receive interim uncompensated care payments during the fiscal year), and for the remaining subsection (d) hospitals and subsection (d) Puerto Rico hospitals that have the potential of receiving a Medicare DSH payment in the event that they receive an empirically justified Medicare DSH payment for the fiscal year as determined at cost report settlement. We note that, as of this proposed rule, the FY 2015 SSI ratios are not yet available. Accordingly, for modeling purposes, we computed Factor 3 using the most recent available data regarding SSI days from the FY 2013 and FY 2014 SSI ratios. However, we expect that the FY 2015 SSI ratios will be available to calculate Factor 3 for the FY 2018 IPPS/LTCH PPS final rule.

We also will publish a supplemental data file containing a list of the mergers that we are aware of and the computed uncompensated care payment for each merged hospital. Hospitals have 60 days from the date of public display of this FY 2018 IPPS/LTCH PPS proposed rule to review the table and supplemental data file published on the CMS Web site in conjunction with the proposed rule and to notify CMS in writing of any inaccuracies. Comments can be submitted to the CMS inbox at Section3133DSH@cms.hhs.gov. We will address these comments as appropriate in the table and the supplemental data file that we publish on the CMS Web site in conjunction with the publication of the FY 2018 IPPS/LTCH PPS final rule. After the publication of the FY 2018 IPPS/LTCH PPS final rule, hospitals will have until August 31, 2017, to review and submit comments on the accuracy of the table and supplemental data file published in conjunction with the final rule. Comments may be submitted to the CMS inbox at Section3133DSH@cms.hhs.gov through August 31, 2017, and any changes to Factor 3 will be posted on the CMS Web site prior to October 1, 2017.

(4) Methodological Considerations for Calculating Factor 3

• **Annualizing short cost reports.** As we explained in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56957 through 56959), we believe that for hospitals that file multiple cost reports beginning in the same year, combining the data from these cost reports has the benefit of supplementing the data of hospitals that filed cost reports that are less than 12 months, such that the basis of their uncompensated care payments and those of hospitals that filed full-year 12-month cost reports would be more equitable. In response to our original proposal in the FY 2017 IPPS/LTCH PPS proposed rule to combine data from multiple cost reports, many hospitals stated that while they were appreciative of CMS’s efforts to provide a more equitable playing field for hospitals that filed short cost reports, they believed that expanding the time period of the data used to calculate Factor 3 as well as combining data across multiple cost reports would not remedy the fact that some hospitals are still disadvantaged by having less than 36 months of data in their Factor 3 calculation (81 FR 56959). Other commenters opposed the use of multiple cost reporting periods if it would result in a hospital having more than 12 months of data in the Factor 3 calculation for a year, and recommended that CMS prorate the data to a 12-month period. Similarly, other commenters recommended that CMS annualize cost report data for any cost reporting period that is less than 12 months. In the FY 2017 IPPS/LTCH PPS final rule, we acknowledged that, although we had not made any proposal in the FY 2017 IPPS/LTCH PPS proposed rule to annualize the cost reports used to calculate Factor 3, the situations presented by commenters, including both long and short cost reporting periods, pose unique challenges in the context of estimating Factor 3. We stated that we intended to consider the issue further and might address the issue in future rulemaking.

For this FY 2018 IPPS/LTCH PPS proposed rule, taking into consideration the feedback from hospitals that have been disadvantaged in the Factor 3 calculation due to cost reports that do not span a full year, we are proposing to annualize Medicaid days if a hospital’s cost report does not equal 12 months of data. We are not proposing to annualize SSI days because we do not obtain these data from hospital cost reports in HCRIS. Rather, we obtain these data from the latest available SSI ratios posted on the Medicare DSH homepage (https://www.cms.gov/medicare/medicare-fee-for-service-payment/acuteinpatientpps/dsh.html), which are aggregated at the hospital level and do not have the information needed to determine if the data should be annualized.

Under this proposal, if the time between the start date of a hospital’s cost reporting year and the end date of its cost reporting year is less than 12 months, we would annualize the Medicaid days so that the hospital has 12 months of data included in its Factor 3 calculation. Conversely, if the time between the aforementioned start date and the end date is greater than 12 months, we would annualize the Medicaid days to achieve 12 months of Medicaid days data. If a hospital files more than one cost report beginning in the same fiscal year, we would first combine the data across the multiple cost reports before determining the difference between the start date and the end date to see if annualization is needed.
To annualize the Medicaid days for a long or short cost reporting year, we would divide the length of a full year (365 or 366 calendar days, as applicable) by the length of the cost reporting year (the number of calendar days in the cost reporting year) and then multiply the quotient by the number of Medicaid days in the cost reporting year.

For instance, a cost reporting year that is 365 calendar days long with 1,200 Medicaid days would be annualized as follows: (365/365) * 1,200 = 1,373 days. A cost reporting year that is 385 calendar days long with 1,200 Medicaid days would be annualized using the same formula: (365/385) * 1,200 = 1,137 days.

Likewise, because long and short cost reporting periods pose the same challenges in the context of estimating Factor 3 using hospital uncompensated care costs, we are proposing to annualize the uncompensated care cost data reported on Worksheet 5–10 for cost reports that do not equal 12 months of data, by dividing the length of a full year (365 or 366 calendar days, as applicable) by the length of the cost reporting year (number of calendar days in the cost reporting year) and then multiplying the quotient by the total reported uncompensated care costs for the cost reporting year.

For instance, a cost reporting year that is 285 calendar days long reporting $10,500,000 in uncompensated care costs would be annualized as follows:

\[(\text{365/285} \times 10,500,000) = 13,447,368.\]

A cost reporting year that is 385 calendar days long reporting $10,500,000 in uncompensated care costs would be annualized using the same formula:

\[(\text{365/385} \times 10,500,000) = 9,954,545.\]

If a hospital files more than one cost report beginning in the same fiscal year, we would first combine the data across the multiple cost reports before determining the length of the cost reporting year to see if annualization is needed.

We are inviting public comment on our proposal to annualize the cost reports used to calculate Factor 3 for FY 2018. In addition, as noted earlier, our proposal to continue calculating a hospital’s share of uncompensated care payments using a time period that includes three cost reporting years is also designed to mitigate undue fluctuations in the amount of uncompensated care payments to hospitals from year to year and smooth over anomalies between cost reporting periods. Given that our proposal to annualize the costs reports used to calculate the Factor 3 for FY 2018 would also mitigate fluctuations in the amount of uncompensated care payments from year to year, we also are seeking public comment on the degree to which the use of three cost reporting years would still be necessary if we were to adopt our proposal to annualize the cost reports used to calculate Factor 3, or if instead the use of a single cost reporting year or two cost reporting years would be appropriate. In order to facilitate public comments, we intend to post on our Web site a data file containing information similar to the information provided in section I.H.5., “Effects of the Proposed Changes to Medicare DSH and Uncompensated Care Payments for FY 2018” of Appendix A of this proposed rule. However, instead of reflecting our proposed approach of calculating Factor 3 using a time period that includes three cost reporting years, it would reflect an alternative approach of calculating Factor 3 using only the most recent year (FY 2014) of our proposed 3-year average. In all other respects, the calculation of Factor 3 would remain the same.

- **Scaling Factor.** Under the methodology adopted in the FY 2017 IPPS/LTCH PPS final rule and that we are proposing to apply in FY 2018, if the hospital does not have data for one or more of the three cost reporting periods, we will compute Factor 3 for the periods available and average those. In other words, we will divide the sum of the individual Factor 3s by the number of cost reporting periods with data so as not to disadvantage hospitals that are missing data for one or more cost reporting periods. Following the publication of the FY 2017 IPPS/LTCH PPS final rule, several hospitals noted that this aspect of the methodology resulted in the Factor 3 values of DSH eligible hospitals in Table 18 and the Medicare DSH Supplemental Data File adding up to slightly greater than one, which results in total uncompensated care payments somewhat exceeding the estimate published in the FY 2017 final rule. Specifically, for hospitals that have fewer than 3 cost reporting years with data, dividing the individual Factor 3s by the number of cost reporting years with data (that is, 2 cost reporting years or 1 cost reporting year) results in a higher average Factor 3 than if the individual Factor 3s were divided by the number of cost reporting years, regardless of whether or not there is data (that is, 3 cost reporting years). For example, a hospital with no data for FY 2011 and a Factor 3 of 0.000051702 for FY 2012 and 0.000049852 for FY 2013 would have an average Factor 3 of 0.000050807 if averaged by 2 but an average Factor 3 of only 0.000033871 if averaged by 3. After reviewing the data in Table 18 and the Medicare DSH Supplemental Data File, which were published in conjunction with the FY 2017 IPPS/LTCH PPS final rule, we concluded that the hospitals’ observations are correct and that an adjustment is needed so that total uncompensated care payments do not exceed the estimate published in section V.G.4.b.(2) of the preamble of this proposed rule.

Accordingly, to address the effects of averaging Factor 3s calculated for three separate fiscal years, we are proposing to apply a scaling factor to the Factor 3 values of all DSH eligible hospitals so that total uncompensated care payments are consistent with the estimated amount available to make uncompensated care payments for FY 2018. Under this proposal, we would first compute the Factor 3 and uncompensated care payments for all hospitals that we anticipate qualifying for Medicare DSH payments in FY 2018. We would then divide 1 (the expected sum of all eligible hospitals’ Factor 3) by the actual sum of all eligible hospitals’ Factor 3 values and multiply the quotient by each hospital’s total uncompensated care payment to obtain scaled uncompensated care payment amounts whose sum is consistent with the estimate of the total amount available to make uncompensated care payments in section V.G.4.b.(2) of the preamble of this proposed rule. The hospital-specific uncompensated care amount would then be divided by a 3-year claims average to obtain the amount of the interim uncompensated care payment the hospital will receive for each claim. As an illustration of the calculation of the scaling factor, applying this proposal to the FY 2017 uncompensated care payments would have resulted in a scaling factor of 0.9992 (1/1.0008). We note that the FY 2017 uncompensated care payments as calculated for the FY 2017 IPPS final rule exceeded the estimated amount by approximately 5% due to the lack of a scaling factor.

We are inviting public comments on our proposal to apply a scaling factor to all hospitals’ Factor 3 values for FY 2018.

(5) Methodological Considerations for Incorporating Worksheet S–10 Data

- **Definition of uncompensated care.** In the FY 2014 IPPS/LTCH PPS rulemaking, we considered three potential definitions of uncompensated care: Charity care; charity care + bad debt; and charity care + bad debt +...
Medicaid shortfalls. As we explained in the FY 2014 IPPS/LTCH PPS final rule (78 FR 50634), we considered proposing to define the amount of uncompensated care for a hospital as the uncompensated care costs of that hospital and considered potential data sources for those costs. We examined the literature on uncompensated care and the concepts of uncompensated care used in various public and private programs, and considered input from stakeholders and public comments in various forums, including the national provider call that we held in January 2013. Our review of the information from these sources indicated that there is some variation in how different States, provider organizations, and Federal programs define “uncompensated care.” However, a common theme of almost all these definitions is that they include both “charity care” and “bad debt” as components of “uncompensated care.” Therefore, a definition that incorporates the most commonly used factors within uncompensated care as reported by stakeholders would include charity care costs and bad debt costs. Worksheet S–10 employs the definition of charity care plus non-Medicare bad debt. Specifically:

- **Cost of Charity Care (Line 23)**
- **Cost of non-Medicare bad debt expense (Line 29)**

Cost of non-Medicare uncompensated care (Line 30)

Where:

- **Cost of charity care = Cost of initial obligation of patients approved for charity care (Line 21) minus partial payment by patients approved for charity care (Line 22).**
- **Cost of non-Medicare bad debt expense = Cost to charge ratio (line 1) times non-Medicare and non-reimbursable bad debt expense (Line 28).**

In the FY 2017 IPPS/LTCH PPS proposed rule (81 FR 25093; 81 FR 56971), we proposed to adopt a definition of uncompensated care costs that included charity care and non-Medicare bad debt. We explained that we believe there are compelling arguments for excluding Medicaid shortfalls from the definition of uncompensated care, including the fact that several government agencies and key stakeholders do not consider Medicaid shortfalls in their definition of uncompensated care and that excluding Medicaid shortfalls from the uncompensated care definition allows Medicare uncompensated care payments to target hospitals that have a disproportionate share of uncompensated care for patients with no insurance coverage. Although we did not finalize the proposed definition of uncompensated care costs as part of the FY 2017 rulemaking, we continue to believe a definition that incorporates the most commonly used factors within uncompensated care as reported by stakeholders would include charity care costs and non-Medicare bad debt costs, which correlates to Line 30 of Worksheet S–10. Therefore, we are again proposing that, for purposes of calculating Factor 3 and uncompensated care costs beginning in FY 2018, “uncompensated care” would be defined as the amount on line 30 of Worksheet S–10, which is the cost of charity care (Line 23) and the cost of non-Medicare bad debt (Line 29). We are inviting public comments on this proposal.

**Trims to apply to CCRs on Line 1 of Worksheet S–10.** As we noted in the FY 2017 IPPS/LTCH proposed and final rules (81 FR 25093; 81 FR 56971), commenters have suggested that uncompensated care costs reported on Worksheet S–10 should be audited due to extremely high values consistently reported by some hospitals. In response to these comments, we have reviewed the Worksheet S–10 data and identified approximately 10 to 20 hospitals that have anomalous uncompensated care costs. We note that many of these hospitals are public hospitals, which can have charging practices that are distinct from other hospital types. We believe that, just as we apply trims to hospitals’ CCRs to eliminate anomalies when calculating outlier payments for extraordinarily high cost cases (§ 412.84(h)(3)(ii)), it is appropriate to apply statistical trims to the CCRs on Worksheet S–10, Line 1 that are considered anomalies. Specifically, § 412.84(h)(3)(ii) states that the Medicare contractor may use a statewide CCR for hospitals whose operating or capital CCR is in excess of 3 standard deviations above the national geometric mean CCR. Based on the information currently available to us, we will remove hospitals that have CCRs above the calculated ceiling of 0.937.

**Step 1:** Remove Maryland hospitals. In addition, we will remove all-inclusive rate providers, as they have charge structures that differ from other IPPS hospitals, and providers that did not report a CCR on Worksheet S–10, Line 1, and assign them the statewide average CCR in step 5 below.

**Step 2:** For hospitals with multiple cost reports included in the 2014 HCRIS data, (a) combine the amounts from Worksheet C, Part I, Line 202, Column 3 from each cost report to calculate total costs, (b) combine the amounts from Worksheet C, Part I, Line 202, Column 8 from each cost report to calculate total charges, and (c) divide the total costs by the total charges to arrive at a recalculated CCR.

**Step 3:** Calculate a CCR “ceiling” using the CCRs reported on Worksheet S–10, Line 1, from all IPPS hospitals that were not removed in Step 1 (including non-DSH eligible hospitals), or the recalculated CCR described in Step 2. The ceiling is calculated as 3 standard deviations above the national geometric mean CCR. This approach is consistent with our calculation of the CCR ceiling used for high-cost outliers. Remove all hospitals that exceed the ceiling so that these aberrant CCRs do not skew the calculation of the statewide average CCR. Based on the information currently available to us, this trim would remove 9 hospitals that have CCRs above the calculated ceiling of 0.937.

**Step 4:** Using the CCRs for the remaining hospitals in Step 3, determine the urban and rural statewide average CCRs using Line 1 of Worksheet S–10 for hospitals within each State (including non-DSH eligible hospitals), weighted by the sum of total inpatient discharges and outpatient visits from Worksheet S–3, Part I, Line 14, Column 14.

**Step 5:** Assign the appropriate statewide average CCR (urban or rural) calculated in Step 4 to all hospitals with a CCR greater than 3 standard deviations above the corresponding national geometric mean (that is, the CCR “ceiling”), as well as to all-inclusive rate providers, and providers that did not report a CCR on Worksheet S–10, Line 1. The statewide average CCR would therefore be applied to 140 hospitals, of which 14 did not report a CCR on Worksheet S–10, Line 1, 9 had a CCR that exceeded the calculated ceiling of 0.937, and 117 are all-inclusive rate providers.

After applying the applicable trims to a hospital’s CCR as appropriate, we are proposing to calculate a hospital's uncompensated care costs as being
equal to Line 30, which is the sum of Line 23 and Line 29, as follows:

Hospital Uncompensated Care Costs =
Line 30 (Line 23 + Line 29), which is equal to—

[(Line 1 CCR (as adjusted, if applicable) × charity care line 20) – (Payments received for charity care Line 22)] + [(Line 1 CCR (as adjusted, if applicable) × Non-Medicare and non-reimbursable Bad Debt Line 28)].

We are inviting public comments on our proposed trim methodology for FY 2018.

**Cost report revisions and Worksheet S–10 audits.** While not directly relevant to our proposal to use FY 2014 Worksheet S–10 data beginning in FY 2018, we note that as part of our ongoing quality control and data improvement measures to continue to improve the Worksheet S–10 data over time, we have made revisions to the cost report instructions and developed an audit process.

With respect to the cost reporting instructions, on November 18, 2016, we issued Transmittal 10 which updated the instructions for Form 2552–10. Specifically, we updated the instructions in Section 4012 of Chapter 40 of the Provider Reimbursement Manual, Part II. The instructions clarify the reporting of charges for charity care. Transmittal 10 can be downloaded from the CMS Web site at: https://www.cms.gov/Regulations-and-Guidance/Guidance/Transmittals/2016-Transmittals-Items/R10P240.html.

With respect to the audit process, in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56694), we stated that we intended to provide standardized instructions to the MACs to guide them in determining when and how often a hospital’s Worksheet S–10 should be reviewed. We indicated that we would not make the MACs’ review protocol public as all CMS desk review and audit protocols are confidential and are for CMS and MAC use only. The instructions for the MACs are still under development and will be provided to the MACs as soon as possible. We refer readers to the FY 2017 IPPS/LTCH PPS final rule for a complete discussion concerning the issues that we are considering in developing the instructions that will be provided to the MACs. We expect that cost reports beginning in FY 2017 will be the first cost reports for which the Worksheet S–10 data will be subject to a desk review. We do not anticipate making any further modifications to the Worksheet S–10 instructions at this time so that hospitals can begin to review and conform to the current instructions in Transmittal 10. Predictability is an important part of the process for reporting data on Worksheet S–10. As a result, we believe it is reasonable to wait until the Worksheet S–10 data have been submitted, the audits have been performed, and the data are available for review before we consider making any further revisions to the Worksheet S–10 instructions.

**H. Medicare-Dependent, Small Rural Hospital (MDH) Program (§ 412.108)**

1. Background for the MDH Program

Section 1886(d)(5)(G) of the Act provides special payment protections, under the IPPS, to a Medicare-dependent, small rural hospital (MDH). (For additional information on the MDH program and the current methodology, we refer readers to the FY 2012 IPPS/LTCH PPS final rule (76 FR 51683 through 51684).) As discussed in section V.B.1. of the preamble of this proposed rule, the MDH program provisions at section 1886(d)(5)(G) of the Act will expire at the end of FY 2017. Beginning with discharges occurring on or after October 1, 2017, all hospitals that previously qualified for MDH status will be paid based on the Federal rate.

Since the extension of the MDH program through FY 2012 provided by section 3124 of the Affordable Care Act, the MDH program had been extended by subsequent legislation as follows:

Section 606 of the ATRA (Pub. L. 112–240) extended the MDH program through FY 2013 (that is, for discharges occurring before October 1, 2013).

Section 1106 of the Pathway for SGR Reform Act of 2013 (Pub. L. 113–67) extended the MDH program through the first half of FY 2014 (that is, for discharges occurring before April 1, 2014).

Section 106 of the PAMA (Pub. L. 113–93) extended the MDH program through the first half of FY 2015 (that is, for discharges occurring before April 1, 2015).

Section 205 of the MACRA (Pub. L. 114–10) extended the MDH program through FY 2017 (that is, for discharges occurring before October 1, 2017). For additional information on the extensions of the MDH program after FY 2017, see the preamble of this proposed rule with comment period (80 FR 49506); and the FY 2017 IPPS/LTCH PPS final rule (81 FR 57054 through 57057).

**b. Expiration of the MDH Program**

Because section 205 of the MACRA extended the MDH program through FY 2017 only, beginning October 1, 2017, the MDH program will no longer be in effect. Because the MDH program is not authorized by statute beyond September 30, 2017, beginning October 1, 2017, all hospitals that previously qualified for MDH status under section 1886(d)(5)(G) of the Act will no longer have MDH status and will be paid based on the IPPS Federal rate.

When the MDH program was set to expire at the end of FY 2012, in the FY 2013 IPPS/LTCH PPS final rule (77 FR 53404 through 53405), we revised our sole community hospital (SCH) policies to allow MDHs to apply for SCH status in advance of the expiration of the MDH program and be paid as such under certain conditions. We codified these changes in the regulations at §412.92(b)(2)(ii) and (b)(2)(v). Specifically, the existing regulations at §412.92(b)(2)(ii) and (b)(2)(v) allow for an effective date of an approval of SCH status that is the day following the expiration date of the MDH program.

We note that these same conditions apply to MDHs that intend to apply for SCH status with the expiration of the MDH program on September 30, 2017. Therefore, in order for an MDH to receive SCH status effective October 1, 2017, the MDH must apply for SCH status at least 30 days before the expiration of the MDH program; that is, the MDH must apply for SCH status by September 1, 2017. The MDH also must request that, if approved as an SCH, the SCH status expires with the expiration of the MDH program; that is, the MDH must request that the SCH status, if approved, be effective October 1, 2017, immediately after its MDH status expires with the expiration of the MDH program on September 30, 2017. We emphasize that an MDH that applies for SCH status in anticipation of the expiration of the MDH program would not qualify for the October 1, 2017 effective date for SCH status if it does not apply by the September 1, 2017 deadline. If the MDH does not apply by the September 1, 2017 deadline, the hospital would instead be subject to the usual effective date for SCH classification; that is, 30 days after the date of CMS’ written notification of approval as specified at §412.92(b)(2)(i).

We note that the regulations governing the MDH program are found...
at §412.108 and the MDH program is also cited in the general payment rules in the regulations at §412.90. As stated earlier, under current law, the MDH program will expire at the end of FY 2017, which is already reflected in §412.108. As such, we are not proposing specific amendments to the regulations at §412.108 to reflect the expiration of the MDH program. However, it has come to our attention that, with the various extensions of the MDH program as noted earlier, we neglected to make conforming changes to the regulation text at §412.90. Therefore, we are proposing to revise the general payment rules under §412.90 to reflect the expiration of the MDH program. However, we are proposing that if the MDH program were to be extended by law, similar to how it was extended through FY 2013, by the ATRA (Pub. L. 112–240); through March 31, 2014, by the Pathway for SGR Reform Act of 2013 (Pub. L. 113–167); through March 31, 2015, by the PAMA (Pub. L. 113–93); and most recently through FY 2017, by the MACRA (Pub. L. 114–10), we would make conforming changes to the regulations governing the MDH program at §412.108(a)(1) and (c)(2)(iii) and the general payment rules at §412.90 to reflect such an extension of the MDH program. These conforming changes would only be made if the MDH program were to be extended by statute beyond September 30, 2017.

1. Hospital Readmissions Reduction Program: Proposed Updates and Changes (§§412.150 Through 412.154)

1. Statutory Basis for the Hospital Readmissions Reduction Program

Section 3025 of the Affordable Care Act, as amended by section 10309 of the Affordable Care Act, added section 1886(q) to the Act, which establishes the "Hospital Readmissions Reduction Program" effective for discharges from "applicable hospitals" beginning on or after October 1, 2012. Under the Hospital Readmissions Reduction Program, payments to applicable hospitals may be reduced to account for certain excess readmissions. We refer readers to section IV.E.1. of the FY 2016 IPPS/LTC PPS final rule (80 FR 49530 through 49531) for a detailed discussion and additional information on the statutory history of the Hospital Readmissions Reduction Program.

On December 13, 2016, the 21st Century Cures Act (Pub. L. 114–255) was enacted. Section 15002 of that law added (D) and (E) to section 1886(q) of the Act, which directs the Secretary to assign hospitals to peer groups, develop a methodology that allows for separate comparisons for hospitals within these groups, and allows for changes in the risk adjustment methodology. The 21st Century Cures Act also directs the Medicare Payment Advisory Commission (MedPAC) to conduct a review of overall hospital readmissions and whether such readmissions are related to any changes in outpatient and emergency services furnished. A report on the study is required to be submitted in the MedPAC's report to Congress no later than June 2018.

Specifically, section 1886(q)(3)(D) of the Act directs the Secretary to develop a transitional methodology that accounts for the percentage of full-benefit dual eligible patients treated by a hospital to determine a hospital's payment adjustment factor. Section 1886(q)(3)(D)(i) of the Act sets forth the requirement that the Secretary assign hospitals to groups and apply a methodology "that allows for separate comparison of hospitals within each such group." This applies to discharges that occur during and after FY 2019 and before the application of section 1886(q)(3)(E)(i) of the Act, which allows the Secretary to take into account the recommendations in the reports required by the IMPACT Act (Pub. L. 113–185) related to risk adjustment and social risk factors. The first of two reports required in the IMPACT Act was released in December of 2016 (available at: https://aspe.hhs.gov/system/files/pdf/253971/ASPEESRTPfinal.pdf), and the second report is required to be completed by October 2019.

The hospital groups in section 1886(q)(3)(D)(ii) of the Act are described as "being based on their overall proportion, of the inpatients who are entitled to, or enrolled for, benefits under Medicare Part A and who are full-benefit dual eligible individuals (as defined in section 1935(c)(6) of the Act)." The Secretary is further required to consult with MedPAC when defining groups and may consider analysis done by MedPAC in preparation for its June 2013 report submitted to Congress. Section 1886(q)(3)(D)(iii) of the Act prevents the imposition of additional reporting requirements in order to carry out subparagraph (D). Section 1886(q)(3)(D)(iv) of the Act requires that the estimated total amount of reductions in payments using the methodology should equal the estimated total amount of reductions in payments if subparagraph (D) did not apply. Section 1886(q)(3)(E) of the Act outlines the considerations the Secretary may take into account with respect to the risk adjustment methodology. Section 1886(q)(3)(E)(i) of the Act allows the Secretary to take into account studies conducted and recommendations made by the Secretary under section 2(d)(1) of the IMPACT Act in the application of risk adjustment methodologies. This does not preclude the consideration of the use of groupings of hospitals. The Secretary is also allowed under section 1886(q)(3)(E)(ii) of the Act to consider the use of "V" or other ICD-related codes for removal of a readmission with respect to discharges occurring after FY 2018. Section 1886(q)(3)(E)(iii) of the Act outlines the considerations the Secretary may make in the removal of certain readmissions. For discharges occurring after FY 2018, the Secretary may consider the removal as a readmission of an admission that is classified within one or more of the following: Transplants; end-stage renal disease; burns; trauma; psychosis; or substance abuse.

2. Regulatory Background

We refer readers to the following past final rules for detailed discussions of the regulatory background and descriptions of the current policies for the Hospital Readmissions Reduction Program: The FY 2012 IPPS/LTC PPS final rule (76 FR 51660 through 51676); the FY 2013 IPPS/LTC PPS final rule (77 FR 53374 through 53401); the FY 2014 IPPS/LTC PPS final rule (78 FR 50649 through 50676); the FY 2015 IPPS/LTC PPS final rule (79 FR 50024 through 50048); the FY 2016 IPPS/LTC PPS final rule (80 FR 49530 through 49543); and the FY 2017 IPPS/LTC PPS final rule (81 FR 56973 through 56979). These policies describe the general framework for the implementation of the Hospital Readmissions Reduction Program, including: (1) The selection of and measures for the applicable conditions; (2) the calculation of the excess readmission ratio, which is used, in part, to calculate the readmissions adjustment factor; (3) the current calculation of the hospital readmission payment adjustment factor, specifically addressing the base operating DRG payment amount, aggregate payments for excess readmissions, and aggregate payments for all discharges; (4) the opportunity for hospitals to review and submit corrections using a process similar to what is currently used for posting results on Hospital Compare; (5) the adoption of an extraordinary circumstances exception policy to address hospitals that experience a disaster or other extraordinary circumstance; (6) the clarification that the public reporting of excess
readmission ratios will be posted on an annual basis to the Hospital Compare Web site as soon as is feasible following the preview period; and (7) the specification that the definition of “applicable hospital” does not include hospitals and hospital units excluded from the IPPS, such as LTCHs, cancer hospitals, children’s hospitals, IRFs, IPFs, CAHs, and hospitals in Puerto Rico.

We also have codified certain requirements of the Hospital Readmissions Reduction Program at 42 CFR 412.152 through 412.154.

3. Maintenance of Technical Specifications for Quality Measures

We refer readers to the FY 2015 IPPS/LTCH PPS final rule (79 FR 50039) for a discussion of the maintenance of technical specifications for quality measures for the Hospital Readmissions Reduction Program. Technical specifications of the readmission measures are provided on our Web site in the Measure Methodology Reports at: http://www.cms.gov/Medicare/Quality- Initiatives-Patient-Assessment- Instruments/HospitalQualityInitiatives/ Measure-Methodology.html. Additional resources about the Hospital Readmissions Reduction Program and measure technical specifications are on the QualityNet Web site on the Resources page at: http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1220772412995

4. Proposed Policies for the Hospital Readmissions Reduction Program

In this proposed rule, we are proposing the following policies for the Hospital Readmissions Reduction Program: (1) The applicable time period for FY 2018; (2) the calculation of aggregate payments for excess readmissions for FY 2018; (3) changes to the payment adjustment factor in accordance with section 15002 of the 21st Century Cures Act for FY 2019; and (4) updates to the Extraordinary Circumstances-Excursion policy beginning in FY 2018 as related to extraordinary circumstances that occur on or after October 1, 2017. These proposals are described in more detail below.

5. Proposed Applicable Period for FY 2018

Under section 1886(q)(5)(D) of the Act, the Secretary has the authority to specify the applicable period with respect to a fiscal year under the Hospital Readmissions Reduction Program. In the FY 2012 IPPS/LTCH PPS final rule (76 FR 51671), we finalized our policy to use 3 years of claims data to calculate the readmission measures. In the FY 2013 IPPS/LTCH PPS final rule (77 FR 53675), we codified the definition of “applicable period” in the regulations at 42 CFR 412.152 as the 3-year period from which data is collected in order to calculate excess readmissions ratios and adjustments for the fiscal year, which includes aggregate payments for excess readmissions and aggregate payments for all discharges used in the calculation of the payment adjustment.

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 56974 through 56975), for FY 2017, consistent with the definition specified at §412.152, we established an “applicable period” for the Hospital Readmissions Reduction Program to be the 3-year period from July 1, 2012 through June 30, 2015. In other words, the excess readmissions ratios and the payment adjustment (including aggregate payments for excess readmissions and aggregate payments for all discharges used in the calculation of the payment adjustment) for FY 2017 were calculated using data from the 3-year time period of July 1, 2012 through June 30, 2015.

In this proposed rule, for FY 2018, consistent with the definition specified at §412.152, we are proposing that the “applicable period” for the Hospital Readmissions Reduction Program would be the 3-year period from July 1, 2013 through June 30, 2016. In other words, we are proposing that the excess readmissions ratios and the payment adjustment (including aggregate payments for excess readmissions and aggregate payments for all discharges) for FY 2018 would be calculated using data from the 3-year time period of July 1, 2013 through June 30, 2016.

We are inviting public comment on this proposal.

6. Proposed Calculation of Aggregate Payments for Excess Readmissions for FY 2018

Section 1886(q)(3)(B) of the Act specifies the ratio used to calculate the adjustment factor under the Hospital Readmissions Reduction Program. It states that the ratio is equal to 1 minus the ratio of—(i) the aggregate payments for excess readmissions and (ii) the aggregate payments for all discharges. For a detailed discussion on the methodology for the calculation of aggregate payments for excess readmissions, we refer readers to the FY 2013 IPPS/LTCH PPS final rule (77 FR 53387 through 53397). We also have codified our calculation of “aggregate payments for excess readmissions” and “aggregate payments for all discharges,” as well as a current methodology for calculating the numerator of the ratio (aggregate payments for excess readmissions) and the denominator of the ratio (aggregate payments for all discharges) at 42 CFR 412.152 through 412.154.

The Hospital Readmissions Reduction Program currently includes the following six applicable conditions: Acute myocardial infarction (AMI); heart failure (HF); pneumonia (PN); total hip arthroplasty/total knee arthroplasty (THA/TKA); chronic obstructive pulmonary disease (COPD); and Coronary Artery Bypass Graft (CABG) Surgery.

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 56975 through 56977), we adopted the methodology to include CABG in the calculation of the readmissions payment adjustment for FY 2017. Specifically, we discussed how the addition of CABG applicable conditions would be included in the calculation of the aggregate payments for excess readmissions (the numerator of the readmissions payment adjustment). We note that this policy did not alter our established methodology for calculating aggregate payments for all discharges (that is, the denominator of the ratio).

When calculating the numerator (aggregate payments for excess readmissions), we determine the base operating DRG payments for the applicable period. To determine the base operating DRG payment amount for an individual hospital for such applicable period for such condition, we use Medicare inpatient claims from the MedPAR file with discharge dates that are within the same applicable period to calculate the excess readmissions ratio. We use MedPAR claims data as our data source for determining aggregate payments for excess readmissions and aggregate payments for all discharges, as this data source is consistent with the claims data source used in IPPS rulemaking to determine IPPS rates.

For FY 2018, we are proposing to use MedPAR claims with discharge dates that are on or after the last day of the applicable period, as our data source (that is, the March updates of the respective Federal fiscal year MedPAR files) for the final rule.

In this proposed rule, for FY 2018, we are proposing to determine aggregate payments for excess readmissions and
aggregate payments for all discharges using data from MedPAR claims with discharge dates that are on or after July 1, 2013, and no later than June 30, 2016. However, we note that, for the purpose of modeling the proposed FY 2018 readmissions payment adjustment factors for this proposed rule, we used excess readmissions ratios for applicable hospitals from the FY 2017 Hospital Readmissions Reduction Program applicable period. For the FY 2018 IPPS/LTCH PPS final rule, applicable hospitals will have had the opportunity to review and correct data from the proposed FY 2018 applicable period of July 1, 2013 to June 30, 2016, before they are made public under our policy regarding the preview and reporting of hospital-specific information, which we discussed in the FY 2013 IPPS/LTCH PPS final rule (77 FR 53374 through 53401).

In this proposed rule, for FY 2018, we are proposing to use MedPAR data from July 1, 2013 through June 30, 2016. Specifically, for this proposed rule, we are using the following MedPAR files:
- March 2014 update of the FY 2013 MedPAR file to identify claims within FY 2013 with discharge dates that are on or after July 1, 2013;
- March 2015 update of the FY 2014 MedPAR file to identify claims within FY 2014;
- March 2016 update of the FY 2015 MedPAR file to identify claims within FY 2015;

For the final rule, we are proposing to use the same MedPAR files as listed above for claims within FY 2013, FY 2014 and FY 2015, and for claims within FY 2016, we are proposing to use the March 2017 update of the FY 2016 MedPAR file.

For a discussion of how we identified the applicable conditions to calculate the aggregate payments for excess readmissions for FY 2017, we refer readers to the FY 2017 IPPS/LTCH PPS final rule (81 FR 56975 through 56977).

Under our current methodology, in identifying the applicable conditions to calculate the aggregate payments for excess readmissions, we apply the same exclusions to the claims in the MedPAR file as are applied in the measure methodology for each of the applicable conditions. In this proposed rule, for FY 2018, we are proposing to continue to apply the same exclusions to the claims in the MedPAR file as we applied for FY 2017, namely, ICD–9–CM codes used to identify each applicable condition, and for CABG and COPD applicable conditions.

For a discussion of how we identified the applicable conditions to calculate the aggregate payments for excess readmissions for FY 2017, we refer readers to the FY 2017 IPPS/LTCH PPS final rules (80 FR 49539; 81 FR 56976) for a list of these exclusions. Updates to these exclusions will be posted on the QualityNet Web site at: http://www.QualityNet.org > Hospital-Inpatient > Claims-Based Measures > Readmission Measures > Measure Methodology.

Furthermore, under our current methodology we only identify Medicare fee-for-service (FFS) claims that meet the criteria described above for each applicable condition to calculate the aggregate payments for excess readmissions (that is, claims paid for under Medicare Part C, Medicare Advantage, are not included in this calculation). This policy is consistent with the methodology to calculate excess readmissions ratios based solely on admissions and readmissions for Medicare FFS patients. Therefore, consistent with our established methodology, for FY 2018, we are proposing to continue to exclude admissions for patients enrolled in Medicare Advantage as identified in the Medicare Enrollment Database.

Under our existing policy, we identify eligible hospitalizations and readmissions of Medicare patients discharged from an applicable hospital having a principal diagnosis for the measured condition in an applicable period (76 FR 51669). As described above, the proposed 3-year applicable period for FY 2018 of July 1, 2013 through June 30, 2016 includes discharges occurring in four Federal FYs (FY 2013, FY 2014, FY 2015, and FY 2016). Diagnoses and procedure codes for discharges occurring prior to October 1, 2015 were reported under the ICD–9–CM code set. Effective with discharges occurring on or after October 1, 2015 (FY 2016), diagnoses and procedure codes are reported under the ICD–10–CM and ICD–10–PCS code sets. Thus, for the proposed FY 2018 applicable period, the discharge diagnoses for each applicable condition would be based on a list of specific ICD–9–CM or ICD–10–CM and ICD–10–PCS code sets, as applicable, for that condition.

In this proposed rule, to identify the discharges for each applicable condition for FY 2018 to calculate the aggregate payments for excess readmissions for an individual hospital, we are proposing to identify each applicable condition, using, for FY 2013, FY 2014 and FY 2015, the appropriate ICD–9–CM codes, and for FY 2016, the appropriate ICD–10–CM and ICD–10–PCS code sets. This proposal is consistent with our methodology to identify the discharges for each applicable condition to calculate the aggregate payments for excess readmissions (76 FR 51673 through 51676). The ICD–9–CM codes for the AMI, HF, PN, THA/TKA, COPD, and CABG applicable conditions can be found on the QualityNet Web site at: http://www.QualityNet.org > Hospital-Inpatient > Claims-Based Measures > Readmission Measures > Measure Methodology. For a complete list of the ICD–9–CM codes we are proposing to use to identify the applicable conditions, we refer readers to the following tables of the measure methodology reports on the QualityNet Web site:
  ++ Table D.1.1—ICD–9–CM Codes for AMI Cohort (page 79).
  ++ Table D.2.1—ICD–9–CM Codes for COPD Cohort (page 83).
  ++ Table D.3.1—ICD–9–CM Codes for Inclusion in HF Cohort (page 89).
  ++ Table D.4.1—ICD–9–CM Codes for Pneumonia Cohort (page 94).
- 2016 Measure Updates: THA/TKA and CABG Readmission (THA and/or TKA-Version 4.0, CABG-Version 2.0: 2016 Procedure-Specific Readmission Measures Updates and Specifications Report)—
  ++ Table D.1.1—ICD–9–CM Codes Used to Identify Eligible CABG Procedures (page 49).
  ++ Table D.2.1—ICD–9–CM Codes Used to Identify Eligible THA/TKA Procedures (page 58).

A detailed list of the condition-specific and procedure-specific reports detailing the ICD–10–CM and ICD–10–PCS code sets we are proposing to use to identify the applicable conditions for the period from October 1, 2015 to June 30, 2016 is not yet publicly available. However, we anticipate the 2017 AMI, HF, Pneumonia, COPD, Stroke, THA/TKA, and CABG Readmission Measures Updates and Specifications Report, will be available by mid-April and can be accessed at: http://www.QualityNet.org > Hospital-Inpatient > Claims-Based Measures > Readmission Measures > Measure Methodology. We are currently making a list of the ICD–10–CM and ICD–10–PCS code sets used to identify the applicable conditions for this proposed rule, titled ICD–10–CM Codes for Inclusion in the Hospital Readmissions Reduction Program Applicable Conditions for FY 2018 Proposed Rule, available on the Hospital Readmissions Reduction Program page on the CMS Web site at: https://
In summary, for FY 2018, we are proposing to calculate aggregate payments for excess readmissions, using MedPAR claims from July 1, 2013 through June 30, 2016, to identify applicable conditions based on the same ICD–9–CM codes or ICD–10–CM and ICD–10–PCS code sets, as applicable, used to identify the conditions for the readmissions measures, and to apply the proposed exclusions for the types of admissions (as previously discussed). We are not proposing any changes to our existing methodology for calculating "aggregate payments for excess readmissions" for each hospital (the numerator of the ratio). Specifically, to calculate aggregate payments for excess readmissions for each hospital, we are proposing to calculate the base operating DRG payment amounts for all claims in the 3-year applicable period for each applicable condition (AMI, HF, PN, COPD, THA/TKA, and CAGB) based on the claims we have identified as described above. Once we have calculated the base operating DRG amounts for all the claims for the six applicable conditions, we are proposing to sum the base operating DRG payments amounts by each condition, resulting in six summed amounts, one amount for each of the six applicable conditions. We are proposing to then multiply the amount for each condition by the respective excess readmissions ratio minus 1 when that excess readmissions ratio is greater than 1, which indicates that a hospital has performed, with respect to readmissions for that applicable condition, worse than the average hospital with similar patients. Each product in this computation represents the payments for excess readmissions for that condition. We are proposing to then sum the resulting products which represent a hospital’s proposed “aggregate payments for excess readmissions” (the numerator of the ratio). Because this calculation is performed separately for each of the six conditions, a hospital’s excess readmissions ratio must be less than or equal to 1 on each measure to avoid CMS’ determination that there were payments made by CMS for excess readmissions (resulting in a payment reduction under the Hospital Readmissions Reduction Program). In other words, in order to avoid a payment adjustment a hospital’s excess readmissions ratio must be less than or equal to 1 on each measure. We note that we are not proposing any changes to our existing methodology to calculate “aggregate payments for all discharges” (the denominator of the ratio).

Section 1886(q)(3)(A) of the Act defines the “adjustment factor” for an applicable hospital for a fiscal year as equal to the greater of: (i) The ratio described in subparagraph (B) for the hospital for the applicable period (as defined in paragraph (5)(D)) for such fiscal year; or (ii) the floor adjustment factor specified in subparagraph (C).

Section 1886(q)(3)(B) of the Act, in turn, describes the ratio used to calculate the adjustment factor. Specifically, it states that the ratio is equal to 1 minus the ratio of—(i) the aggregate payments for excess readmissions and (ii) the aggregate payments for all discharges. The calculation of this ratio is codified at §412.154(c)(1) of the regulations and the floor adjustment factor is codified at §412.154(c)(2) of the regulations. Section 1886(q)(3)(C) of the Act specifies the floor adjustment factor at 0.970 for FY 2015 and subsequent fiscal years.

Consistent with section 1886(q)(3)(C) of the Act, codified at §412.154(c)(2), for FY 2015 and subsequent years, the adjustment factor is either the greater of the ratio or the floor adjustment factor of 0.9700. In other words, a hospital subject to the Hospital Readmissions Reduction Program will have an adjustment factor that is between 1.0000 (no reduction) and 0.9700 (greatest possible reduction). Under our established policy, the ratio is rounded to the fourth decimal place.


a. Background

As described above, section 1886(q)(3)(D) of the Act requires the Secretary to group hospitals and apply a methodology that allows for separate comparisons of hospitals within groups in determining a hospital’s adjustment factor for payments of discharges beginning in FY 2019. Furthermore, section 1886(q)(3)(D) of the Act directs the Secretary to define groups of hospitals, based on their overall proportion, of the inpatients who are entitled to, or enrolled for, benefits under part A, and who are full-benefit dual eligible individuals (as defined in section 1935(c)(6)(A) of the Act).38 Under these statutory requirements, hospitals are grouped based on the proportion or ratio of full-benefit dual eligible patients (numerator) to the hospital’s Medicare inpatient stays (denominator). The Act specifies that in defining groups, the Secretary shall consult the MedPAC and may consider the analysis done by MedPAC in preparing the portion of its

As described above, section 1886(q)(3)(D) of the Act requires the Secretary to group hospitals and apply a methodology that allows for separate comparisons of hospitals within groups in determining a hospital’s adjustment factor for payments of discharges beginning in FY 2019. Furthermore, section 1886(q)(3)(D) of the Act directs the Secretary to define groups of hospitals, based on their overall proportion, of the inpatients who are entitled to, or enrolled for, benefits under part A, and who are full-benefit dual eligible individuals (as defined in section 1935(c)(6)(A) of the Act).38 Under these statutory requirements, hospitals are grouped based on the proportion or ratio of full-benefit dual eligible patients (numerator) to the hospital’s Medicare inpatient stays (denominator). The Act specifies that in defining groups, the Secretary shall consult the MedPAC and may consider the analysis done by MedPAC in preparing the portion of its

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38 Section 1935(c)(6)(A) of the Act defines “full-benefit dual eligible individual” as, for a State for a month, an individual who—(i) has coverage for the month for covered part D drugs under a prescription drug plan under part D of title XVIII, or under an MA–PD plan under part C of such title; and (ii) is determined eligible by the State for medical assistance for full benefits under this title for such month under section 1902(a)(10)(A) or 1902(a)(10)(C) [of the Act], by reason of section 1902(l) [of the Act], or under any other category of eligibility for medical assistance for full benefits under this title, as determined by the Secretary.
report submitted to Congress in June 2013 relating to readmissions.

b. Proposed Data Sources Used To Determine Dual Eligibility

In this proposed rule, we are proposing to identify full-benefit dual status (numerator) using dual eligibility status data where the original data source is the State Medicare Modernization Act (MMA) file of dual eligibility which States submit to CMS monthly. The State MMA file is considered the most current and most accurate source of data for identifying dual eligible beneficiaries since it is also used for operational purposes related to the administration of Part D benefits. Under our proposal, an individual would be counted as a full-benefit dual patient if the beneficiary was identified as full-benefit dual status in the State MMA files for the month he/she was discharged from the hospital.

We are inviting public comment on this proposal.

For this proposed rule, we considered two alternative definitions of total number of Medicare patients (denominator) that could be used to calculate each hospital's proportion of dual eligible patients. We are proposing to define the proportion of full-benefit dual eligible beneficiaries as the proportion of dual eligible patients among all Medicare FFS and Medicare Advantage stays. This is our preferred approach because using the proportion of dual eligible patients calculated among all Medicare FFS and managed care patients more accurately represents the proportion of dual eligible patients served by the hospital, particularly for hospitals in States with high managed care penetration rates. For example, Hospital A located in Arizona has a high managed care penetration rate. When stratified based on the proportion of dual eligibles, calculated among Medicare FFS and managed care patients, Hospital A was assigned to the top quintile of proportion of dual eligibles and its payment adjustment calculated based on its ERR relative to the threshold for the top quintile. When stratified based on the proportion of dual eligible among only Medicare FFS patients, Hospital A was assigned to the second quintile and its payment adjustment calculated relative to the threshold of the second quintile. Its classification when managed care patients are included more accurately identifies the social risk of the patients Hospital A serves, compared to its classification if only the FFS population is included.

However, because the Hospital Readmissions Reduction Program payment adjustment is only applied to Medicare FFS payments, and is based on excess readmissions among Medicare FFS patients only, we are including an alternative to define the proportion of full-benefit dual eligible beneficiaries as only Medicare FFS stays. Under both approaches, we are proposing to use the MedPAR files, the same data source used to calculate the payment adjustment factors, to identify total hospital stays as this is the best available claims data that is readily publicly available. However, in developing our proposal, we also considered using other data sources such as the CMS integrated data repository (IDR), which may incorporate managed care claims more consistently to calculate total hospital stays, but it is currently not readily available to the public. We are inviting stakeholder input on the most appropriate data source to identify total hospital stays and whether such stays should include all Medicare FFS and Medicare Advantage stays or only Medicare FFS stays.

We are inviting public comment on our preferred proposals and alternative considerations.

c. Proposed Data Period Used To Define Dual Eligibility

Consistent with the requirement of the statute, we are proposing to group or stratify hospitals based on the proportion of full-benefit dual eligible patients determined under the proposals discussed above and are proposing to define the proportion of full-benefit dual eligible beneficiaries as the number of dual eligible patients discharged during the 3-year applicable period under the Hospital Readmissions Reduction Program. For this proposed rule, we considered two alternatives for the data period used to define dual eligibility, a 3-year period corresponding to the performance period, and a 1-year period, which would be calculated over the most recent year for which complete data is available.

While both data periods would include the most recently available data to define dual eligibility, our proposal to use a 3-year period accounts for the influence of social risk factors on the excess readmissions ratio (ERR) because the proportion of dual eligible patients is measured over the full period when they influenced the likelihood of excess readmissions. However, the most recent 1-year period would capture the most recent population served by the hospital and may enable a more accurate stratification to calibrate the impact of payment adjustments to the proportion of dual eligible patients that the hospital currently serves.

We are inviting public comment on our preferred proposal and alternative considerations.

9. Provisions for the Proposed Payment Adjustment Methodology for FY 2019:

Payment Adjustment Methodology for Assigning Hospitals to Peer Groups

For this proposed rule, we considered three alternative methodologies for assigning hospitals to peer groups. For the reasons discussed below, our preferred approach is to stratify hospitals into quintiles (five peer groups). However, we are also seeking public comment on stratifying hospitals into two and 10 peer groups.

To understand the impact on payment adjustments of stratifying hospitals into different number of peer groups, we conducted an analysis that estimated payment adjustments when stratifying hospitals into two, five (quintiles), or 10 (deciles) peer groups. Two and 10 peer groups were considered to align with previous research conducted by MedPAC and ASPE that assessed impacts from stratifying hospitals into two or 10 groups. MedPAC's analysis stratified hospitals into 10 peer groups when setting the target rate used to compare hospital performance. ASPE's analysis stratified hospitals into two and 10 peer groups to calculate payment adjustments. Our analysis showed that using five peer groups allows for more precisely defined peer groups than is possible with a grouping of two, while ensuring that the number of hospitals is sufficient to represent a peer group, even for measures, like CABG, in which only a minority of hospitals are subject to a payment adjustment.

We note, as the number of groupings increase, hospitals became more similar within their peer groups with respect to proportion of dual eligible patients in their patient population. Hence, payment adjustments are more closely related to the proportion of dual eligibles, and to the possible influence on the likelihood of readmission resulting from small variations in patient populations. We also observed that increasing the number of peer groups also increases the likelihood that hospitals with similar exposure to dual eligible patients will be compared to different thresholds in the payment adjustment formula. Deciles cover a narrow range of dual eligible patient proportions in each peer group, so small differences in proportion are likely to result in differences in peer group assignment and corresponding comparison thresholds used in the payment adjustment formula. This
problem is compounded by the small number of hospitals in deciles. When
the number of hospitals is small, peer group thresholds or distributions and
the resulting payment adjustments are less predictable.

Stratifying hospitals into two peer groups is a simpler method and reduces
the likelihood that similar hospitals are assigned different payment adjustments.
However, this approach yields peer groups with a more heterogeneous mix of
hospitals assigned to each group and weakens the relationship between the
payment adjustment and the hospital’s patient population. When the impact on
payments of different peer group definitions was tested using the various
methods of incorporating stratification into the payment formula, we found a
substantial reduction in penalties (measured as the share of payment
adjustments as a percentage of total payments) to safety-net hospitals,
defined as hospitals in the highest quintile for disproportionate share (DSH) payments, from stratification into quintiles compared to stratification into
two groups. Furthermore, our analysis found a similar impact on the share of
total payments borne as payment adjustments by safety-net hospitals from
stratifying hospitals into quintiles and deciles, suggesting that the benefit to
safety-net hospitals from increasing the number of strata would be small. For
example, using the preferred modified payment formula, proposed below,
across the current set of six conditions, we found that for safety-net hospitals,
payment adjustment as a proportion of total payments decreased from a
baseline of 0.64 percent to 0.59 percent with two groups, 0.55 percent with
quintiles and 0.54 percent with deciles.

Based on the analysis described above, we are proposing to stratify
hospitals into quintiles (five peer groups) because it creates peer groups
that accurately reflect the relationship between the proportion of dual eligibles
in the hospital’s population without the disadvantage of establishing a larger
number of peer groups.

In this proposed rule, our preferred proposal and alternative
calculation methodologies for calculating the payment adjustment factor. Our
preferred approach is assessing performance compared to the peer
group median ERR, rather than the current threshold of 1.0000, and scaling
hospital payment adjustments by a neutrality modifier. However, we are
seeking public comment on three additional approaches—using the mean
ERR plus a neutrality modifier, a budget neutralizing ERR, and a standardized
ERR plus a neutrality modifier.

(1) Median ERR Plus a Neutrality Modifier

In this proposed rule, our preferred approach is using the median ERR plus
a neutrality modifier. We would use the median ERR for the hospital’s peer
group in place of 1.0000, which is the approximate mean and median of the
baseline distribution, in the current payment adjustment formula. The
payment adjustment formula would then be:

\[
P = 1 - \min\{0.3, \sum_{dx} Payment(dx) \cdot \max\{(ERR(dx) - 1.0000), 0\} \}
\]

where \(dx\) is AMI, HF, pneumonia, COPD, THA/TKA or CABG. In our
analyses, we modified the payment adjustment formula by replacing the
current threshold ERR of 1.0000 with a peer group specific threshold.

In adopting a methodology for achieving budget neutrality, our priority
is to adopt a simplified and well-known metric that allows us to be more
transparent in our methodology and reduces the penalty on safety-net
hospitals, while not disproportionally increasing the penalty to non-safety-net
hospitals. In developing policy options to implement the budget neutrality
requirement, we analyzed the following alternatives to evaluate the financial
impacts:

- Using the median ERR for the hospital’s peer group in place of 1.0000
  in the payment adjustment formula and applying a uniform modifier to maintain
  budget neutrality;
- Using the mean ERR for the hospital’s peer group in place of 1.0000
  in the payment adjustment formula and applying a uniform modifier to maintain
  budget neutrality;

We are inviting public comment on
our preferred proposal and alternative
considerations.

10. Provisions for the Proposed Payment Adjustment Methodology for FY 2019:
Proposed Payment Adjustment Formula Calculation Methodology

a. Background

As described above, section 1886(q)(3)(D)(iv) of the Act requires the
Secretary to design the methodology to implement this subsection so that the
estimated total amount of Medicare savings under this subsection (stratified
methodology) equals the estimated total amount of Medicare savings that would
otherwise occur under this subsection (current methodology) if this
subsection did not apply (that is, maintain budget neutrality).

We analyzed several modifications of the payment adjustment formula to
assess payment reductions based on a hospital’s performance compared to
performance of other hospitals in its peer group. The current readmissions
payment adjustment can be written as

\[
1 - \min\{0.3, \sum_{dx} Payment(dx) \cdot \max\{(ERR(dx) - 1.0000), 0\} \}
\]

We are inviting public comment on
our preferred proposal and alternative
considerations.
The payment reduction \((1-P)\) resulting from use of the median ERR for the peer group is scaled by a neutrality modifier \((N_M)\) to achieve budget neutrality. To calculate the neutrality modifier, we estimate total Medicare savings across all hospitals under the current method and under the proposed stratified method, in the absence of a modifier. We then calculate a multiplicative factor that, when applied to each hospital’s adjustment calculated using the stratified method, would equate total Medicare savings from that method to total Medicare savings under the current method. Total Medicare savings and the neutrality modifier will be calculated using the same payment data. These data will consist of the most recently available full year of MedPAR data. For example, if the payment reduction for a hospital \((1-P)\) equals 0.00748 when using the median threshold, then under the median plus neutrality modifier method it would equal \(NM*0.00748 = 0.9545\) * 0.00748 = 0.00714, where the neutrality modifier was equal to 0.9545. Thus, the hospital’s payment adjustment factor \((P)\) would equal 0.9929 \((1-0.00748)\) in the absence of the neutrality modifier, and 0.9929 \((1-0.00714)\) when the modifier is added.

(2) Mean ERR Plus a Neutrality Modifier

We also analyzed the use of the mean ERR plus a neutrality modifier to calculate the readmissions adjustment factor. Just like the median ERR plus neutrality modifier approach mentioned above, the mean ERR for the hospital’s peer group would be used in place of 1.0000 in the payment adjustment formula. The payment adjustment formula would then be:

\[
P = 1 - \min \left\{ 0.03, \sum_{dx} K_p \cdot \frac{\text{Payment}(dx) \cdot \max \left( \left( \text{ERR}(dx) - \text{Mean peer group ERR}(dx) \right), 0 \right)}{\text{All payments}} \right\}
\]

(3) Budget Neutralizing ERR

We also analyzed using a budget neutralizing ERR in which penalties are assessed based on the difference between the hospital’s ERR and the budget neutralizing ERR. The payment adjustment formula would be:

\[
P = 1 - \min \left\{ 0.03, \sum_{dx} \frac{\text{Payment}(dx) \cdot \max \left( \left( \text{ERR}(dx) - \text{budget neutralizing ERR}(dx) \right), 0 \right)}{\text{All payments}} \right\}
\]

(4) Standardized ERR Plus a Neutrality Modifier

We also analyzed using a standardized ERR in which penalties are assessed by determining the mean and standard deviation of the ERRs across all hospitals. The payment adjustment formula would be calculated by dividing hospitals into strata based on a hospital’s proportion of dual eligible patients. The current ERRs would then be transformed to create a new standardized distribution of ERRs within each stratum with the same mean and standard deviation as the original mean and standard deviation across all hospitals.

\[
P = 1 - \min \left\{ 0.03, \sum_{dx} \frac{\text{Payment}(dx) \cdot \max \left( \left( \text{ERR}(dx) - \text{standard neutralizing ERR}(dx) \right), 0 \right)}{\text{All payments}} \right\}
\]

where \(S_p(dx)\) and \(\mu_p(dx)\) are the standard deviation and mean of the current ERR distribution for a condition \((dx)\), and \(S_p(dx)\) and \(\mu_p(dx)\) are the standard deviation and mean of the peer group ERR distribution for that \(dx\). The standardized ERRs have a mean of 1 and a standard deviation equal to the standard deviation of ERRs across all hospitals in the peer group for that condition. The standardized ERRs are compared to 1.0000 in the payment adjustment formula to determine excess readmissions. The payment reduction \((1-P)\) resulting from use of the standardized ERR is then scaled by a neutrality modifier \((N_M)\) to achieve budget neutrality.

c. Analysis

As mentioned above, in adopting a methodology for achieving budget neutrality, our priority is to adopt a simplified and well-known metric that allows us to be more transparent in our methodology and reduces the penalty on safety-net hospitals, while not disproportionately increasing the penalty to non-safety-net hospitals. To assess the expected impact on hospital payments resulting from the changes to the formula, we simulated hospitals’ readmission adjustment factors under different stratified thresholds. Readmissions adjustment factors were calculated using total base operating DRG payment amounts for each hospital as well as total base DRG payment amounts for each of the six measure cohorts (AMI, HF, pneumonia, COPD, CABG, THA/TKA) included in the FY 2018 program. We used DRG payment information for the period July 1, 2012 through June 30, 2015. Furthermore, to estimate the dollar amount of the penalty and the share of payments the penalty represents, we used total base operating DRG payments among Medicare FFS claims from the FY 2015 MedPAR data file.

All four methods support the agency’s efforts to reduce the payment adjustment for safety-net hospitals. We are proposing to use the median ERR plus a neutrality modifier because it creates a standard where a hospital’s ERR is subject to payment reduction when a hospital’s performance as measured by the ERR is worse than that of half the other hospitals in its peer group. The median ERR plus neutrality modifier is preferred to the mean ERR plus neutrality modifier because the median represents a consistent standard (that is, 50th percentile) for the hospital’s rank within its peer group, while the rank corresponding to the mean changes between years, cohorts and peer groups. The median ERR plus neutrality modifier substantially reduces the penalty as a share of total payments (from 0.64 percent to 0.55 percent with quintile peer groups) and penalty per discharge (from $157 to $135) for safety-net hospitals while not disproportionately increasing the payment reduction amount for non-safety-net hospitals (from 0.61 percent...
to 0.63 percent as share of total payments). The median ERR plus neutrality modifier is also preferred because it achieves more precise budget neutrality than the budget neutralizing ERR. Below we show the estimated total Medicare savings under the current and stratified methodology used to assess budget neutrality.

<table>
<thead>
<tr>
<th>Method</th>
<th>Estimated total medicare savings</th>
<th>Difference between stratified and current methodology</th>
<th>Percentage difference between stratified and current methodology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Current methodology</td>
<td>$532,948,318</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Mean plus neutrality modifier</td>
<td>532,949,006</td>
<td>$688</td>
<td>&lt;0.00</td>
</tr>
<tr>
<td>Median plus neutrality modifier</td>
<td>532,946,272</td>
<td>($2,046)</td>
<td>&lt;0.00</td>
</tr>
<tr>
<td>Budget neutralizing ERR</td>
<td>533,199,304</td>
<td>250,985</td>
<td>0.05</td>
</tr>
<tr>
<td>Standardized ERR plus neutrality modifier</td>
<td>532,948,288</td>
<td>($30)</td>
<td>&lt;0.00</td>
</tr>
</tbody>
</table>

Source: FY 2017 Hospital Readmissions Reduction Program Final Rule Results. Results are based on July 1, 2012, through June 30, 2015, discharges among subsection (d) and Maryland hospitals only. Although data from all subsection (d) and Maryland hospitals are used in calculations of each hospital’s Excess Readmission Ratio (ERR), this table does not include results for Maryland hospitals. Hospital Characteristics are based on the FY 2017 final rule Impact File. Hospitals are stratified into quintiles based on the proportion of dual-eligible beneficiaries among Medicare fee-for-service and managed care patients discharged between July 1, 2012, through June 30, 2015.

When we analyzed the other options, we found that the mean threshold permits a higher standard to be set if hospitals in the peer group have performance well above the midpoint but not far below, or a lower standard if hospitals are more likely to have very high rates. In our testing, the mean plus modifier resulted in lower penalties for safety-net hospitals (0.52 percent as a share of total payments compared to 0.55 percent for the median plus modifier). However, our preferred approach of the median is based on the judgment that the standard reflected by the threshold should not be affected by hospitals with unusually strong or weak performance in the peer group. Like the median, the budget neutralizing ERR threshold approach imposes a consistent rank-based standard across peer groups. However, this method is not preferred since it is more complex, less intuitive and results in greater divergence between total payment adjustments under the stratified and current methodologies than approaches using a neutrality modifier (differing from the current methodology by approximately 0.05 percent of total payments when simulated with quintile peer groups). The median uses the original distribution of hospital ERR estimates, based on their relationship to a national standard, and represents the most precise possible measures of their performance under that standard. Using a standardized ERR within each peer group compares a hospital’s performance to other hospitals in the peer group. In contrast, using the mean or median threshold adjusts penalties based on a hospital’s relative performance within the peer group, but the performance indicator of the ERR retains the comparison to the mean performance of all hospitals across all peer groups. However, comparing the ERR to the mean or median for each peer group is a more straightforward methodology than re-standardizing ERRs. The median is preferred to the standardized ERR because, as with the budget neutralizing ERR, the median is less complex and more intuitive. Using a less complex and well-known metric, will create a more transparent methodology since it will be easier for hospitals and other stakeholders to replicate the calculation of the median ERRs.

The impact of the proposed changes to the payment adjustment formula for the budget neutral considered methods, by peer group options, for safety-net and non-safety-net hospitals is shown in the table below. The table includes three penalty metrics: Average payment reduction, total Medicare savings, and share of payment adjustments as a percentage of total payments. The average payment reduction shows the average reduction in Medicare DRG payments for safety-net and non-safety-net hospitals. The total Medicare savings column shows the total estimated penalties borne by safety-net and non-safety-net hospitals under each approach. Since the payment reduction is applied to hospitals’ base DRG payments, hospitals with more discharges will contribute a larger amount of Medicare savings to the group total of Medicare savings. Furthermore, since there are fewer safety-net than non-safety-net hospitals, as safety-net is defined as hospitals in the top quintile of DSH patient percentage, the total Medicare savings for non-safety-net hospitals are inherently much larger than for safety-net hospitals. Therefore, to compare the financial impact of the program on hospitals in each group we calculated the payment adjustment as a proportion of DRG payments. Using this metric allows comparison across the different methodologies where the total base operating DRG payments are different between different groups of hospitals and is a more accurate indication of the financial impact on the group. For example, under the current methodology, the payment adjustment as a proportion of all DRG payments among safety-net hospitals is 0.64 percent.
## Comparison of Penalty Metrics by Threshold Methods and Peer Group Options for All Hospitals, Safety-Net, and Non-Safety-Net Hospitals

<table>
<thead>
<tr>
<th>Stratification approach and payment formula methodology</th>
<th>Average payment reduction ((1-P)_a) (percent)</th>
<th>Total Medicare savings</th>
<th>Payment adjustment as a proportion of all DRG payments (percent)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Current methodology:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Safety-net hospitals</td>
<td>0.62</td>
<td>$109,142,525</td>
<td>0.64</td>
</tr>
<tr>
<td>Non-safety-net hospitals</td>
<td>0.61</td>
<td>429,805,793</td>
<td>0.61</td>
</tr>
<tr>
<td>Approach 1: Two equal peer groups based on the proportion of dual-eligible beneficiaries</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median plus neutrality modifier (neutrality modifier = 0.9558):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Safety-net hospitals</td>
<td>0.56</td>
<td>100,205,115</td>
<td>0.59</td>
</tr>
<tr>
<td>Non-safety-net hospitals</td>
<td>0.61</td>
<td>432,741,358</td>
<td>0.62</td>
</tr>
<tr>
<td>Mean plus neutrality modifier (neutrality modifier = 1.0191):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Safety-net hospitals</td>
<td>0.54</td>
<td>97,837,278</td>
<td>0.57</td>
</tr>
<tr>
<td>Non-safety-net hospitals</td>
<td>0.61</td>
<td>435,112,491</td>
<td>0.63</td>
</tr>
<tr>
<td>Budget neutralizing ERR:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Safety-net hospitals</td>
<td>0.55</td>
<td>98,208,670</td>
<td>0.58</td>
</tr>
<tr>
<td>Non-safety-net hospitals</td>
<td>0.61</td>
<td>435,216,961</td>
<td>0.63</td>
</tr>
<tr>
<td>Standardized ERR plus neutrality modifier (neutrality modifier = 0.9796):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Safety-net hospitals</td>
<td>0.55</td>
<td>98,468,430</td>
<td>0.58</td>
</tr>
<tr>
<td>Non-safety-net hospitals</td>
<td>0.61</td>
<td>434,478,852</td>
<td>0.63</td>
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<tr>
<td>Approach 2: Quintiles based on the proportion of dual-eligible beneficiaries</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median plus neutrality modifier (neutrality modifier = 0.9546):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Safety-net hospitals</td>
<td>0.52</td>
<td>93,878,536</td>
<td>0.55</td>
</tr>
<tr>
<td>Non-safety-net hospitals</td>
<td>0.62</td>
<td>439,067,736</td>
<td>0.63</td>
</tr>
<tr>
<td>Mean plus neutrality modifier (neutrality modifier = 1.0135):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Safety-net hospitals</td>
<td>0.49</td>
<td>89,182,424</td>
<td>0.52</td>
</tr>
<tr>
<td>Non-safety-net hospitals</td>
<td>0.62</td>
<td>443,766,582</td>
<td>0.64</td>
</tr>
<tr>
<td>Budget neutralizing ERR:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Safety-net hospitals</td>
<td>0.49</td>
<td>88,510,157</td>
<td>0.52</td>
</tr>
<tr>
<td>Non-safety-net hospitals</td>
<td>0.62</td>
<td>444,689,147</td>
<td>0.64</td>
</tr>
<tr>
<td>Standardized ERR plus neutrality modifier (neutrality modifier = 0.9710):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Safety-net hospitals</td>
<td>0.50</td>
<td>91,686,964</td>
<td>0.54</td>
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<tr>
<td>Non-safety-net hospitals</td>
<td>0.62</td>
<td>441,261,324</td>
<td>0.64</td>
</tr>
<tr>
<td>Approach 3: Deciles based on the proportion of dual-eligible beneficiaries</td>
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<tr>
<td>Median plus neutrality modifier (neutrality modifier = 0.9555):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Safety-net hospitals</td>
<td>0.51</td>
<td>91,881,047</td>
<td>0.54</td>
</tr>
<tr>
<td>Non-safety-net hospitals</td>
<td>0.62</td>
<td>441,068,999</td>
<td>0.64</td>
</tr>
<tr>
<td>Mean plus neutrality modifier (neutrality modifier = 1.0148):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Safety-net hospitals</td>
<td>0.48</td>
<td>87,289,962</td>
<td>0.51</td>
</tr>
<tr>
<td>Non-safety-net hospitals</td>
<td>0.62</td>
<td>445,653,065</td>
<td>0.64</td>
</tr>
<tr>
<td>Budget neutralizing ERR:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Safety-net hospitals</td>
<td>0.47</td>
<td>86,671,374</td>
<td>0.51</td>
</tr>
<tr>
<td>Non-safety-net hospitals</td>
<td>0.62</td>
<td>446,299,290</td>
<td>0.64</td>
</tr>
<tr>
<td>Standardized ERR plus neutrality modifier (neutrality modifier = 0.9713):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Safety-net hospitals</td>
<td>0.49</td>
<td>90,058,433</td>
<td>0.53</td>
</tr>
<tr>
<td>Non-safety-net hospitals</td>
<td>0.62</td>
<td>442,888,696</td>
<td>0.64</td>
</tr>
</tbody>
</table>

Notes: Results based on July 1, 2012 through June 30, 2015 discharges among subsection (d) and Maryland hospitals only. Although data from subsection (d) and Maryland hospitals are used in calculations of each hospital’s ERR, this table does not include results for Maryland hospitals. Hospitals are stratified based on the proportion of duals calculated among Medicare FFS and managed care patients for the FY 2017 performance period. Safety-net hospitals are defined as hospitals in the top quintile of DSH patient percentage. DSH patient percentage was calculated among all hospitals with a positive DSH value (including hospitals not eligible for DSH payments). a. The payment reduction shows what percentage of DRG payments hospitals will lose as a result of the program. This is slightly different than the adjustment factor that CMS applies, which is 1 minus the number reported here (that is, ranges from 0.97 to 1). b. Total Medicare savings is estimated by multiplying the payment reduction by total base operating DRG payments from July 1, 2014 through June 30, 2015. c. The group share of payment adjustments as a percentage of all DRG payments is calculated as the sum of total Medicare savings for the group of hospitals (that is, safety-net hospitals or non-safety-net hospitals) divided by total base operating DRG payments from July 1, 2014 through June 30, 2015 for the group of hospitals.

Our analysis also assesses the impact of the proposed changes to the payment adjustment formula on additional groups of hospitals. Variation in the impact of the proposed changes by hospital characteristics on the share of payment adjustments as a percentage of all DRG payments for the FY 2019 Hospital Readmissions Reduction Program, is shown in the table below. The table is based on results when hospitals are stratified into quintiles.
based on the proportion of dual eligible beneficiaries among Medicare FFS and managed care patients discharged between July 1, 2012, through June 30, 2015, our preferred approaches. The table shows the average share of payment adjustments as a percentage of all DRG payments for each group of hospitals. The group average is calculated as the sum of penalties for all hospitals with that characteristic over the sum of all DRG payments for those hospitals between July 1, 2014 and June 30, 2015. For example, under the current methodology, the average share of payment adjustments as a percentage of all DRG payments for urban hospitals is 0.61 percent. This means that total penalties for all urban hospitals is 0.61 percent of total payments for urban hospitals (that is the ratio of total penalties to total DRG payments is 0.61 percent). This metric allows us to compare the financial impact of the different methods for assessing penalties between hospitals with different number of beds even though larger hospitals tend to generate higher total Medicare savings since their payment reduction is applied to more DRG payments. Measuring the financial impact on hospitals as a proportion of total DRG payments allows us to account for differences in the amount of DRG payments for hospitals when comparing the financial impact of the program on different groups of hospitals, and allows comparison across the different methodologies between groups of hospitals with different numbers of eligible hospitals.

### Average Share of Payment Adjustments as a Percentage of All DRG Payments for Considered Approaches for the Hospital Readmissions Reduction Program, by Hospital Characteristic

<table>
<thead>
<tr>
<th>Hospital characteristics</th>
<th>Number of hospitals with characteristic</th>
<th>Current methodology (percent)</th>
<th>Median Plus neutrality modifier (neutrality modifier = 0.9546) (percent)</th>
<th>Mean plus neutrality modifier (neutrality modifier = 1.0135) (percent)</th>
<th>Budget neutralizing ERR (percent)</th>
<th>Standardized ERR plus neutrality modifier (neutrality modifier = 0.9710) (percent)</th>
</tr>
</thead>
<tbody>
<tr>
<td>All Hospitals</td>
<td>3,096</td>
<td>0.62</td>
<td>0.62</td>
<td>0.62</td>
<td>0.62</td>
<td>0.62</td>
</tr>
<tr>
<td>Geographic Location:</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Urban</td>
<td>2,304</td>
<td>0.61</td>
<td>0.62</td>
<td>0.62</td>
<td>0.62</td>
<td>0.62</td>
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<tr>
<td>Rural</td>
<td>792</td>
<td>0.65</td>
<td>0.62</td>
<td>0.60</td>
<td>0.60</td>
<td>0.60</td>
</tr>
<tr>
<td>Bed size:</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1–99 beds</td>
<td>1,113</td>
<td>0.57</td>
<td>0.57</td>
<td>0.56</td>
<td>0.56</td>
<td>0.57</td>
</tr>
<tr>
<td>100–199 beds</td>
<td>886</td>
<td>0.70</td>
<td>0.70</td>
<td>0.70</td>
<td>0.70</td>
<td>0.70</td>
</tr>
<tr>
<td>200–299 beds</td>
<td>453</td>
<td>0.65</td>
<td>0.66</td>
<td>0.66</td>
<td>0.66</td>
<td>0.66</td>
</tr>
<tr>
<td>300–399 beds</td>
<td>278</td>
<td>0.64</td>
<td>0.63</td>
<td>0.63</td>
<td>0.63</td>
<td>0.63</td>
</tr>
<tr>
<td>400–499</td>
<td>155</td>
<td>0.53</td>
<td>0.54</td>
<td>0.54</td>
<td>0.54</td>
<td>0.54</td>
</tr>
<tr>
<td>500 or more beds</td>
<td>211</td>
<td>0.57</td>
<td>0.57</td>
<td>0.57</td>
<td>0.57</td>
<td>0.57</td>
</tr>
<tr>
<td>By DSH Payment Eligibility:</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Not eligible</td>
<td>474</td>
<td>0.55</td>
<td>0.61</td>
<td>0.65</td>
<td>0.64</td>
<td>0.64</td>
</tr>
<tr>
<td>DSH payment eligible</td>
<td>2,622</td>
<td>0.63</td>
<td>0.62</td>
<td>0.61</td>
<td>0.61</td>
<td>0.61</td>
</tr>
<tr>
<td>By Teaching Status:</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-teaching</td>
<td>2,076</td>
<td>0.66</td>
<td>0.67</td>
<td>0.67</td>
<td>0.67</td>
<td>0.67</td>
</tr>
<tr>
<td>Teaching</td>
<td>1,020</td>
<td>0.59</td>
<td>0.58</td>
<td>0.58</td>
<td>0.58</td>
<td>0.58</td>
</tr>
<tr>
<td>Fewer than 100 residents</td>
<td>772</td>
<td>0.59</td>
<td>0.60</td>
<td>0.60</td>
<td>0.61</td>
<td>0.60</td>
</tr>
<tr>
<td>100 or more residents</td>
<td>248</td>
<td>0.57</td>
<td>0.55</td>
<td>0.54</td>
<td>0.54</td>
<td>0.55</td>
</tr>
<tr>
<td>By Type of Ownership:</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Government</td>
<td>490</td>
<td>0.54</td>
<td>0.53</td>
<td>0.53</td>
<td>0.53</td>
<td>0.53</td>
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<tr>
<td>Proprietary</td>
<td>779</td>
<td>0.79</td>
<td>0.79</td>
<td>0.80</td>
<td>0.80</td>
<td>0.79</td>
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<tr>
<td>Voluntary</td>
<td>1,827</td>
<td>0.59</td>
<td>0.59</td>
<td>0.59</td>
<td>0.59</td>
<td>0.59</td>
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<tr>
<td>DSH patient percentage:</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>1st</td>
<td>547</td>
<td>0.54</td>
<td>0.60</td>
<td>0.63</td>
<td>0.63</td>
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<tr>
<td>2nd</td>
<td>635</td>
<td>0.66</td>
<td>0.71</td>
<td>0.72</td>
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<tr>
<td>3rd</td>
<td>646</td>
<td>0.60</td>
<td>0.61</td>
<td>0.62</td>
<td>0.62</td>
<td>0.61</td>
</tr>
<tr>
<td>4th</td>
<td>642</td>
<td>0.61</td>
<td>0.60</td>
<td>0.59</td>
<td>0.59</td>
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</tr>
<tr>
<td>5th</td>
<td>626</td>
<td>0.64</td>
<td>0.55</td>
<td>0.52</td>
<td>0.52</td>
<td>0.54</td>
</tr>
<tr>
<td>MCR Percent:</td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
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<tr>
<td>0–24</td>
<td>410</td>
<td>0.42</td>
<td>0.40</td>
<td>0.40</td>
<td>0.39</td>
<td>0.39</td>
</tr>
<tr>
<td>25–49</td>
<td>2,081</td>
<td>0.63</td>
<td>0.63</td>
<td>0.63</td>
<td>0.63</td>
<td>0.63</td>
</tr>
<tr>
<td>50 and over</td>
<td>590</td>
<td>0.72</td>
<td>0.73</td>
<td>0.74</td>
<td>0.74</td>
<td>0.74</td>
</tr>
<tr>
<td>Region:</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>New England</td>
<td>130</td>
<td>0.68</td>
<td>0.64</td>
<td>0.63</td>
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</tr>
<tr>
<td>Middle Atlantic</td>
<td>354</td>
<td>0.86</td>
<td>0.83</td>
<td>0.83</td>
<td>0.83</td>
<td>0.83</td>
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<tr>
<td>South Atlantic</td>
<td>512</td>
<td>0.74</td>
<td>0.76</td>
<td>0.78</td>
<td>0.78</td>
<td>0.77</td>
</tr>
<tr>
<td>East North Central</td>
<td>482</td>
<td>0.63</td>
<td>0.63</td>
<td>0.63</td>
<td>0.63</td>
<td>0.63</td>
</tr>
<tr>
<td>East South Central</td>
<td>290</td>
<td>0.76</td>
<td>0.79</td>
<td>0.80</td>
<td>0.80</td>
<td>0.79</td>
</tr>
<tr>
<td>West North Central</td>
<td>252</td>
<td>0.39</td>
<td>0.41</td>
<td>0.41</td>
<td>0.41</td>
<td>0.41</td>
</tr>
<tr>
<td>West South Central</td>
<td>487</td>
<td>0.46</td>
<td>0.48</td>
<td>0.48</td>
<td>0.48</td>
<td>0.47</td>
</tr>
<tr>
<td>Mountain</td>
<td>223</td>
<td>0.36</td>
<td>0.39</td>
<td>0.40</td>
<td>0.40</td>
<td>0.39</td>
</tr>
</tbody>
</table>
We are inviting public comment on our preferred proposal and alternative considerations.

11. Accounting for Social Risk Factors in the Hospital Readmissions Reduction Program

We understand that social risk factors such as income, education, race and ethnicity, employment, disability, community resources, and social support (certain factors of which are also sometimes referred to as socioeconomic status (SES) factors or socio-demographic status (SDS) factors) play a major role in health. One of our core objectives is to improve beneficiary outcomes, including reducing health disparities, and we want to ensure that all beneficiaries, including those with social risk factors, receive high quality care. In addition, we seek to ensure that the quality of care furnished by providers and suppliers is assessed as fairly as possible under our programs, while ensuring that beneficiaries have adequate access to excellent care.

We have been reviewing reports prepared by the Office of the Assistant Secretary for Planning and Evaluation (ASPE) and the National Academies of Sciences, Engineering, and Medicine on the issue of accounting for social risk factors in CMS’ value-based purchasing programs and quality reporting programs, and considering options on how to address the issue in these programs. On December 21, 2016, ASPE submitted a report to Congress on a study it was required to conduct under section 2(d) of the Improving Medicare Post-Acute Care Transformation (IMPACT) Act of 2014. The study analyzed the effects of certain social risk factors in Medicare beneficiaries on quality measures and measures of resource use used in one or more of nine Medicare value-based purchasing programs, including the Hospital Readmissions Reduction Program.42

The report also included considerations for strategies to account for social risk factors in these programs. In a January 10, 2017 report released by the National Academies of Sciences, Engineering, and Medicine, that body provided various potential methods for measuring and accounting for social risk factors, including stratified public reporting.42

As noted in the FY 2017 IPPS/LTCH PPS final rule, the NQF has undertaken a 2-year trial period in which certain new measures, measures undergoing maintenance review, and measures endorsed with the condition that they enter the trial period can be assessed to determine whether risk adjustment for selected social risk factors is appropriate for these measures. This trial entails temporarily allowing inclusion of social risk factors in the risk-adjustment approach for these measures. At the conclusion of the trial, NQF will issue recommendations on the future inclusion of social risk factors in risk adjustment for these quality measures, and we will closely review its findings.

As we continue to consider the analyses and recommendations from these reports and await the results of the NQF trial on risk adjustment for quality measures, we are continuing to work with stakeholders in this process. As we have previously communicated, we are concerned about holding providers to different standards for the outcomes of their patients with social risk factors because we do not want to mask potential disparities or minimize incentives to improve the outcomes for disadvantaged populations. Keeping this concern in mind, while we sought input on this topic previously, we continue to seek public comment on whether we should account for social risk factors in the Hospital Readmissions Reduction Program and, if so, what method or combination of methods would be most appropriate for accounting for social risk factors. Examples of methods include: confidential reporting of stratified measure rates to providers; public reporting of stratified measure rates; risk adjustment of a particular measure as appropriate based on data and evidence; developing readmission measures or statistical approaches that are suitable for the reporting of performance on readmissions; providing financial incentives for achievement of low readmission rates for beneficiaries with social risk factors; and using a hospital-wide readmissions measure. While we consider whether and to what extent we currently have statutory authority to implement one or more of the above-described methods, we are seeking comments on whether any of these methods should be considered, and if so, which of these methods or combination of methods would best account for social risk factors in the Hospital Readmissions Reduction Program.

In addition, we are also seeking public comment on which social risk factors might be most appropriate for stratifying measure scores and/or potential risk adjustment of a particular


Source: FY 2017 Hospital Readmissions Reduction Program Final Rule Results. Results are based on July 1, 2012, through June 30, 2015, discharges among subsection (d) and Maryland hospitals only. Although data from all subsections (d) and Maryland hospitals are used in calculations of each hospital’s Excess Readmission Ratio (ERR), this table does not include results for Maryland hospitals. This table only includes results for hospitals who are eligible for a penalty under the program on the basis of having at least 25 eligible discharges for at least one measure. Hospital Characteristics are based on the FY 2017 final rule Impact File. There were 15 hospitals that did not have MCR percentages in the FY 2017 final rule Impact File. To calculate the payment adjustment as a proportion of total base operating DRG payments, this analysis used MedPAR data to calculate the total base operating DRG payments from July 1, 2014 through June 30, 2015. The group average share of payment adjustments as a percentage of all DRG payments is calculated as the sum of all Medicare savings for the group of hospitals divided by total base operating DRG payments for all hospitals in that group.

<table>
<thead>
<tr>
<th>Hospital characteristics</th>
<th>Number of hospitals with characteristic</th>
<th>Current methodology (percent)</th>
<th>Median plus neutrality modifier (neutrality modifier = 0.9546) (percent)</th>
<th>Mean plus neutrality modifier (neutrality modifier = 1.0135) (percent)</th>
<th>Budget neutralizing ERR (percent)</th>
<th>Standardized ERR plus neutrality modifier (neutrality modifier = 0.9710) (percent)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pacific</td>
<td>366</td>
<td>0.42</td>
<td>0.37</td>
<td>0.34</td>
<td>0.34</td>
<td>0.36</td>
</tr>
</tbody>
</table>
measure. Examples of social risk factors include, but are not limited to, dual eligibility/low-income subsidy, race and ethnicity, and geographic area of residence. We are seeking comments on which of these factors, including current data sources where this information would be available, could be used alone or in combination, and whether other data should be collected to better capture the effects of social risk. We will take commenters’ input into consideration as we continue to assess the appropriateness and feasibility of accounting for social risk factors in the Hospital Readmissions Reduction Program. We note that any such changes would be proposed through future notice-and-comment rulemaking.

We look forward to working with stakeholders as we consider the issue of accounting for social risk factors and reducing health disparities in CMS programs. Of note, implementing any of the above methods would be taken into consideration in the context of how this and other CMS programs operate (for example, data submission methods, availability of data, statistical considerations relating to reliability of data calculations, among others), so we also welcome comment on operational considerations. CMS is committed to ensuring that its beneficiaries have access to and receive excellent care, and that the quality of care furnished by providers and suppliers is assessed fairly in CMS programs.

12. Extraordinary Circumstance Exception (ECE) Policy

Many of our quality reporting and value-based purchasing programs share a common process for requesting an exception from program reporting due to an extraordinary circumstance not within a provider’s control. The Hospital IQR, Hospital OQR, IPFQR, Ambulatory Surgical Center Quality Reporting (ASCQR), PCHQR Programs, as well as the HAC Reduction Program, and the Hospital Readmissions Reduction Program, share common processes for ECE requests. In reviewing the policies for these programs, we recognized that there are five areas in which these programs have variance regarding ECE requests. These are: (1) Allowing the facilities or hospitals to submit a form signed by the facility’s or hospital’s CEO versus CEO or designated personnel; (2) requiring the form be submitted within 30 days following the date that the extraordinary circumstance occurred versus within 90 days following the date the extraordinary circumstance occurred; (3) inconsistency regarding specification of a timeline for us to provide our formal response notifying the facility or hospital of our decision; (4) inconsistency regarding specification of our authority to grant ECEs due to CMS data system issues; and (5) referring to the program as “extraordinary extensions/exemptions” versus as “extraordinary circumstances exceptions.” We believe addressing these five areas, as appropriate, can improve administrative efficiencies for affected facilities or hospitals.

In the FY 2016 IPPS/LTCH PPS final rule (80 FR 40542 through 40543), we adopted an ECE policy for the Hospital Readmissions Reduction Program beginning in FY 2016. This policy was similar to the ECE policy for the Hospital IQR Program, as finalized in the FY 2012 IPPS/LTCH PPS final rule (76 FR 51651), modified in the FY 2014 IPPS/LTCH PPS final rule (78 FR 50836) (designation of a non-CEO hospital contact), and further modified in the FY 2015 IPPS/LTCH PPS final rule (79 FR 50277) (amended 42 CFR 412.140(c)(2) to refer to “extension or exemption” instead of the former “extension or waiver”).

We are proposing to update these policies by: (1) Allowing the facility to submit a form signed by the facility’s CEO or designated personnel; (2) clarifying that we will strive to provide our formal response notifying the facility of our decision within 90 days of receipt of the facility’s request; and (3) allowing CMS to have the authority to grant ECEs due to CMS data system issues which affect data submission. These proposed policies generally align with policies in the Hospital IQR Program (76 FR 51651 through 51652), (78 FR 50836 through 50837) and (81 FR 57181 through 57182), Hospital OQR Program (77 FR 66849 and 81 FR 79795), as well as other quality reporting programs. We are proposing that these policies would apply beginning in FY 2018 as related to extraordinary circumstances that occur on or after October 1, 2017.

We note that there may be circumstances in which it is not feasible for a facility’s CEO to sign the ECE request form. In these circumstances, we believe that facilities affected by such circumstances should be able to submit ECE forms regardless of the CEO’s availability to sign. This proposed change would allow hospitals to designate an appropriate, non-CEO, contact at its discretion. This individual would be responsible for the submission, and would be the one signing the form. Therefore, we are proposing to accept ECE forms which have been signed by designated personnel.

We also believe that it is important for facilities to receive timely feedback regarding the status of ECE requests. We strive to complete our review of each ECE request as quickly as possible. However, we recognize that the number of requests we receive, and the complexity of the information provided impacts the actual timeframe to make ECE determinations. To improve transparency of our process, we believe it is appropriate to clarify that we will strive to complete our review of each request within 90 days of receipt.

Although we do not anticipate this situation will happen on a regular basis, there may be times where CMS experiences issues with its data systems that directly affects facilities’ abilities to submit data. In these cases, we believe it would be inequitable to require facilities to report. Therefore, we are proposing to allow CMS to grant ECEs to facilities if we determine that a systemic problem with one of our data collection systems directly affected the ability of the facilities to submit data. If we make the determination to grant ECEs, we are proposing to communicate this decision through routine communication channels.

We are inviting public comment on these proposed modifications to the Extraordinary Circumstance Exception policy.

13. Timeline for Public Reporting of Excess Readmission Ratios on Hospital Compare for the FY 2018 Payment Determination

Section 1886(q)(6) of the Act requires the Secretary to make information available to the public regarding readmission rates of each subsection (d) hospital under the program, and states that such information shall be posted on the Hospital Compare Internet Web site in an easily understandable format. Accordingly, in the FY 2013 IPPS/LTCH PPS final rule (77 FR 53401), we indicated that public reporting for excess readmission ratios could be available on the Hospital Compare Web site as early as mid-October. In the FY 2017 IPPS/LTCH PPS final rule (81 FR 56978 through 56979), we clarified that public reporting of excess readmission ratios will be posted on an annual basis to the Hospital Compare Web site as soon as is feasible following the review period. This may occur as early as October, but it could occur later for a particular year in order to streamline reporting and align with other hospital quality reporting and performance programs.
Section 1886(o) of the Act, as added by section 3001(a)(1) of the Affordable Care Act, requires the Secretary to establish a hospital value-based purchasing program (the Hospital VBP Program) under which value-based incentive payments are made in a fiscal year (FY) to hospitals that meet performance standards established for a performance period for such fiscal year. Both the performance standards and the performance period for a fiscal year are to be established by the Secretary. For more of the statutory background and descriptions of our current policies for the Hospital VBP Program, we refer readers to the Hospital Inpatient VBP Program final rule (76 FR 26490 through 26547); the FY 2012 IPPS/LTCH PPS final rule (76 FR 51653 through 51660); the CY 2013 OPPS/ASC final rule with comment period (76 FR 74527 through 74547); the FY 2013 IPPS/LTCH PPS final rule (77 FR 53567 through 53614); the CY 2014 OPPS/ASC final rule (78 FR 50676 through 50707); the CY 2014 OPPS/ASC final rule (78 FR 75120 through 75121); the CY 2015 IPPS/LTCH PPS final rule (79 FR 50048 through 50087); the FY 2016 IPPS/LTCH PPS final rule with comment period (80 FR 49544 through 49570); the FY 2017 IPPS/LTCH PPS final rule (81 FR 56979 through 57011); and the CY 2017 OPPS/ASC final rule with comment period (81 FR 79855 through 79862).

We have also codified certain requirements for the Hospital VBP Program at 42 CFR 412.160 through 412.167.

b. FY 2018 Program Year Payment Details

Section 1886(o)(7)(B) of the Act instructs the Secretary to reduce the base operating DRG payment amount for a hospital for each discharge in a fiscal year by an applicable percent. Under section 1886(o)(7)(A) of the Act, the sum total of these reductions in a fiscal year must equal the total amount available for value-based incentive payments for all eligible hospitals for the fiscal year, as estimated by the Secretary. We finalized details on how we would implement these provisions in the FY 2013 IPPS/LTCH PPS final rule (77 FR 53571 through 53573) and refer readers to that rule for further details.

Under section 1886(o)(7)(C)(iv) of the Act, the applicable percent for the FY 2018 program year is 2.00 percent. Using the methodology we adopted in the FY 2013 IPPS/LTCH PPS final rule (77 FR 53571 through 53573), we estimate that the total amount available for value-based incentive payments for FY 2018 is approximately $1.9 billion, based on the December 2016 update of the FY 2016 MedPAR file. We intend to update this estimate for the FY 2018 IPPS/LTCH PPS final rule using the March 2017 update of the FY 2016 MedPAR file.

As finalized in the FY 2013 IPPS/LTCH PPS final rule (77 FR 53573 through 53576), we will utilize a linear exchange function to translate this estimated amount available into a value-based incentive payment percentage for each hospital, based on its Total Performance Score (TPS). We will then calculate a value-based incentive payment adjustment factor that will be applied to the base operating DRG payment amount for each discharge occurring in FY 2018, on a per-claim basis. We are publishing proxy value-based incentive payment adjustment factors in Table 16 associated with this proposed rule (which is available via the Internet on the CMS Web site). The proxy factors are based on the TPS from the FY 2017 program year. These FY 2017 performance scores are the most recently available performance scores hospitals have been given the opportunity to review and correct. The slope of the linear exchange function used to calculate those proxy value-based incentive payment adjustment factors is 3.0692781725. This slope, along with the estimated amount available for value-based incentive payments, is also published in Table 16.

We intend to update this table as Table 16A in the final rule (which will be available on the CMS Web site) to reflect changes based on the March 2017 update to the FY 2016 MedPAR file. We also intend to update the slope of the linear exchange function used to calculate those updated proxy value-based incentive payment adjustment factors. The updated proxy value-based incentive payment adjustment factors for FY 2018 will continue to be based on historic FY 2017 program year TPSs because hospitals will not have been given the opportunity to review and correct their actual TPSs for the FY 2018 program year until after the FY 2018 IPPS/LTCH PPS final rule is published.

After hospitals have been given an opportunity to review and correct their actual TPSs for FY 2018, we will add Table 16B (which will be available via the Internet on the CMS Web site) to display the actual value-based incentive payment adjustment factors, exchange function slope, and estimated amount available for the FY 2018 program year. We expect Table 16B will be posted on the CMS Web site in the fall of 2017.

2. Accounting for Social Risk Factors in the Hospital VBP Program

We understand that social risk factors such as income, education, race and ethnicity, employment, disability, community resources, and social support (certain factors of which are also sometimes referred to as socioeconomic status (SES) factors or socio-demographic status (SDS) factors) play a major role in health. One of our core objectives is to improve beneficiary outcomes, including reducing health disparities, and we want to ensure that all beneficiaries, including those with social risk factors, receive high quality care. In addition, we seek to ensure that the quality of care furnished by providers and suppliers is assessed as fairly as possible under our programs while ensuring that beneficiaries have adequate access to excellent care.

We have been reviewing reports prepared by the Office of the Assistant Secretary for Planning and Evaluation (ASPE)43 and the National Academies of Sciences, Engineering, and Medicine on the issue of accounting for social risk factors in CMS’ value-based purchasing and quality reporting programs, and considering options on how to address the issue in these programs. On December 21, 2016, ASPE submitted a Report to Congress on a study it was required to conduct under section 2(d) of the Improving Medicare Post-Acute Care Transformation (IMPACT) Act of 2014. The study analyzed the effects of certain social risk factors in Medicare beneficiaries on quality measures and measures of resource use used in one or more of nine Medicare value-based purchasing programs, including the Hospital VBP Program.44 The report also included considerations for strategies to account for social risk factors in these programs. In a January 10, 2017 report released by the National Academies of Sciences, Engineering, and Medicine, that body provided various potential methods for measuring and accounting for social risk factors, including stratified public reporting.45

In the ASPE report noted above, there is an analysis of and focus on the Medicare Spending Per Beneficiary (MSPB) measure, which was adopted by the Hospital VBP Program beginning with the FY 2015 program year.46 We note that the MSPB measure is currently undergoing endorsement review for NQF, as part of the 2-year socioeconomic trial period described below.47 ASPE’s December 2016 Report to Congress did not include an analysis of the effect of social risk factors on hospital performance on any condition-specific payment measures that are currently adopted for the Hospital VBP Program. As noted in the FY 2017 IPPS/LTC PPS final rule, the MSPB has undertaken a 2-year trial period in which certain new measures, measures undergoing maintenance review, and measures endorsed with the condition that they enter the trial period can be assessed to determine whether risk adjustment for selected social risk factors is appropriate for these measures. This trial entails temporarily allowing inclusion of social risk factors in the risk-adjustment approach for these measures. At the conclusion of the trial, NQF will issue recommendations on the future inclusion of social risk factors in risk adjustment for these quality measures, and we will closely review its findings.

We note that the AMI Payment and HF Payment measures adopted in the FY 2017 IPPS/LTC PPS final rule (81 FR 56986 through 56990 and 81 FR 56990 through 56992, respectively) were included in the NQF’s trial. Based on its review of these measures during the trial, the NQF re-endorsed these measures without modifications to their risk adjustment methodologies for social risk factors. We are proposing to adopt the AMI Payment measure beginning with the FY 2022 program year for the Hospital VBP Program and, if so, what method or combination of methods would be most appropriate for accounting for social risk factors because we do not want to mask potential disparities or minimize incentives to improve the outcomes for disadvantaged populations. Keeping this concern in mind, while we sought input on this topic previously, we continue to seek public comment on whether we should account for social risk factors in the Hospital VBP Program, and if so, what method or combination of methods would be most appropriate for accounting for social risk factors. Examples of methods include: adjustment of the payment adjustment methodology under the Hospital VBP Program; adjustment of provider provider performance scores for instances, stratifying providers based on the proportion of their patients who are dual eligible); confidential reporting of stratified measure rates to providers; public reporting of stratified measure rates; risk adjustment of a particular measure based on data and evidence; and redesigning payment incentives (for instance, rewarding improvement for providers caring for patients with social risk factors or incentivizing providers to achieve health equity.

We note that in section V.I.9 of the preamble of this rule, we discuss considerations for stratifying hospitals in peer groups for purposes of assessing payment adjustments under the Hospital Readmissions Reduction Program, as required by the 21st Century Cures Act. We refer readers to that section for a detailed discussion of these alternatives; while this discussion and corresponding proposal are specific to the Hospital Readmissions Reduction Program, they reflect the level of analysis we would undertake when evaluating methods and combinations of methods for accounting for social risk factors in CMS’ other value-based purchasing programs, such as the Hospital VBP Program. While we consider whether and to what extent we currently have statutory authority to implement one or more of the above-described methods, we are seeking comments on whether any of these methods should be considered, and if so, which of these methods or combination of methods would best account for social risk factors in the Hospital VBP Program.

In addition, we are also seeking public comment on which social risk factors might be most appropriate for stratifying measure scores and/or potential risk adjustment of a particular measure. Examples of social risk factors include, but are not limited to, dual eligibility/low-income subsidy, race and ethnicity, and geographic area of residence. We are seeking comments on which of these factors, including current data sources where this information would be available, could be used alone or in combination, and whether other data should be collected to better capture the effects of social risk. We will take commenters’ input into consideration as we continue to assess the appropriateness and feasibility of accounting for social risk factors in the Hospital VBP Program. We note that any such changes would be proposed through future notice-and-comment rulemaking.

We look forward to working with stakeholders as we consider the issue of accounting for social risk factors and reducing health disparities in CMS programs. Of note, implementing any of the above methods would be taken into consideration in the context of how this and other CMS programs operate (for example, data submission methods, availability of data, statistical considerations relating to reliability of data calculations, among others), we also welcome comment on operational considerations. CMS is committed to ensuring that its beneficiaries have access to and receive excellent care, and that the quality of care furnished by providers and suppliers is assessed fairly in CMS programs.
3. Retention and Removal of Quality Measures for the FY 2019 Program Year
   a. Retention of Previously Adopted Hospital VBP Program Measures

   In the FY 2013 IPPS/LTCH PPS final rule (77 FR 53592), we finalized a policy to retain measures from prior program years for each successive program year, unless otherwise proposed and finalized. We are not proposing any changes to this policy.

   b. Proposed Removal of the PSI 90 Measure

   In the FY 2017 IPPS/LTCH PPS final rule (81 FR 56979 through 56981), we finalized our proposal to shorten the performance period for the current PSI 90 measure for the FY 2018 program year due to concerns associated with combining measure performance data that use both ICD–9 and ICD–10 data in calculating performance scores under the measure. In that final rule, we explained our system requires an ICD–10 risk-adjusted version of the AHRQ PSI software in order to calculate scores using ICD–10 codes, and AHRQ needs a full year of nationally representative ICD–10 coded data before it can complete development of risk-adjusted models based on a national reference population for this software. This means the AHRQ PSI software will not be available for us to calculate scores until late CY 2017. More importantly, we noted an ICD–10 version of the current PSI 90 measure is not being developed (81 FR 56980), nor will ICD–10 AHRQ QI software be available to calculate performance scores for the FY 2019 program year (81 FR 56981). As a result, we will not be able to calculate performance scores for the current PSI 90 measure for the FY 2019 program year because these scores would include ICD–10 data. Based on these concerns, in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56981), we signaled our intent to propose to remove the current PSI 90 measure from the Hospital VBP Program beginning with the FY 2019 program year. We are now proposing in this FY 2018 IPPS/LTCH PPS proposed rule to remove the current PSI 90 measure from the Hospital VBP Program beginning with the FY 2019 program year.

   We are inviting public comment on this proposal. We also refer readers to section V.J.4.b. of the preamble of this proposed rule where we are proposing to adopt the modified version of the PSI 90 measure for the Hospital VBP Program beginning with the FY 2023 program year.

   In summary, for the FY 2019 and FY 2020 program years, we have finalized the following measure set and are proposing to remove the current PSI 90 measure, as indicated:

PREVIOUSLY ADOPTED MEASURES AND PROPOSED MEASURE FOR REMOVAL FOR THE FY 2019 AND FY 2020 PROGRAM YEARS

<table>
<thead>
<tr>
<th>Measure short name</th>
<th>Domain/measure name</th>
<th>NQF #</th>
</tr>
</thead>
<tbody>
<tr>
<td>HCAHPS</td>
<td>Hospital Consumer Assessment of Healthcare Providers and Systems (HCAHPS)** (including Care Transition Measure).</td>
<td>0166 (0228)</td>
</tr>
<tr>
<td>MORT–30–AMI</td>
<td>Hospital 30-Day, All-Cause, Risk-Standardized Mortality Rate (RSMR) Following Acute Myocardial Infarction (AMI) Hospitalization.</td>
<td>0230</td>
</tr>
<tr>
<td>MORT–30–HF</td>
<td>Hospital 30-Day, All-Cause, Risk-Standardized Mortality Rate (RSMR) Following Heart Failure (HF) Hospitalization.</td>
<td>0229</td>
</tr>
<tr>
<td>MORT–30–PN</td>
<td>Hospital 30-Day, All-Cause, Risk-Standardized Mortality Rate (RSMR) Following Pneumonia Hospitalization.</td>
<td>0468</td>
</tr>
<tr>
<td>THA/TKA</td>
<td>Hospital-Level Risk-Standardized Complication Rate (RSCR) Following Elective Primary Total Hip Arthroplasty (THA) and/or Total Knee Arthroplasty (TKA).</td>
<td>1550</td>
</tr>
<tr>
<td>CAUTI</td>
<td>National Healthcare Safety Network (NHSN) Catheter-Associated Urinary Tract Infection (CAUTI) Outcome Measure.</td>
<td>0138</td>
</tr>
<tr>
<td>CLABSI</td>
<td>National Healthcare Safety Network (NHSN) Central Line-Associated Bloodstream Infection (CLABSI) Outcome Measure.</td>
<td>0139</td>
</tr>
<tr>
<td>Colon and Abdominal Hysterectomy SSI.</td>
<td>American College of Surgeons—Centers for Disease Control and Prevention (ACS–CDC) Harmonized Procedure Specific Surgical Site Infection (SSI) Outcome Measure.</td>
<td>0753</td>
</tr>
<tr>
<td>MRSA Bacteremia</td>
<td>National Healthcare Safety Network (NHSN) Facility-wide Inpatient Hospital-onset Methicillin-resistant Staphylococcus aureus (MRSA) Bacteremia Outcome Measure.</td>
<td>1716</td>
</tr>
<tr>
<td>CDI</td>
<td>National Healthcare Safety Network (NHSN) Facility-wide Inpatient Hospital-onset Clostridium difficile Infection (CDI) Outcome Measure.</td>
<td>1717</td>
</tr>
<tr>
<td>PSI 90***</td>
<td>Patient Safety for Selected Indicators (Composite Measure)</td>
<td>0531</td>
</tr>
<tr>
<td>PC–01</td>
<td>Elective Delivery</td>
<td>0469</td>
</tr>
</tbody>
</table>

The “current” PSI 90 measure refers to the version of the PSI 90 measure previously finalized for use in the Hospital VBP Program in the FY 2013 IPPS/LTCH PPS final rule (78 FR 50694).

The AHRQ QI Software is the software used to calculate PSIs and the composite measure. More information is available at: http://www.qualityindicators.ahrq.gov/Downloads/
4. Proposed New Measures for the FY 2022 Program Year, FY 2023 Program Year, and Subsequent Years

We consider measures for adoption based on the statutory requirements, including specification under the Hospital IQR Program, posting dates on the Hospital Compare Web site, and our priorities for quality improvement as outlined in the current CMS Quality Strategy, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/QualityInits/HospitalQualityInitiatives.html. Due to the time necessary to adopt measures, we often adopt policies for the Hospital VBP Program well in advance of the program year for which they will be applicable.

a. Proposed New Measure for the FY 2022 Program Year and Subsequent Years: Hospital-Level, Risk-Standardized Payment Associated With a 30-Day Episode-of-Care for Pneumonia (PN Payment)

(1) Measure Proposal

Hospital-Level, Risk-Standardized Payment Associated with a 30-Day Episode-of-Care for Pneumonia (PN Payment) is a measure assessing hospital risk-standardized payment associated with a 30-day episode-of-care for pneumonia. We adopted this measure in the Hospital IQR Program in the FY 2015 IPPS/LTCH PPS final rule (79 FR 50227 through 50231), and we adopted an updated version of the measure, with an expanded cohort and modified risk-adjustment model, in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57125 through 57128). For purposes of describing this measure, the “cohort” is the set of hospitalizations, or “index admissions,” that meet all of the measure’s inclusion and exclusion criteria and, thus, are used to calculate the total payments Medicare makes on behalf of these Medicare beneficiaries for a 30-day episode-of-care. The cohort for the expanded version of the PN Payment measure includes Medicare FFS patients aged 65 or older with: (1) A principal hospital discharge diagnosis of pneumonia, including not only viral or bacterial pneumonia but also aspiration pneumonia; or (2) a principal discharge diagnosis of sepsis (but not severe sepsis) with a secondary diagnosis of pneumonia (including viral or bacterial pneumonia and aspiration pneumonia) coded as present on admission. The measure calculates payments for these patients over a 30-day episode-of-care, beginning with the index admission, using administrative claims data. In general, the measure uses the same approach to risk-adjustment as 30-day outcome measures previously adopted for the Hospital VBP Program, including the 30-day PN mortality measure, MORT–30–PN. Initial measure data collected under the Hospital IQR Program for the expanded PN Payment cohort and modified risk-adjustment model will be posted on Hospital Compare in July 2017, and the full measure specifications are available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInitiatives/HospitalQualityInits/Measure-Methodology.html.

Promoting high-value care is an essential part of our mission to provide better health care for individuals, better health for populations, and lower costs for health care. Our aim is to encourage a higher value care where there is the greatest number of patients to benefit from improvements, and the largest sample size to ensure reliability. Pneumonia is one of the leading causes of hospitalization for Americans aged 65 and over, and pneumonia patients incur roughly $10 billion in aggregate health care costs. There is evidence of variation in payments at hospitals for pneumonia patients in the proposed PN Payment measure; median 30-day risk-adjusted payment among Medicare FFS patients aged 65 or older hospitalized for pneumonia was $15,988 and ranged from $9,193 to $26,546 for the July 2011 through June 2014 reporting period in the Hospital IQR Program. This variation in payment suggests there is opportunity for improvement. We believe it is important to adopt the PN Payment measure for the Hospital VBP Program because variation in payment may reflect differences in care decision-making and resource utilization (for example, treatment, supplies, or services) for patients with pneumonia both during hospitalization and immediately post-discharge. The PN Payment measure specifically addresses the NQS priority and CMS Quality Strategy goal to make quality care more affordable.

We recognize high or low payments to hospitals are difficult to interpret in isolation. Some high payment hospitals may produce better clinical outcomes when compared with low payment hospitals, while other high payment hospitals may not produce better outcomes. For this reason, payment measure results viewed in isolation are not necessarily an indication of quality. However, by viewing such information along with quality measure results, consumers, payers, and providers would be able to better assess the value of care. In order to incentivize innovation that promotes high-quality care at high value, it is important to examine measures of payment and patient outcomes concurrently. The proposed PN Payment measure is intended to be paired with the MORT–30–PN measure in the Hospital VBP Program, thereby


52 The Hospital VBP Program first adopted the MORT–30–PN measure for the FY 2014 program year in the Hospital Inpatient Value-Based Purchasing Program final rule (76 FR 26497 through
directly linking payment to quality by the alignment of comparable populations and risk-adjustment methodologies to facilitate the assessment of efficiency and value of care. We believe adopting the PN Payment measure will create stronger incentives for appropriately reducing practice pattern variation to achieve the aim of lowering the cost of care and creating better coordinated care for Medicare beneficiaries.

We are proposing to adopt the PN Payment measure beginning with the FY 2022 program year. The PN Payment measure would be added to the Efficiency and Cost Reduction domain. The proposed measure fulfills all of the statutory requirements for the Hospital VBP Program based on our adoption of the measure in the Hospital IQR Program, and our anticipated posting of measure data for the refined PN Payment measure, with the expanded cohort and modified risk-adjustment model, on Hospital Compare beginning July 2017, which would be at least one year before the beginning of the proposed performance period of August 1, 2018. We refer readers to sections V.J.5.c.(3) through V.J.5.c.(5) of the preamble of this proposed rule where we are proposing baseline periods and performance periods for this measure if adopted for the Hospital VBP Program.

The proposed PN Payment measure (MUC15–378) was reviewed by the MAP in December 2015 and did not receive support for adoption into the Hospital VBP Program.53 The result of the MAP vote was 31 percent support, 15 percent conditional support, and 54 percent do not support. The MAP’s decision of “do not support” for the proposed PN Payment measure was based on concerns that the measure may overlap with and thereby double count services that are already captured in the MSPB measure. In addition, some MAP members expressed a desire to have more experience with the measure in the Hospital IQR Program to understand whether there may be unintended consequences or a need to adjust for social risk factors. We note some MAP members expressed support for the proposed PN Payment measure and other condition-specific payment measures, expressing that the increased granularity provided by condition-specific payment measures will provide valuable feedback to hospitals for targeted improvement.

With respect to MAP stakeholder concerns that treatment- or condition-specific payment measures may overlap and double count services, we note that the proposed PN Payment measure addresses a topic of critical importance to quality improvement in the inpatient hospital setting. As discussed above, we selected the PN Payment measure because we believe it is appropriate to provide stronger incentives for hospitals to provide high-value and efficient care, especially for a high-volume condition such as pneumonia. We acknowledge that hospitals that do not perform well on the PN Payment measure may also perform poorly on the MSPB measure and potentially receive a lower incentive payment, depending upon their performance on other measures. However, because admissions for pneumonia make up only a part of all admissions included in the MSPB measure, a hospital’s results on the MSPB measure may not be the same as their result on the PN Payment measure. In other words, a hospital’s results for one measure are not deterministic of its results of the other, so we cannot state conclusively that if a hospital performs well (or poorly) on one measure, that they will also perform well (or poorly) on the second measure. Hospitals would perform differently on the MSPB and PN Payment measures because their measures evaluate performance on different metrics. For example, some hospitals with poorer results on the MSPB measure may have better results on the PN Payment measure allowing them to improve their overall score. In addition, the overlap between the MSPB and PN Payment measures may result in some hospitals receiving an increased benefit by performing well on both measures. Furthermore, if a hospital does not perform as well on the MSPB measure relative to other hospitals but performs very well with respect to its pneumonia patients on the proposed PN Payment measure, that hospital would have the opportunity to earn a higher score in the Efficiency and Cost Reduction domain.

Regarding MAP stakeholder concerns for the need to adjust for social risk factors, we note the proposed PN Payment measure already incorporates a risk-adjustment methodology that accounts for age and comorbidities. We understand the important role social risk factors play in the care of patients, routinely monitor the impact of social risk factors on hospitals’ results on our measures, and will continue to do so. In addition, as discussed in section V.J.3. of the preamble of this proposed rule, the original PN Payment measure using the previous measure cohort (Hospital-level, risk-standardized payment associated with a 30-day episode-of-care for pneumonia (NQF #2579)), as well as the AMI Payment and HF Payment measures adopted in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56987 through 56990 and 81 FR 56990 through 56992, respectively), which use the same measurement methodology as the proposed PN Payment measure, recently underwent successful NQF re-endorsement following enrollment in the NQF’s trial. The NQF re-endorsed these measures without requesting modifications to their risk adjustment methodologies for adjustment by social risk factors. The proposed PN Payment measure includes an updated risk-adjustment model that accounts for patient comorbidities, and we intend to submit to NQF that risk adjustment model as part of the overall proposed PN Payment measure specifications during the next Cost and Resource Use project.

As noted above, some MAP members expressed support for the proposed PN Payment measure and other condition-specific payment measures, agreeing the increased granularity provided by condition-specific payment measures will provide valuable feedback to hospitals for targeted improvement. In addition, a NQF-commissioned white paper also supports the position that cost or payment measures should be interpreted in the context of quality measures and that measures which link cost and quality are the preferred method of assessing hospital efficiency.54 The PN Payment measure, which directly pairs with the MORT–30–PN measure in the Hospital VBP Program, follows this recommended approach. Based on our analysis of the issues surrounding condition-specific payment measures, we believe the benefits of adopting the PN Payment measure outweigh any potential risks; however, we also remain committed to monitoring for unintended consequences.

We are inviting public comment on this proposal.

(2) Proposed Scoring Methodology for the PN Payment Measure

We are proposing to calculate the PN Payment measure using the same

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28511. We subsequently expanded the measure cohort beginning with the FY 2021 program year in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56994 through 56996).


methodology we use to score the MSPB measure and, as finalized in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56992 through 56993), the AMI Payment and HF Payment measures so that all measures in the Efficiency and Cost Reduction domain are scored in the same manner. We note for these measures that lower values represent better performance.

For achievement points, we are proposing to calculate a spending ratio of PN spending for each hospital to the median PN spending across all hospitals during the performance period. We would then use each hospital’s PN spending ratio to calculate between 0 and 10 achievement points. We are proposing to set the achievement threshold at the median PN spending ratio across all hospitals during the performance period. Because lower values represent better performance under the proposed PN Payment measure, we are proposing to set the benchmark at the mean of the lowest decile of the PN spending ratios during the performance period. Therefore, if a hospital’s individual PN spending ratio falls above the achievement threshold, the hospital would score 0 achievement points on the measure. If a hospital’s individual PN spending ratio falls at or below the benchmark, the hospital would score the maximum 10 achievement points on the measure. If a hospital’s individual PN spending ratio falls at or below the achievement threshold but above the benchmark, the hospital would score between 1 and 9 points according to the following formula:

\[ (10 \times \left( \frac{\text{Hospital baseline period ratio}}{\text{Hospital performance period ratio}} \right) - 1) \times 0.5 \]

For improvement points, we are proposing to calculate a spending ratio of PN spending for each hospital to the median PN spending across all hospitals during the performance period. We would then use each hospital’s PN spending ratio to calculate between 0 and 9 improvement points on the measure. If a hospital’s score on the measure during the performance period is less than its baseline period score but above the benchmark, the hospital would receive a score of 0 to 9 according to the following formula:

\[ (10 \times \left( \frac{\text{Hospital baseline period ratio} - \text{Hospital performance period ratio}}{(\text{Hospital baseline period ratio} - \text{benchmark})} \right) - 0.5 \]

We note that if a hospital scores at or below the benchmark on the achievement scoring methodology, that hospital will receive the maximum 10 points for this measure. As a result, the hospital would not receive an improvement score for this measure.

For more information about the proposed scoring methodology for the proposed PN Payment measure, we refer readers to section IV.B.3.b. of the preamble of the FY 2012 IPPS/LTCH PPS final rule (76 FR 51654 through 51656) where we discuss the MSPB measure’s identical scoring methodology in detail.

We are inviting public comment on the proposed scoring methodology for the proposed PN Payment measure.

b. Proposed New Measure for the FY 2023 Program Year and Subsequent Years: Patient Safety and Adverse Events (Composite) (NQF #0531)

The current PSI 90 measure previously adopted for the Hospital VBP Program underwent NQF maintenance review and re-endorsement in 2015, leading to several substantive measure changes.55 Due to statutory requirements in the Hospital VBP Program,56 we were unable to adopt the newly re-endorsed version of the PSI 90 measure in the FY 2017 IPPS/LTCH PPS final rule (81 FR 50981), but stated our intent to propose to adopt the modified version of the PSI 90 measure in future rulemaking. In section V.J.3.b. of the preamble of this proposed rule, we are proposing to remove the current PSI 90 measure from the Hospital VBP Program.

55 National Quality Forum QFS Measure Description for “Patient Safety for Selected Indicators (modified version of PSI 90 [Composite Measure])” found at: https://www.qualityforum.org/ QFS/MeasureDetails.aspx?standardID=321&print=0 &entityTypeID=3; and PSI 90 Fact Sheet found at: http://www.qualityindicators.ahrq.gov/News/ PSI90_Factsheet_FAQ_v2.pdf (we note that this fact sheet is written from an all-payer perspective, and is therefore not limited to the measure as used in the Medicare FFS population).
56 First, section 1886(o)(2)(A) of the Act requires the Hospital VBP Program to select measures that have been specified for the Hospital IQR Program. Second, section 1886(o)(2)(C)(i) of the Act requires the Hospital VBP Program to refrain from beginning the performance period for a new measure until the measure have been posted on Hospital Compare for at least one year. Finally, section 1886(o)(3)(C) of the Act requires that the Hospital VBP Program establish performance standards for each measure not later than 60 days prior to the beginning of the performance period.

The Hospital IQR Program adopted this measure in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57128 through 57133),57 beginning with the FY 2018 payment determination, and we intend to publicly report initial measure data on the measure on Hospital Compare on or around July 2017. The full measure specifications are available at: https://www.qualityindicators.ahrq.gov/ Modules/PSI_TechSpec_ICD09_v60.aspx.

The Patient Safety and Adverse Events (Composite) measure is a weighted average of the reliability-adjusted, indirectly standardized, observed-to-expected ratios for the following 10 individual PSI component indicators:

- PSI 03 Pressure Ulcer Rate;
- PSI 06 Iatrogenic Pneumothorax Rate;
- PSI 08 In-Hospital Fall with Hip Fracture Rate; 58
- PSI 09 Perioperative Hemorrhage or Hematoma Rate;* 59
- PSI 10 Postoperative Acute Kidney Injury Requiring Dialysis Rate; 59
- PSI 11 Postoperative Respiratory Failure Rate;*
- PSI 12 Perioperative Pulmonary Embolism (PE) or Deep Vein Thrombosis (DVT) Rate;
- PSI 13 Postoperative Sepsis Rate;
- PSI 14 Postoperative Wound Dehiscence Rate; and
- PSI 15 Unrecognized Abdominopelvic Accidental Puncture/ Laceration Rate.60 61

(* Denotes new component for the Patient Safety and Adverse Events (Composite) measure)

55 We note that the HAC Reduction Program also adopted this measure in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57033 through 57030).
60 61 Previously titled “Postoperative Hip Fracture” prior to v6.0.
59 Previously titled “Postoperative Physiologic and Metabolic Derangement” prior to v6.0.
60 Previously titled “Accidental Puncture or Laceration Rate” prior to v6.0.
61 Available at: http://www.qualityforum.org/ QFS/0531.
The Patient Safety and Adverse Events (Composite) measure no longer includes PSI 07 Central Venous Catheter-Related Blood Stream Infection Rate, because of potential overlap with the CLABSI measure (NQF #0139), which has been included in the Hospital VBP Program since the FY 2013 IPPS/LTCH PPS final rule (77 FR 53597 through 53598).

The measure is calculated using administrative claims data. Like the previously adopted PSI 90 measure, under the Patient Safety and Adverse Events (Composite) measure, the predicted value for each case is computed using a Generalized Estimating Equation hierarchical modeling approach that adjusts for demographic and clinical characteristics. The expected rate for each of the indicators is computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest (that is, the hospital). The risk-adjusted rate for each of the indicators is computed using indirect standardization as the observed rate divided by the expected rate, multiplied by the reference population rate.62

As stated above, the previously adopted eight-indicator version of the PSI 90 measure underwent an extended NQF maintenance re-endorsement in the 2014 NQF Patient Safety Committee due to concerns with the underlying component indicators and their composite weights. In its final report, the NQF Patient Safety Committee deferred their final decision for the PSI 90 measure until the following measure evaluation cycle.63 Following this report, AHRQ worked to address many of the NQF stakeholders’ concerns about the PSI 90 measure, and subsequently completed NQF maintenance re-review and received re-endorsement on December 10, 2015. As a result of this process, the current PSI 90 measure’s NQF maintenance re-endorsement led to several changes to the measure, specifically: A change to the measure name; the addition of three indicators; the removal of one indicator; the re-specification of two indicators; and a revision to the weighting of component indicators.64 For more information on the proposed Patient Safety and Adverse Events (Composite) measure and component indicators, we refer readers to the Quality Indicators Empirical Methods available at:


We continue to believe the PSI 90 measure is an important measure of patient safety, addressing the NQS priority and CMS Quality Strategy goal to make care safer, and that these modifications help broaden and strengthen the measure. We expect inclusion of the Patient Safety and Adverse Events (Composite) measure in the Hospital VBP Program will encourage improvement in patient safety over the long-term for all hospitals. Conditions such as central line-associated blood stream infections, catheter-associated urinary tract infections, pressure ulcers, and other complications or conditions that arise after a patient was admitted to the hospital for the treatment of another condition are often preventable, and cost Medicare and the private sector billions of dollars each year and take a significant toll on patients and families. In most cases, hospitals can prevent these conditions when they follow protocols, procedures, and evidence-based guidelines. We anticipate the Patient Safety and Adverse Events (Composite) measure will provide actionable information and specific direction for prevention of patient safety events, because hospitals can track and monitor individual PSI rates and develop targeted improvements to patient safety using this measure data.65

We are proposing to adopt the Patient Safety and Adverse Events (Composite) measure for the Hospital VBP Program beginning with the FY 2023 program year because we believe the measure would continue to create strong incentives for hospitals to ensure that patients are not harmed by the medical care they receive, which is a critical consideration in quality improvement. We also are proposing that the measure would be added to the Safety domain, like the previously adopted PSI 90 measure that we are proposing to remove in section V.J.3.b. of the preamble of this proposed rule. The Patient Safety and Adverse Events (Composite) measure fulfills all statutory requirements for the Hospital VBP Program based on our adoption of that measure in the Hospital IQR Program and the anticipated posting of measure data on Hospital Compare at least 1 year prior to the start of the proposed measure performance period. The Patient Safety and Adverse Events (Composite) measure (MUC15–604) was included on the “List of Measures Under Consideration for December 1, 2015”66 and received support from the MAP, which noted the importance of safety measures for the Hospital VBP Program.67 Therefore, we are proposing to add the Patient Safety and Adverse Events (Composite) measure to the Safety domain for the FY 2023 program year and subsequent years.

We are inviting public comment on this proposal.

5. Previously Adopted and Proposed Baseline and Performance Periods

a. Background

Section 1886(o)(4) of the Act requires the Secretary to establish a performance period for the Hospital VBP Program that begins and ends prior to the beginning of such fiscal year. We refer readers to the FY 2016 IPPS/LTCH PPS final rule (80 FR 49561 through 49562) for the baseline and performance periods for the Clinical Care, Person and Community Engagement, Safety, and Efficiency and Cost Reduction domains that we have adopted for the FY 2018 program year. We refer readers to the FY 2017 IPPS/LTCH PPS final rule (81 FR 56998 through 57003) for additional baseline and performance periods that we have adopted for the FY 2018, FY 2019, FY 2020, FY 2021 and FY 2022 program years. Although in past rulemaking we have proposed and adopted a new baseline and performance period for each program year for each measure in each final rule, in the FY 2017 IPPS/LTCH PPS final rule, we finalized a schedule for all future baseline and performance periods.

b. Person and Community Engagement Domain

Since the FY 2015 program year, we have adopted a 12-month baseline
period and 12-month performance period for measures in the Person and Community Engagement domain (previously referred to as the Patient- and Caregiver-Centered Experience of Care/Care Coordination domain) (77 FR 53598; 78 FR 50692; 79 FR 50072; 80 FR 49561). In the FY 2017 IPPS/LTCH PPS final rule (81 FR 56998), we finalized our proposal to adopt a 12-month performance period for the Person and Community Engagement domain that runs on the calendar year two years prior to the applicable program year and a 12-month baseline period that runs on the calendar year four years prior to the applicable program year, for the FY 2019 program year and subsequent years.

We are not proposing any changes to these policies.

c. Efficiency and Cost Reduction Domain

(1) MSPB Measure

Since the FY 2016 program year, we have adopted a 12-month baseline period and 12-month performance period for the MSPB measure in the Efficiency and Cost Reduction domain.

In the FY 2017 IPPS/LTCH PPS final rule, we finalized our proposal to adopt a 12-month performance period for the MSPB measure that runs on the calendar year two years prior to the applicable program year and a 12-month baseline period that runs on the calendar year four years prior to the applicable program year for the FY 2019 program year and subsequent years (81 FR 56998). We are not proposing any changes to these policies.

(2) AMI Payment and HF Payment Measures

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 56999), we adopted a 24-month performance period and a 36-month baseline period for the AMI Payment and HF Payment measures for the FY 2022 program year (81 FR 57000). We are not proposing any changes to the length of these performance or baseline periods for the FY 2022 program year.

In the FY 2017 IPPS/LTCH PPS final rule, we also adopted a 36-month performance period and 36-month baseline period for the AMI Payment and HF Payment measures for the FY 2022 program year.

We are not proposing any changes to the length of these performance or baseline periods for the FY 2022 program year. We are not proposing any changes to the length of these performance or baseline periods for the FY 2022 program year.

For the FY 2022 program year and subsequent years, we conclude it would be appropriate to use a 36-month performance period and 36-month baseline period for the AMI Payment and HF Payment measures as we have adopted for the FY 2022 program year. Therefore, for the FY 2023 program year and subsequent years, we are proposing to adopt a 36-month performance period that runs from July 1st five years prior to the applicable fiscal program year to June 30th two years prior to the applicable fiscal program year. We also are proposing to adopt a 36-month baseline period that runs from July 1, 10 years prior to the applicable fiscal program year, to June 30, 7 years prior to the applicable fiscal program year.

We are inviting public comment on these proposals.

(3) Proposed PN Payment Measure in the FY 2022 Program Year

As discussed in section V.J.4.a. of the preamble of this proposed rule, we are proposing to adopt the PN Payment measure beginning with the FY 2022 program year. In order to adopt this measure as early as feasible into the Hospital VBP Program, we are proposing to adopt a 36-month baseline period and a 23-month performance period. We are proposing to adopt a 23-month performance period because we anticipate that the refined measure will not be posted on Hospital Compare for one year until July 2017. Therefore, for the FY 2022 program year, we are proposing to adopt a 23-month performance period that runs from August 1, 2018 to June 30, 2020 and a baseline period that runs from July 1, 2013 to June 30, 2016.

We believe that, using a 23-month performance period for the proposed PN Payment measure, rather than a 36-month performance period, in the FY 2022 program year would accurately assess the quality of care provided by hospitals and would not substantially change hospitals’ performance on the measure. To determine the viability of using a 23-month performance period to calculate the proposed PN Payment measure’s scores, we compared the measure score reliability for a 24-month and 36-month performance period. We calculated the Intraclass Correlation Coefficient (ICC) to determine the extent to which assessment of a hospital using different but randomly selected subsets of patients produces similar measures of hospital performance. 68 We calculated the risk-standardized payment (RSP) using a random split-sample of a 36-month performance period (we used July 1, 2013 through June 30, 2016) and a random split-sample of a 24-month performance period (we used July 1, 2013 through June 30, 2015).

For both the 36-month and 24-month performance period, we obtained two RSPs for each hospital, using an entirely distinct set of patients from the same time period. If the RSPs for both the 36-month and 24-month performance periods agree, we can demonstrate that the measure assesses the quality of the hospital rather than the types of patients treated. To calculate agreement between these measure subsets, we calculated the ICC (2,1) 69 for both the 36-month and 24-month performance periods.

For the proposed PN Payment measure, there were 1,170,762 index admissions and 3,242 hospitals that met the minimum case threshold for reporting a measure result (at least 25 cases) in the 36-month performance period. There were 787,817 index admissions and 1,218 hospitals that met the minimum case threshold for reporting a measure result in the 24-month performance period.

For the 36-month performance period, the ICC for the two independent assessments of each hospital was 0.868. For the 24-month performance period, the ICC for the two independent assessments of each hospital was 0.834.

Therefore, the data subsets showcase “substantial” agreement of hospital performance, and we can demonstrate that, even with a shortened performance period, the proposed PN Payment measure assesses the quality of care provided at a hospital rather than the types of patients that these hospitals treat.

To assess whether using fewer than 36 months of data change the performance in the same hospital, we compared the percent change in a hospital’s predicted/expected (P/E) ratio using 24 months of data. For hospitals that met the minimum case threshold in the 24-month performance period, the median percent change was 0.11 percent (with an interquartile range of −1.5 percent to 0.07 percent). These results suggest minimal difference in same-hospital

performance when using a 24-month measurement period. Based on these analyses, we are confident that using a 23-month performance period will result in reliable measure scores because our analysis demonstrates strong reliability at 24 months and we believe the change in available data due to a one month difference in the performance period is insufficient to substantially impact the measure’s reliability.

In summary, based on the analysis described above, we are confident that using a 23-month performance period, rather than a 35-month performance period, for the initial performance period for this measure would accurately assess the quality of care provided by that hospital and would not substantially change the hospital’s performance on that measure.

We are inviting public comment on these proposals.

(4) Proposed PN Payment Measure in the FY 2023 Program Year

We have stated in past rules that we would strive to adopt 36-month performance periods and baseline periods when possible to accommodate the time needed to process measure data and to ensure that we collect enough measure data for reliable performance scoring for all mortality measures (78 FR 50074; 79 FR 50057; and 80 FR 49588). While we cannot adopt a 36-month performance period for the FY 2023 program year because we anticipate that the refined measure will not be posted on Hospital Compare for 1 year until July 2017, we could lengthen the PN Payment measure performance period from 23 months to 35 months. As demonstrated above, our analysis of the proposed PN Payment measure indicates that the measure would produce reliable measure scores using 24 months of data as well as 36 months of data. As such, we are confident they will also be reliable when calculated using 35 months of data for the performance period for the FY 2023 program year. Therefore, for the FY 2023 program year, we are proposing to adopt a 35-month performance period that runs from August 1, 2018 to June 30, 2021 and a 36-month baseline period that runs from July 1, 2013 to June 30, 2016.

We are inviting public comment on these proposals.

(5) Proposed PN Payment Measure in the FY 2024 Program Year and Subsequent Years

For the FY 2024 program year and subsequent years, we believe it would be appropriate to use a 36-month performance period and 36-month baseline period for the PN Payment measure. Therefore, for the FY 2024 program year and subsequent years, we are proposing to adopt a 36-month baseline period and a 36-month performance period for the proposed PN Payment measure. Specifically, we are proposing to adopt a 36-month performance period that runs from July 1, 5 years prior to the applicable fiscal program year, to June 30, 2 years prior to the applicable fiscal program year and a 36-month baseline period that runs from July 1, 10 years prior to the applicable fiscal program year, to June 30, 7 years prior to the applicable fiscal program year.

We are inviting public comment on these proposals.

d. Safety Domain

(1) Previously Adopted Measures in the Safety Domain

Since the FY 2016 program year, we have adopted a 12-month baseline period and 12-month performance period for all measures in the Safety domain, with the exception of the PSI 90 measure (78 FR 50692; 79 FR 50071; 80 FR 49562). In the FY 2017 IPPS/LTCPPS final rule, we finalized our proposal to adopt a performance period for all measures in the Safety domain—with the exception of the PSI 90 measure, as discussed in more detail below—that runs on the calendar year 2 years prior to the applicable program year and a baseline period that runs on the calendar year 4 years prior to the applicable program year for the FY 2019 program year and subsequent program years (81 FR 57000).

We are not proposing any changes to these policies.

(2) Proposed Patient Safety and Adverse Events (Composite) Measure in the FY 2023 Program Year

As discussed above in section V.J.3.b. of the preamble of this proposed rule, we are proposing to remove the currently adopted PSI 90 measure beginning with the FY 2019 program year, and in section V.J.4.b. of the preamble of this proposed rule, we are proposing to adopt the Patient Safety and Adverse Events (Composite) measure beginning with the FY 2023 program year. In order to adopt the Patient Safety and Adverse Events (Composite) measure as early as feasible into the Hospital VBP Program, we are proposing to adopt a 21-month baseline period and 24-month performance period for the measure for the FY 2023 program year. Specifically, we are proposing to adopt a performance period that runs from July 1, 2019 to June 30, 2021, and a baseline period that runs from October 1, 2015 to June 30, 2017. The 21-month baseline period would only apply to the FY 2023 program year and would only use ICD–10 data.

Prior to deciding to propose an abbreviated baseline period for the FY 2023 program year, we took several factors into consideration, including the recommendations of the measure steward, the feasibility of using a combination of ICD–9 and ICD–10 data without the availability of the appropriate measure software, minimizing provider burden, program implementation timelines, and the reliability of using a shortened baseline period. We believe using a 21-month baseline period for the Patient Safety and Adverse Events (Composite) measure for the FY 2023 program year best serves the need to provide important information on hospital patient safety and adverse events by allowing sufficient time to process the claims data and calculate measure scores, while minimizing reporting burden and program disruption. We also believe that measure scores would continue to be reliable for the above proposed baseline period because the NQF, which re-endorsed the modified version of the measure that we are now proposing, found it to be reliable using 12 months of data.74

We are inviting public comment on these proposals.

(3) Proposed Patient Safety and Adverse Events (Composite) Measure in the FY 2024 Program Year and Subsequent Years

For the FY 2024 program year and subsequent years, we are proposing to lengthen the Patient Safety and Adverse Events (Composite) measure baseline period to 24 months and continue to adopt a 24-month performance period because we believe the measure is most reliable with a 24-month baseline period. For the FY 2024 program year, the baseline period would run from July 1, 2016 to June 30, 2018. Therefore, we are proposing to adopt a performance period that runs from July 1, 4 years prior to the applicable fiscal program year, to June 30, 2 years prior to the applicable fiscal program year, and a baseline period that runs from July 1, 8 years prior to the applicable program year, to June 30, 6 years prior to the applicable program year.

We are inviting public comment on these proposals.

e. Clinical Care Domain
(1) Previously Adopted Measures in the Clinical Care Domain

For the FY 2019, FY 2020, and FY 2021 program years, we adopted a 36-month baseline period and 36-month performance period for measures in the Clinical Care domain (78 FR 50692 through 50694; 79 FR 50073; 80 FR 49563). In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57001), we finalized our proposal to adopt a 36-month performance period and 36-month baseline period for the FY 2022 program year for each of the previously finalized measures in the Clinical Care domain—that is, the MORT–30–AMI, MORT–30–HF, MORT–30–COPD, THA/TKA, and MORT–30–CABG measures. We are now proposing to adopt a 36-month performance period and 36-month baseline period for these measures for the FY 2023 program year and subsequent years.

Specifically, for the mortality measures (MORT–30–AMI, MORT–30–HF, MORT–30–COPD, and MORT–30–CABG), the performance period would run for 36 months from July 1, 5 years prior to the applicable fiscal program year, to June 30, 2 years prior to the applicable fiscal program year, and the baseline period would run for 36 months from July 1, 10 years prior to the applicable fiscal program year, to June 30, 7 years prior to the applicable fiscal program year. For the THA/TKA measure, the performance period would run for 36 months from April 1, 5 years prior to the applicable fiscal program year, to March 31, 2 years prior to the applicable fiscal program year, and the baseline period would run for 36 months from April 1, 10 years prior to the applicable fiscal program year, to March 31, 7 years prior to the applicable fiscal program year.

We are inviting public comment on these proposals.

(2) MORT–30–PN (Updated Cohort) Measure

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57001), we adopted a 22-month performance period for the MORT–30–PN (updated cohort) measure and a 36-month baseline period for the FY 2021 program year. In the same final rule, we adopted a 34-month performance period and 36-month baseline period for the MORT–30–PN (updated cohort) measure for the FY 2022 program year. We are not proposing any changes to the length of these performance or baseline periods for the FY 2021 and FY 2022 program years.

f. Summary of Previously Adopted and Proposed Baseline and Performance Periods for the FY 2019 through FY 2023 Program Years

The tables below summarize the baseline and performance periods that we have previously adopted and are proposing to adopt in this proposed rule.

### PREVIOUSLY ADOPTED BASELINE AND PERFORMANCE PERIODS FOR THE FY 2019 PROGRAM YEAR

<table>
<thead>
<tr>
<th>Domain</th>
<th>Baseline period</th>
<th>Performance period</th>
</tr>
</thead>
<tbody>
<tr>
<td>Person and Community Engagement:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Clinical Care:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• THA/TKA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Safety: *</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• PC–01 and NHSN measures (CAUTI, CLABSI, SSI, CDI, MRSA)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Efficiency and Cost Reduction:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• MSPB</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* As discussed in section V.I.3.b. of the preamble of this proposed rule, we are proposing to remove the current PSI 90 measure beginning with the FY 2019 program year. As a result, the previously finalized performance and baseline periods for this measure are not included in this table.

### PREVIOUSLY ADOPTED BASELINE AND PERFORMANCE PERIODS FOR THE FY 2020 PROGRAM YEAR

<table>
<thead>
<tr>
<th>Domain</th>
<th>Baseline period</th>
<th>Performance period</th>
</tr>
</thead>
<tbody>
<tr>
<td>Person and Community Engagement:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• HCAHPS Survey</td>
<td>January 1, 2016–December 31, 2016</td>
<td>January 1, 2018–December 31, 2018</td>
</tr>
<tr>
<td>Clinical Care:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• THA/TKA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Safety: *</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• PC–01 and NHSN measures (CAUTI, CLABSI, SSI, CDI, MRSA)</td>
<td></td>
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</tr>
</tbody>
</table>

* The THA/TKA measure was added for the FY 2019 program year with a 36-month baseline period and a 24-month performance period (79 FR 50072), but we have since adopted 36-month baseline and performance periods for the FY 2021 program year (80 FR 49563). We intend to continue having 36-month baseline periods and 36-month performance periods in the future for all measures in the Clinical Care domain.
PREVIOUSLY ADOPTED BASELINE AND PERFORMANCE PERIODS FOR THE FY 2020 PROGRAM YEAR—Continued

<table>
<thead>
<tr>
<th>Domain</th>
<th>Baseline period</th>
<th>Performance period</th>
</tr>
</thead>
<tbody>
<tr>
<td>Efficiency and Cost Reduction:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• MSPB</td>
<td>January 1, 2016–December 31, 2016</td>
<td>January 1, 2018–December 31, 2018</td>
</tr>
</tbody>
</table>

*As discussed in section V.J.3.b. of the preamble of this proposed rule, we are proposing to remove the current PSI 90 measure beginning with the FY 2019 program year. As a result, the previously finalized performance and baseline periods for this measure are not included in this table.

PREVIOUSLY ADOPTED BASELINE AND PERFORMANCE PERIODS FOR THE FY 2021 PROGRAM YEAR

| Domain                                  | Baseline period                  | Performance period                |
|-----------------------------------------|----------------------------------|                                   |
| Person and Community Engagement:        |                                  |                                   |
| • HCAHPS Survey                         | January 1, 2017–December 31, 2017| January 1, 2019–December 31, 2019 |
| Clinical Care:                          |                                  |                                   |
| • THA/TKA                               | April 1, 2011–March 31, 2014     | April 1, 2016–March 31, 2019.    |
| Safety:                                 |                                  |                                   |
| • PC–01 and NHSN measures (CAUTI, CLABSI, SSI, CDI, MRSA) | January 1, 2017–December 31, 2017 | January 1, 2019–December 31, 2019 |
| Efficiency and Cost Reduction:          |                                  |                                   |
| • MSPB                                  | January 1, 2017–December 31, 2017| January 1, 2019–December 31, 2019 |
| • Payment (AMI Payment and HF Payment)  | July 1, 2012–June 30, 2015       | January 1, 2019–December 31, 2019 |

*As discussed in section V.J.3.b. of the preamble of this proposed rule, we are proposing to remove the current PSI 90 measure beginning with the FY 2019 program year. As a result, the previously finalized performance and baseline periods for this measure are not included in this table.

PREVIOUSLY ADOPTED AND PROPOSED BASELINE AND PERFORMANCE PERIODS FOR THE FY 2022 PROGRAM YEAR

| Domain                                  | Baseline period                  | Performance period                |
|-----------------------------------------|----------------------------------|                                   |
| Person and Community Engagement:        |                                  |                                   |
| • HCAHPS Survey                         | January 1, 2018–December 31, 2018| January 1, 2020–December 31, 2020 |
| Clinical Care:                          |                                  |                                   |
| • THA/TKA                               |                                  |                                   |
| Safety:                                 |                                  |                                   |
| • PC–01 and NHSN measures (CAUTI, CLABSI, SSI, CDI, MRSA) | January 1, 2018–December 31, 2018 | January 1, 2020–December 31, 2020 |
| Efficiency and Cost Reduction:          |                                  |                                   |
| • MSPB                                  | January 1, 2018–December 31, 2018| January 1, 2020–December 31, 2020 |
| • Payment (AMI Payment, HF Payment)     | July 1, 2012–June 30, 2015       | January 1, 2020–December 31, 2020 |

*As discussed in section V.J.3.b. of the preamble of this proposed rule, we are proposing to remove the current PSI 90 measure beginning with the FY 2019 program year. As a result, the previously finalized performance and baseline periods for this measure are not included in this table.** As discussed in section V.J.4.a. of the preamble of this proposed rule, we are proposing to adopt the PN Payment measure beginning with the FY 2021 program year.

PREVIOUSLY ADOPTED AND PROPOSED BASELINE AND PERFORMANCE PERIODS FOR THE FY 2023 PROGRAM YEAR

| Domain                                  | Baseline period                  | Performance period                |
|-----------------------------------------|----------------------------------|                                   |
| Person and Community Engagement:        |                                  |                                   |
| • HCAHPS Survey                         | January 1, 2019–December 31, 2019| January 1, 2021–December 31, 2021 |
| Clinical Care:                          |                                  |                                   |
| Safety:                                 |                                  |                                   |
| • PC–01 and NHSN measures (CAUTI, CLABSI, SSI, CDI, MRSA) | January 1, 2019–December 31, 2019 | January 1, 2021–December 31, 2021 |
| • Patient Safety and Adverse Events     | October 1, 2015–June 30, 2017    | January 1, 2019–December 31, 2021 |
| (Composite) *                          |                                  |                                   |
| Efficiency and Cost Reduction:          |                                  |                                   |

| Efficiency and Cost Reduction:          |                                  |                                   |
| • MSPB                                  | January 1, 2018–December 31, 2018| January 1, 2020–December 31, 2020 |
| • Payment (AMI Payment, HF Payment)     | July 1, 2012–June 30, 2015       | January 1, 2020–December 31, 2020 |
6. Proposed Performance Standards for the Hospital VBP Program

a. Background

Section 1886(o)(3)(A) of the Act requires the Secretary to establish performance standards for the measures selected under the Hospital VBP Program for a performance period for the applicable fiscal year. The performance standards must include levels of achievement and improvement, as required by section 1886(o)(3)(B) of the Act, and must be established no later than 60 days before the beginning of the performance period for the fiscal year involved, as required by section 1886(o)(3)(C) of the Act. We refer readers to the Hospital Inpatient VBP Program final rule (76 FR 26511 through 26513) for further discussion of the general methodology used in the Hospital VBP Program.

In addition, when establishing the performance standards, section 1886(o)(3)(D) of the Act requires the Secretary to consider appropriate factors, such as: (1) Practical experience with the measures, including whether a significant proportion of hospitals failed to meet the performance standard during previous performance periods; (2) historical performance standards; (3) improvement rates; and (4) the opportunity for continued improvement.

We refer readers to the FY 2013, FY 2014, and FY 2015 IPPS/LTCH PPS final rules (77 FR 53604 through 53605; 78 FR 50694 through 50698; and 79 FR 50077 through 50079, respectively) for a more detailed discussion of the general scoring methodology used in the Hospital VBP Program.

We note that the performance standards for the following measures are calculated with lower values representing better performance:

- The NHNS measures (the CLABSI, CAUTI, CDI, Colon and the Abdominal Hysterectomy SSI, and MRSA Bacteremia measures);
- The THA/TKA measure;
- The PC–01 measure;
- The MSPB measure;
- The HF and AMI Payment measures;
- The proposed PN Payment measure; and
- The proposed Patient Safety and Adverse Events (Composite) measure.

This distinction is made in contrast to other measures for which higher values indicate better performance.\(^7\)\(^3\) As discussed further in the FY 2014 IPPS/LTCH PPS final rule (78 FR 50684), the performance standards for the Colon and Abdominal Hysterectomy SSI measure are computed separately for each procedure stratum, and we first award achievement and improvement points to each stratum separately, then compute a weighted average of the points awarded to each stratum by predicted infections.

b. Previously Adopted and Proposed Performance Standards for the FY 2020 Program Year

In accordance with our finalized methodology for calculating performance standards (discussed more fully in the Hospital Inpatient VBP Program final rule (76 FR 26511 through 26513)), we are proposing to adopt additional performance standards for the FY 2020 program year. We note that the numerical values for the performance standards displayed in this proposed rule, below, represented estimates based on the most recently available data, and we intend to update the numerical values in the FY 2018 IPPS/LTCH PPS final rule. We note further that the MSPB measure’s performance standards are based on performance period data; therefore, we are unable to provide numerical equivalents for the standards at this time. These previously adopted and newly proposed performance standards for the measures in the FY 2020 program year are set out in the tables below.

### PREVIOUSLY ADOPTED AND PROPOSED PERFORMANCE STANDARDS FOR THE FY 2020 PROGRAM YEAR: SAFETY, CLINICAL CARE, AND EFFICIENCY AND COST REDUCTION DOMAINS

<table>
<thead>
<tr>
<th>Measure short name</th>
<th>Achievement threshold</th>
<th>Benchmark</th>
</tr>
</thead>
<tbody>
<tr>
<td>CAUTI *</td>
<td>0.806</td>
<td>0.000</td>
</tr>
<tr>
<td>CLABSI *</td>
<td>0.797</td>
<td>0.000</td>
</tr>
<tr>
<td>CDI *</td>
<td>0.876</td>
<td>0.090</td>
</tr>
<tr>
<td>MRSA Bacteremia *</td>
<td>0.794</td>
<td>0.000</td>
</tr>
<tr>
<td>Colon and Abdominal Hysterectomy SSI *</td>
<td>0.784</td>
<td>0.000</td>
</tr>
<tr>
<td>PC–01 *</td>
<td>0.005952</td>
<td>0.875869</td>
</tr>
</tbody>
</table>

\(^7\)\(^3\) We note that the mortality measures in the Hospital VBP Program use survival rates rather than mortality rates; as a result, higher values indicate better performance on these measures.
PREVIOUSLY ADOPTED AND PROPOSED PERFORMANCE STANDARDS FOR THE FY 2020 PROGRAM YEAR: SAFETY, CLINICAL CARE, AND EFFICIENCY AND COST REDUCTION DOMAINS—Continued

<table>
<thead>
<tr>
<th>Measure short name</th>
<th>Achievement threshold</th>
<th>Benchmark</th>
</tr>
</thead>
<tbody>
<tr>
<td>MORT–30–HF ±</td>
<td>0.881090</td>
<td>0.906086.</td>
</tr>
<tr>
<td>MORT–30–PN ±</td>
<td>0.882266</td>
<td>0.909532.</td>
</tr>
<tr>
<td>THA/TKA ±</td>
<td>0.032229</td>
<td>0.023178.</td>
</tr>
</tbody>
</table>

### Efficiency and Cost Reduction Domain

<table>
<thead>
<tr>
<th>MSPB ±</th>
<th>Median Medicare Spending Per Beneficiary ratio across all hospitals during the performance period</th>
<th>Mean of the lowest decile Medicare Spending Per Beneficiary ratios across all hospitals during the performance period</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
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</tbody>
</table>

As discussed in section V.J.3.b. of the preamble of this proposed rule, we are proposing to remove the current PSI 90 measure beginning with the FY 2019 program year. As a result, the previously finalized performance standards for this measure are not included in this table.

- The performance standards displayed in this table for the Safety domain measures were calculated using one quarter (Q4) CY 2015 data and three quarters (Q1, Q2, and Q3) CY 2016 data. We will update this table’s performance standards using four quarters of CY 2016 data in the final rule.

- In section III.F.2.e. of preamble of the FY 2016 IPPS/LTCH PPS final rule (80 FR 49544 through 49555), we finalized our proposal to use the CDC’s new standard population data to calculate performance standards for the NHSN measures beginning with the FY 2019 program year. We refer readers to that final rule for additional information regarding the NHSN measures’ standard population data. In addition, we note that a technical update was released for these measures for the FY 2019 program year in order to ensure that hospitals have the correct performance standards for the applicable performance period.

- Lower values represent better performance.
- Previously adopted performance standards.

In the CY 2017 OPPS/ASC final rule with comment period (81 FR 79857), we discussed how the removal of the Pain Management dimension of the HCAHPS Survey, beginning with the FY 2018 program year, affects the scoring of the Person and Community Engagement domain. The eight dimensions of the HCAHPS measure are calculated to generate the HCAHPS Base Score. For each of the eight dimensions, Achievement Points (0–10 points) and Improvement Points (0–9 points) are calculated, the larger of which is then summed across the eight dimensions to create the HCAHPS Base Score (0–80 points). Each of the eight dimensions is of equal weight, thus the HCAHPS Base Score ranges from 0 to 80 points. HCAHPS Consistency Points are then calculated, which range from 0 to 20 points. The Consistency Points take into consideration the scores of all eight Person and Community Engagement dimensions; as noted above, the Pain Management dimension is not included in the scoring of this Domain. The final element of the scoring formula is the summation of the HCAHPS Base Score and the HCAHPS Consistency Points, which results in the Person and Community Engagement Domain score that ranges from 0 to 100 points.

PROPOSED PERFORMANCE STANDARDS FOR THE FY 2020 PROGRAM YEAR: PERSON AND COMMUNITY ENGAGEMENT DOMAIN ±

<table>
<thead>
<tr>
<th>HCAHPS survey dimension</th>
<th>Floor (percent)</th>
<th>Achievement threshold (percent)</th>
<th>Benchmark (percent)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Communication with Nurses</td>
<td>49.26</td>
<td>78.99</td>
<td>87.17</td>
</tr>
<tr>
<td>Communication with Doctors</td>
<td>46.91</td>
<td>80.31</td>
<td>88.56</td>
</tr>
<tr>
<td>Responsiveness of Hospital Staff</td>
<td>35.92</td>
<td>65.16</td>
<td>80.05</td>
</tr>
<tr>
<td>Communication about Medicines</td>
<td>23.44</td>
<td>63.41</td>
<td>73.94</td>
</tr>
<tr>
<td>Hospital Cleanliness &amp; Quietness</td>
<td>37.21</td>
<td>65.81</td>
<td>79.29</td>
</tr>
<tr>
<td>Discharge Information</td>
<td>65.60</td>
<td>87.36</td>
<td>92.04</td>
</tr>
<tr>
<td>Care Transition</td>
<td>21.20</td>
<td>51.12</td>
<td>62.56</td>
</tr>
<tr>
<td>Overall Rating of Hospital</td>
<td>35.46</td>
<td>71.35</td>
<td>85.01</td>
</tr>
</tbody>
</table>

± We renamed this domain from Patient- and Caregiver-Centered Experience of Care/Care Coordination domain to Person and Community Engagement domain beginning with the FY 2019 program year, as discussed in the FY 2017 IPPS/LTCH PPS final rule (81 FR 55984).

The performance standards displayed in this table were calculated using one quarter (Q4) CY 2015 data and three quarters (Q1, Q2, and Q3) CY 2016 data. We will update this table’s performance standards using four quarters of CY 2016 data in the final rule.

We are inviting public comments on these proposed performance standards for the FY 2020 program year.

c. Previously Adopted Performance Standards for Certain Measures for the FY 2021 Program Year

As discussed above, we have adopted certain measures for the Clinical Care and Efficiency and Cost Reduction domains for future program years in order to ensure that we can adopt baseline and performance periods of sufficient length for performance scoring purposes. In the FY 2016 IPPS/LTCH PPS final rule (80 FR 49567), we adopted performance standards for the FY 2021 program year for the Clinical Care domain measures (THA/TKA, MORT–30–HF, MORT–30–AMI, and MORT–30–COPD). In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57008), we adopted performance standards for the MORT–30–PN (updated cohort) measure (81 FR 57008) and the AMI Payment and HF Payment measures for the FY 2021 program year. We note that the performance standards for the MSPB, AMI Payment, and HF Payment measures are based on performance.
Per Beneficiary ratios across all hospitals during the performance period.

Median Medicare Spending Per Beneficiary ratio across all hospitals during the performance period.

Preceding the performance period.

Per Beneficiary ratios across all hospitals during the performance period.

Median Medicare Spending Per Beneficiary ratio across all hospitals during the performance period.

Mean of the lowest decile Medicare Spending Per Beneficiary ratios across all hospitals during the performance period.

Mean of the lowest decile Medicare Spending Per Beneficiary ratios across all hospitals during the performance period.

Mean of the lowest decile Hospital-Level, Risk-Standardized Payment Associated with a 30-Day Episode-of-Care across all hospitals during the performance period.

Mean of the lowest decile Hospital-Level, Risk-Standardized Payment Associated with a 30-Day Episode-of-Care across all hospitals during the performance period.

Previously Adopted and Proposed Performance Standards for the FY 2022 Program Year

As discussed above, we have adopted certain measures for the Clinical Care and Efficiency and Cost Reduction domains for future program years in order to ensure that we can adopt baseline and performance periods of sufficient length for performance scoring purposes. In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57009), we adopted performance standards for the FY 2022 program year for the Clinical Care domain measures (THA/TKA, MORT–30–HF, MORT–30–AMI, MORT–30–PN (updated cohort), MORT–30–COPD, and MORT–30–CABG) and the Efficiency and Cost Reduction domain measures (AMI Payment and HF Payment). In section V.J.4.a. of the preamble of this proposed rule, we are proposing to add one measure, the PN Payment measure, beginning with the FY 2022 program year. We note that the performance standards for the MSPB, AMI Payment, HF Payment, and PN Payment measures are based on performance period data; therefore, we are unable to provide numerical equivalents for the standards at this time. The previously adopted and newly proposed performance standards for these measures are set out in the table below.

PREVIOUSLY ADOPTED AND PROPOSED PERFORMANCE STANDARDS FOR THE FY 2022 PROGRAM YEAR

<table>
<thead>
<tr>
<th>Measure short name</th>
<th>Achievement threshold</th>
<th>Benchmark</th>
</tr>
</thead>
<tbody>
<tr>
<td>MORT–30–AMI *</td>
<td>0.861793</td>
<td>0.861305</td>
</tr>
<tr>
<td>MORT–30–HF *</td>
<td>0.879869</td>
<td>0.903608</td>
</tr>
<tr>
<td>MORT–30–PN (updated cohort) *</td>
<td>0.836122</td>
<td>0.870506</td>
</tr>
<tr>
<td>MORT–30–COPD *</td>
<td>0.920058</td>
<td>0.936662</td>
</tr>
<tr>
<td>MORT–30–CABG *</td>
<td>0.968210</td>
<td>0.979000</td>
</tr>
<tr>
<td>THA/TKA *</td>
<td>0.029833</td>
<td>0.021493</td>
</tr>
</tbody>
</table>
### PREVIOUSLY ADOPTED AND PROPOSED PERFORMANCE STANDARDS FOR THE FY 2022 PROGRAM YEAR—Continued

<table>
<thead>
<tr>
<th>Measure short name</th>
<th>Achievement threshold</th>
<th>Benchmark</th>
</tr>
</thead>
<tbody>
<tr>
<td>MSPB *</td>
<td>Median Medicare Spending Per Beneficiary ratio across all hospitals during the performance period.</td>
<td>Mean of the lowest decile Medicare Spending Per Beneficiary ratios across all hospitals during the performance period.</td>
</tr>
<tr>
<td>AMI Payment *</td>
<td>Median Hospital-Level, Risk-Standardized Payment Associated with a 30-Day Episode-of-Care across all hospitals during the performance period.</td>
<td>Mean of the lowest decile Hospital-Level, Risk-Standardized Payment Associated with a 30-Day Episode-of-Care across all hospitals during the performance period.</td>
</tr>
<tr>
<td>HF Payment *</td>
<td>Median Hospital-Level, Risk-Standardized Payment Associated with a 30-Day Episode-of-Care across all hospitals during the performance period.</td>
<td>Mean of the lowest decile Hospital-Level, Risk-Standardized Payment Associated with a 30-Day Episode-of-Care across all hospitals during the performance period.</td>
</tr>
<tr>
<td>PN Payment *</td>
<td>Median Hospital-Level, Risk-Standardized Payment Associated with a 30-Day Episode-of-Care across all hospitals during the performance period.</td>
<td>Mean of the lowest decile Hospital-Level, Risk-Standardized Payment Associated with a 30-Day Episode-of-Care across all hospitals during the performance period.</td>
</tr>
</tbody>
</table>

*Previously adopted performance standards.

† After publication of the FY 2017 IPPS/LTC PPS final rule, we determined there was a display error in the performance standards for this measure. Specifically, the Achievement Threshold and Benchmark values, while accurate, were presented in the wrong categories. We have corrected this issue in the table above, and the correct performance standards are displayed here.

* Lower values represent better performance.

 Scored the same as the MSPB, AMI Payment, and HF Payment measures, as discussed in section V.J.4.a.(2) of the preamble of this proposed rule.

We are inviting public comment on the proposed PN Payment measure performance standards for the FY 2022 program year.

e. Proposed Performance Standards for Certain Measures for the FY 2023 Program Year

As discussed above, we have adopted certain measures for the Clinical Care and Efficiency and Cost Reduction domains for future program years in order to ensure that we can adopt baseline and performance periods of sufficient length for performance scoring purposes. We are proposing the following performance standards for the FY 2023 program year for the Clinical Care domain measures (THA/TKA, MORT–30–AMI, MORT–30–HF, MORT–30–PN (updated cohort), MORT–30–COPD, and MORT–30–CABG) and for the Efficiency and Cost Reduction domain measures (MSPB, AMI Payment, HF Payment, and the proposed PN Payment measure). Although we are proposing to adopt the Patient Safety and Adverse Events (Composite) measure beginning with the FY 2023 program year, we do not currently have data available to calculate the performance standards; we therefore intend to propose the FY 2023 performance standards for this measure in next year’s rulemaking. We note that the performance standards for the MSPB, AMI Payment, HF Payment, and PN Payment measures are based on performance period data; therefore, we are unable to provide numerical equivalents for the standards at this time. These newly proposed performance standards for these measures are set out in the table below.

### PROPOSED PERFORMANCE STANDARDS FOR THE FY 2023 PROGRAM YEAR

<table>
<thead>
<tr>
<th>Measure short name</th>
<th>Achievement threshold</th>
<th>Benchmark</th>
</tr>
</thead>
<tbody>
<tr>
<td>MORT–30–AMI</td>
<td>0.866548</td>
<td>0.85499.</td>
</tr>
<tr>
<td>MORT–30–HF</td>
<td>0.881939</td>
<td>0.906798.</td>
</tr>
<tr>
<td>MORT–30–PN (updated cohort)</td>
<td>0.840138</td>
<td>0.871741.</td>
</tr>
<tr>
<td>MORT–30–COPD</td>
<td>0.919769</td>
<td>0.936349.</td>
</tr>
<tr>
<td>MORT–30–CABG</td>
<td>0.966747</td>
<td>0.979620.</td>
</tr>
<tr>
<td>THA/TKA*</td>
<td>0.027428</td>
<td>0.019779.</td>
</tr>
</tbody>
</table>
Hospitals must receive a minimum of one measure score within the Efficiency and Cost Reduction domain.

b. Proposed Domain Weighting for the FY 2019 Program Year and Subsequent Years for Hospitals Receiving Scores on Fewer Than Four Domains

For the FY 2017 program year and subsequent years, we adopted a policy that hospitals must receive domain scores on at least three of four quality domains in order to receive a TPS, and hospitals with sufficient data on only three domains will have their TPSs proportionately reweighted (79 FR 50084 through 50085). We are not proposing any changes to these domain weights for the FY 2019 program year or subsequent years.

For a hospital to receive a TPS for the FY 2019 program year and subsequent years:

• Hospitals must report a minimum number of 100 completed HCAHPS surveys for a hospital to receive a Person and Community Engagement domain score.
• Hospitals must receive a minimum of two measure scores within the Clinical Care domain.
• Hospitals must receive a minimum of two measure scores within the Safety domain.

We are proposing two changes to our domain scoring policies for the FY 2019 program year and subsequent years. We are proposing to change the minimum number of measures scores a hospital must receive to receive a score on the Safety domain from three measures to two measures. Second, we are proposing that hospitals must receive a minimum of one measure score within the Efficiency and Cost Reduction domain to receive a domain score rather than requiring that hospitals meet the
requirements to receive a MSPB measure score.

The proposed change to the Safety domain minimum number of measure scores is based on our proposal to remove the current PSI 90 measure from the Hospital VBP Program beginning with the FY 2019 program year. Based on our analyses, removing this measure but maintaining the requirement that a hospital receive three measure scores in order to receive a Safety Domain score would have a significant impact on the number of hospitals eligible to receive a Safety domain score. Therefore, in order to include the greatest number of hospitals in the Hospital VBP Program possible while ensuring the need for TPSs to be sufficiently reliable, we are proposing to reduce the minimum number of required measure scores within the Safety domain from three measures to two.

In addition, we note that we are not proposing to reduce the number of measures a hospital must receive a score on in order to receive an Efficiency and Cost Reduction domain score. Under the current program requirements (79 FR 50086), a hospital must be eligible to receive a score on the MSPB measure in order to receive a score for this domain. In the FY 2017 IPPS/LTCH PPS final rule (81 FR 56987 through 56990 and 81 FR 56990 through 56992), we adopted two condition-specific payment measures, the AMI Payment and HF Payment measures, beginning with the FY 2021 program year, and in section V.J.4.a. of the preamble of this proposed rule we are proposing to adopt one additional condition-specific payment measure, the PN Payment measure. We are therefore proposing to require that hospitals must be eligible to receive a score on at least one measure within the Efficiency and Cost Reduction domain, rather than on the MSPB measure specifically, to reflect this expansion of the domain’s measure set.

We believe these proposed changes reflect the evolution of the Hospital VBP Program measure set, and we continue to believe that these requirements appropriately balance our desire to enable as many hospitals as possible to participate in the Hospital VBP Program and the need for TPSs to be sufficiently reliable to provide meaningful distinctions between hospitals’ performance on quality measures.

We are inviting public comment on these proposals.

c. Minimum Numbers of Cases for Hospital VBP Program Measures for the FY 2019 Program Year and Subsequent Years

(1) Background

Section 1886(o)(1)(C)(ii)(IV) of the Act requires the Secretary to exclude for the fiscal year hospitals that do not report a minimum number (as determined by the Secretary) of cases for the measures that apply to the hospital for the performance period for the fiscal year. Under section 1886(o)(1)(C)(iii) of the Act, in determining the minimum number of reported cases for a given measure, the Secretary must conduct an independent analysis of what minimum numbers would be appropriate. For additional discussion of the previously finalized minimum numbers of cases for measures under the Hospital VBP Program, we refer readers to the Hospital Inpatient VBP Program final rule (76 FR 26527 through 26531); the CY 2012 OPPS/ASC final rule (76 FR 74532 through 74534); the FY 2013 IPPS/LTCH PPS final rule (77 FR 53608 through 53609); the FY 2015 IPPS/LTCH PPS final rule (79 FR 50085); the FY 2016 IPPS/LTCH PPS final rule (80 FR 49570); and the FY 2017 IPPS/LTCH PPS final rule (81 FR 57011).

(2) Person and Community Engagement Domain

In the Hospital Inpatient VBP Program final rule (76 FR 26527 through 26531), we adopted a minimum number of 100 completed HCAHPS Surveys for a hospital to receive a score on the HCAHPS measure.

We are not proposing any changes to this policy.

(3) Clinical Care Domain

In the CY 2012 OPPS/ASC final rule with comment period (76 FR 74532 through 74534), we adopted a minimum number of 10 cases for the PC–01 measure.74 In the CY 2014 IPPS/LTCH PPS final rule (77 FR 53608 through 53609), we adopted a minimum number of 25 cases in order to receive a score on the PC–01 measure.

In the FY 2015 IPPS/LTCH PPS final rule (79 FR 50085), we adopted a minimum of one predicted infection for NHSN-based surveillance measures (that is, the CAUTI, CLABSI, CDI, MRSA, and SSI measures) based on CDC’s minimum case criteria. In the FY 2015 IPPS/LTCH PPS final rule (79 FR 50085), we adopted this case minimum for the NHSN-based surveillance measures FY 2016 Hospital VBP Program and subsequent years. In the FY 2012 IPPS/LTCH PPS final rule (76 FR 26530), we adopted a minimum of 10 cases for the PC–01 measure.74

Beginning with the FY 2023 program year, we are proposing that hospitals must report a minimum of three eligible cases on any one underlying indicator during the baseline period in order to receive an improvement score and three eligible cases on any one underlying indicator during performance period in order to receive an achievement score on the Patient Safety and Adverse Events (Composite) measure. For the purposes of the Patient Safety and Adverse Events (Composite) measure, a case is “eligible” for a given indicator if it meets the criterion for inclusion in the indicator measure population. This minimum number of cases is based on AHRQ’s methodology for scoring performance on the Patient Safety and Adverse Events (Composite) measure. We note that these proposed minimum data requirements for the Patient Safety and Adverse Events (Composite) measure are the same as those previously finalized for the current PSI 90 measure.

We are inviting public comment on our proposal regarding the minimum number of cases for the Patient Safety and Adverse Events (Composite) measure.

(5) Efficiency and Cost Reduction Domain

In the FY 2013 IPPS/LTCH PPS final rule (77 FR 53608 through 53610), we adopted a minimum of 25 cases in order to receive a score for the MSPB measure. In the FY 2015 IPPS/LTCH PPS final rule (79 FR 50085 through 50086), we retained the same MSPB measure case minimum for the FY 2016 program year and subsequent years. In the FY 2017 IPPS/LTCH PPS final rule (81 FR 56990 through 56992, respectively),

74 We note that the PC–01 measure was previously included in the Clinical Care—Process domain. In the FY 2016 IPPS/LTCH PPS final rule (80 FR 49553 through 49554), we re-categorized this measure as a Safety domain measure beginning with the FY 2016 program year.
we adopted the AMI Payment and HF Payment measures in the Efficiency and Cost Reduction domain for the FY 2021 program year and subsequent years. In section V.I.4.a. of the preamble of this proposed rule, we are proposing to adopt the PN Payment measure in the Efficiency and Cost Reduction domain for the FY 2022 program year and subsequent years. For these condition-specific payment measures (namely, the AMI Payment and HF Payment measures, as well as the proposed PN Payment measure, if finalized), we are proposing that hospitals must report a minimum number of 25 cases per measure in order to receive a measure score for the FY 2021 program year. FY 2022 program year, and subsequent years. We believe this minimum number of cases is appropriate because it balances our interest in allowing the maximum possible number of hospitals the opportunity to receive a score on the measure and maintaining sufficiently reliable scores. As we noted in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56992), we expect this case minimum will ensure that each hospital’s payment measure rate is sufficiently reliable to generate a score that meaningfully distinguishes hospital performance on the measures. In addition, the statistical model that CMS uses to calculate the payment measures allows for the inclusion of hospitals with relatively few cases by taking into account the uncertainty associated with sample size.

We are inviting public comment on our proposal regarding the minimum number of cases for the AMI, HF, and PN Payment measures.

(6) Summary of Previously Adopted and Proposed Minimum Numbers of Cases for the FY 2019 Program Year and Subsequent Years

These previously adopted and newly proposed minimum numbers of cases for these measures are set forth in the table below.

PREVIOUSLY ADOPTED AND PROPOSED MINIMUM CASE NUMBER REQUIREMENTS FOR THE FY 2019 PROGRAM YEAR AND SUBSEQUENT YEARS

<table>
<thead>
<tr>
<th>Measure short name</th>
<th>Minimum number of cases</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Person and Community Engagement Domain</strong></td>
<td></td>
</tr>
<tr>
<td>HCAHPS</td>
<td>Hospitals must report a minimum number of 100 completed HCAHPS surveys.</td>
</tr>
<tr>
<td><strong>Clinical Care Domain</strong></td>
<td></td>
</tr>
<tr>
<td>MORT–30–AMI</td>
<td>Hospitals must report a minimum number of 25 cases.</td>
</tr>
<tr>
<td>MORT–30–HF</td>
<td>Hospitals must report a minimum number of 25 cases.</td>
</tr>
<tr>
<td>MORT–30–PN (updated cohort)</td>
<td>Hospitals must report a minimum number of 25 cases.</td>
</tr>
<tr>
<td>MORT–30–COPD</td>
<td>Hospitals must report a minimum number of 25 cases.</td>
</tr>
<tr>
<td>MORT–30–CABG</td>
<td>Hospitals must report a minimum number of 25 cases.</td>
</tr>
<tr>
<td>THA/TKA</td>
<td>Hospitals must report a minimum number of 25 cases.</td>
</tr>
<tr>
<td><strong>Safety Domain</strong></td>
<td></td>
</tr>
<tr>
<td>CAUTI</td>
<td>Hospitals have a minimum of 1,000 predicted infections as calculated by the CDC.</td>
</tr>
<tr>
<td>CLABSI</td>
<td>Hospitals have a minimum of 1,000 predicted infections as calculated by the CDC.</td>
</tr>
<tr>
<td>Colon and Abdominal Hysterectomy SSI</td>
<td>Hospitals have a minimum of 1,000 predicted infections as calculated by the CDC.</td>
</tr>
<tr>
<td>MRSA Bacteremia</td>
<td>Hospitals have a minimum of 1,000 predicted infections as calculated by the CDC.</td>
</tr>
<tr>
<td>CDI</td>
<td>Hospitals have a minimum of 1,000 predicted infections as calculated by the CDC.</td>
</tr>
<tr>
<td>Patient Safety and Adverse Events (Composite)</td>
<td>Hospitals must report a minimum of three eligible cases on any one underlying indicator.</td>
</tr>
<tr>
<td>PC–01</td>
<td>Hospitals must report a minimum of 10 cases.</td>
</tr>
<tr>
<td><strong>Efficiency and Cost Reduction Domain</strong></td>
<td></td>
</tr>
<tr>
<td>MSPB</td>
<td>Hospitals must report a minimum number of 25 cases.</td>
</tr>
<tr>
<td>AMI Payment</td>
<td>Hospitals must report a minimum number of 25 cases.</td>
</tr>
<tr>
<td>HF Payment</td>
<td>Hospitals must report a minimum number of 25 cases.</td>
</tr>
<tr>
<td>PN Payment*</td>
<td>Hospitals must report a minimum number of 25 cases.</td>
</tr>
</tbody>
</table>

*As discussed in section V.I.3.b. of the preamble of this proposed rule, we are proposing to remove the current PSI 90 measure beginning with the FY 2019 program year. As discussed in section V.I.4.b. of the preamble of this proposed rule, we are proposing to adopt the Patient Safety and Adverse Events (Composite) measure beginning with the FY 2023 program year.

*As discussed in section V.I.4.a. of the preamble of this proposed rule, we are proposing to adopt the PN Payment measure beginning with the FY 2022 program year.

d. Weighting Measures Within the Efficiency and Cost Reduction Domain

In the FY 2012 IPPS/LTCH PPS final rule (76 FR 51618 through 51627), we adopted the MSPB measure for the Hospital VBP Program beginning with the FY 2015 program year. MSPB is the only cost measure in the Hospital VBP Program through the FY 2020 program year; as a result, hospitals’ Efficiency and Cost Reduction domain scores are currently based solely on their MSPB measure scores. In the FY 2017 IPPS/LTCH PPS final rule, we adopted two condition-specific cost measures, the AMI Payment and HF Payment measures, beginning with the FY 2021 program year (81 FR 56987 through 56990 and 81 FR 56990 through 56992, respectively). In addition, as discussed in section V.I.4.a. of the preamble of this proposed rule, we are proposing to adopt an additional condition-specific cost measure, the PN Payment measure, beginning with the FY 2022 program year. Based on this evolution of the Hospital VBP Program measure set, we believe it is appropriate to address measure score weighting within the Efficiency and Cost Reduction domain.

In determining how to weight measures in the Efficiency and Cost Reduction domain, we took into consideration hospitals’ experience with the measures and the measures’ ability to incentivize greater coordination among hospitals, physicians, and providers of post-acute care services to optimize the value of care they provide...
to Medicare beneficiaries. Therefore, we are proposing to weight the measures within the Efficiency and Cost Reduction domain such that the MSPB measure comprises 50 percent of a hospital’s domain score and the other condition-specific payment measures, weighed equally, comprise the remaining 50 percent of a hospital’s domain score, beginning with the FY 2021 program year and for subsequent years. We further are proposing that:

- If a hospital meets the case minimum to receive a score on the MSPB measure but does not meet the minimum number of cases for any other measures in the Efficiency and Cost Reduction domain, its domain score will be based solely on its MSPB score;
- If a hospital does not meet the case minimum to receive a score on the MSPB measure but meets the minimum number of cases for any other measure or measures within the Efficiency and Cost Reduction domain, its domain score will be based on its scores on the other payment measures, weighted equally (that is, the MSPB measure’s weight will be redistributed equally among the Efficiency and Cost Reduction domain measures for which the hospital is eligible receive a score); and
- If a hospital meets the case minimum to receive a score on the MSPB measure and one or more other measures within the Efficiency and Cost Reduction domain, but not all measures within this domain, the hospital’s MSPB measure score will comprise 50 percent of its domain score and the remaining 50 percent will be divided equally among the measures for which the hospital is eligible to receive a score.

Under our proposed weighting scheme, a hospital’s MSPB measure score could constitute between 12.5 percent and 25 percent of the hospital’s TPS. We believe the proposed weighting is appropriate because the MSPB measure is an overall spending measure and is therefore more broadly applicable than the condition-specific payment measures. In addition, hospitals have the most familiarity with this measure because it has been in the program the longest. We also considered proposing to weight all measures within the Efficiency and Cost Reduction domain equally. However, we determined this weighting may not reflect the broader applicability of the MSPB measure and its importance in ensuring that hospitals monitor the overall costs of care they provide to a larger subset of Medicare beneficiaries during an inpatient hospitalization and are involved in the coordination of beneficiaries’ care immediately prior to hospitalization and post-discharge.

We are inviting public comment on these proposals.

### K. Proposed Changes to the Hospital-Acquired Condition (HAC) Reduction Program

#### 1. Background

We refer readers to section V.I.1.a. of the FY 2014 IPPS/LTCH PPS final rule (78 FR 50707 through 50708) for a general overview of the HAC Reduction Program. For a detailed discussion of the statutory basis of the HAC Reduction Program, we refer readers to section V.I.2. of the FY 2014 IPPS/LTCH PPS final rule (78 FR 50708 through 50709). For a further description of our previously finalized policies for the HAC Reduction Program, we refer readers to the FY 2014 IPPS/LTCH PPS final rule (78 FR 50707 through 50729), the FY 2015 IPPS/LTCH PPS final rule (79 FR 50087 through 50104), the FY 2016 IPPS/LTCH PPS final rule (80 FR 49570 through 49581), and the FY 2017 IPPS/LTCH PPS final rule (81 FR 57011 through 57026). These policies describe the general framework for implementation of the HAC Reduction Program, including:

- (a) The relevant definitions applicable to the program;
- (b) the payment adjustment under the program;
- (c) the measure selection process and conditions for the program, including a risk-adjustment and scoring methodology;
- (d) performance scoring;
- (e) the process for making hospital-specific performance information available to the public, including the opportunity for a hospital to review the information and submit corrections; and
- (f) limitation of administrative and judicial review.

We also have codified certain requirements of the HAC Reduction Program at 42 CFR 412.170 through 412.172.

#### 2. Implementation of the HAC Reduction Program for FY 2018

In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50717), we finalized the following Centers for Disease Control and Prevention (CDC) National Healthcare Safety Network (NHSN) measures for Domain 2 for use in the FY 2015 program and subsequent years: CLABSI, CAUTI, Colon and Abdominal Hysterectomy SSI, MRSA Bacteremia, and CDI. In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57020), we finalized the use of the Patient Safety and Adverse Events Composite (PSI 90) measures for use in the FY 2018 program and subsequent years for Domain 1. These previously finalized measures are shown in the table below.

### HAC Reduction Program Measures for FY 2018

<table>
<thead>
<tr>
<th>Short name</th>
<th>Measure name</th>
<th>NQF #</th>
</tr>
</thead>
<tbody>
<tr>
<td>PSI 90</td>
<td>Patient Safety and Adverse Events Composite</td>
<td>0531</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Domain 1</th>
</tr>
</thead>
</table>

**Domain 2**

<table>
<thead>
<tr>
<th>Measure name</th>
<th>NQF #</th>
</tr>
</thead>
<tbody>
<tr>
<td>CAUTI</td>
<td>0138</td>
</tr>
<tr>
<td>CDI</td>
<td>1717</td>
</tr>
<tr>
<td>CLABSI</td>
<td>0139</td>
</tr>
<tr>
<td>Colon and Abdominal Hysterectomy SSI</td>
<td>0753</td>
</tr>
<tr>
<td>MRSA Bacteremia</td>
<td>1716</td>
</tr>
</tbody>
</table>

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57022), we finalized a 15-month performance period from July 1, 2014 through September 30, 2015, for the Domain 1 measure (PSI 90 Patient Safety and Adverse Events Composite) and a 24-month performance period from January 1, 2015 through December 31, 2016 (CYs 2015 and 2016) for...
Domain 2 measures (CLABSI, CAUTI, Colon and Abdominal Hysterectomy SSI, MRSA Bacteremia, and CDI) for the FY 2018 HAC Reduction Program. We anticipate we will be able to provide hospitals with their confidential hospital-specific reports and discharge level information used in the calculation of their FY 2018 Total HAC Score in late summer 2017 via the QualityNet Secure Portal. In order to access their hospital-specific reports, hospitals must register for a QualityNet Secure Portal account. We did not make any changes to the review and correction policies for FY 2017. Hospitals have a period of 30 days after the information is posted to the QualityNet Secure Portal to review and submit corrections for the calculation of their HAC Reduction Program measure scores, domain scores, and Total HAC Score for the fiscal year.

In this proposed rule, for the HAC Reduction Program, we are: (1) Proposing to specify the dates of the time period used to calculate hospital performance for the FY 2020 HAC Reduction Program; (2) requesting comments on additional measures for potential future adoption; (3) requesting comments on social risk factors; (4) requesting comments on accounting for disability and medical complexity in the CDC NHSN measures in Domain 2; and (5) proposing to update the Extraordinary Circumstance Exception policy beginning in FY 2018 as related to extraordinary circumstances that occur on or after October 1, 2017. These proposals are described in more detail below.

3. Proposed Data Collection Time Periods for the FY 2020 HAC Reduction Program

Section 1886(p)(4) of the Act gives the Secretary the statutory authority to determine the “applicable period” during which data are collected for the HAC Reduction Program. In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50717), we finalized and codified at 42 CFR 412.170 that we would use a 2-year time period of performance data to calculate the Total HAC Score. In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57020), we finalized a truncated data collection period for Domain 1, shorter than the previous 2-year data collection period for calculating the Total HAC Score for the FY 2018 and FY 2019 HAC Reduction Programs, in order to accommodate the transition to the ICD-10 classification system. We also changed the definition of “applicable period,” in 42 CFR 412.170, to reflect this change.

For the FY 2020 program, we are proposing to return to a two-year time period for the calculation of HAC Reduction Program measure results. We believe that using 2 years of data for both domains balances the needs of the program and allows for sufficient time to process the claims data and calculate the measure results. The 2-year time period allows time to complete the complex calculation process for the measures, to perform comprehensive quality assurance to enhance the accuracy of measure results, and to disseminate confidential reports on hospital-level results to individual hospitals. For the Domain 1 measure (Patient Safety and Adverse Events Composite), we are proposing to use the 24-month period from July 1, 2016 through June 30, 2018. The claims for all Medicare Fee-for-Service beneficiaries discharged during this period would be included in the calculations of measure results for Domain 1 for the FY 2020 program. For the CDC NHSN measures in Domain 2 (CLABSI, CAUTI, Colon and Abdominal Hysterectomy SSI, MRSA Bacteremia, and CDI), we are proposing to use data from CYs 2017 and 2018, that is January 1, 2017—December 31, 2018, for the FY 2020 program.

4. Request for Comments on Additional Measures for Potential Future Adoption

In the FY 2017 IPPS/LTCH PPS proposed rule (81 FR 25123), we welcomed public comment and suggestions for additional HAC Reduction Program measures. We believe that our continued efforts to reduce HACs are vital to improving patients’ quality of care and reducing complications and mortality, while simultaneously decreasing costs. The reduction of HACs is an important marker of quality of care and has a positive impact on both patient outcomes and cost of care. Our goal for the HAC Reduction Program is to heighten the awareness of HACs and reduce the number of incidences that occur.

As part of our ongoing efforts to evaluate and strengthen the HAC Reduction Program, we are conducting a review of patient safety measures to include in Domain 1. We seek to adopt outcomes-focused patient-safety measures that focus on topic areas including but not limited to: Falls with injury, adverse drug events (ADEs), glycemic events and ventilator associated events (VAEs). NQF identified these as gap areas for the HAC Reduction Program.

We note that falls are frequent in the inpatient setting. An estimated 700,000 to 1 million inpatients fall each year in U.S. hospitals. These falls can result in further health care complications for patients, and add costs by increasing the need for expensive imaging, like head computed-tomography scans. Risk assessment is the primary tool for preventing falls and research has indicated that inpatient fall prevention programs with patient education components are effective in reducing fall rates. ADEs are a frequent and preventable occurrence among hospital inpatients. They pose serious threats to patient safety and can result in prolonged hospitalization, increased morbidity and higher health care costs. Glycemic events, a common occurrence among inpatients, are associated with a greater risk of negative health outcomes. Many guidelines exist to support glycemic control in hospitalized patients. The most common guideline recommendations include documenting diabetes diagnosis, obtaining a hemoglobin A1C on admission, use of the “basal-bolus” method for insulin delivery, discontinuation of noninsulin agents for non-ICU patients with type 2 diabetes, and use of standardized order sets.
Mechanically ventilated patients are at greater risk for VAEs, which can result in morbidity and death. VAEs include ventilator associated pneumonia (VAP) and preventable adverse events, such as pulmonary edema and acute respiratory distress syndrome. VAP continues to rank among the most common HACs and effective prevention strategies for VAP include early removal of invasive devices and strict infection control and prevention efforts should target these high-risk groups.

Our overarching purpose is to support the National Quality Strategy’s three-part aim of better health care for individuals, better health for populations, and lower costs for health care. To the extent practicable, HAC Reduction Program measures should be nationally endorsed by a multi-stakeholder organization. Measures should be aligned with best practices among other payers and the needs of the end users of the measures. Measures should take into account widely accepted criteria established in medical literature.

We welcome public comments and suggestions on these measure areas, as well as additional outcome-based patient-safety measures that will help achieve the program goals.

5. Accounting for Social Risk Factors in the HAC Reduction Program

We understand that social risk factors such as income, education, race and ethnicity, employment, disability, community resources, and social support (certain factors of which are also sometimes referred to as socioeconomic status (SES) factors or socio-demographic status (SDS) factors) play a major role in health. One of our core objectives is to improve beneficiary outcomes, including reducing health disparities, and we want to ensure that all beneficiaries, including those with social risk factors, receive high quality care. In addition, we seek to ensure that the quality of care furnished by providers and suppliers is assessed as fairly as possible under our programs while ensuring that beneficiaries have adequate access to excellent care.

We have been reviewing reports prepared by the Office of the Assistant Secretary for Planning and Evaluation (ASPE) and the National Academies of Sciences, Engineering, and Medicine on the issue of accounting for social risk factors in CMS’ value-based purchasing and quality reporting programs, and considering options on how to address the issue in these programs. On December 21, 2016, ASPE submitted a report to Congress on a study it was required to conduct under section 2(d) of the Improving Medicare Post-Acute Care Transformation (IMPACT) Act of 2014. The study analyzed the effects of certain social risk factors in Medicare beneficiaries on quality measures and measures of resource use used in one or more of nine Medicare value-based purchasing programs, including the HAC Reduction Program. The report also included considerations for strategies to account for social risk factors in these programs. In a January 2017 report released by the National Academies of Sciences, Engineering, and Medicine, that body provided various potential methods for measuring and accounting for social risk factors, including stratified public reporting.

As noted in the FY 2017 IPPS/LTC PPS final rule, the NQF has undertaken a 2-year trial period in which certain new measures, measures undergoing maintenance review, and measures endorsed with the condition that they enter the trial period can be assessed to determine whether risk adjustment for selected social risk factors is appropriate for these measures. This trial entails temporarily allowing inclusion of social risk factors in the risk-adjustment approach for these measures. At the conclusion of the trial, NQF will issue recommendations on the future inclusion of social risk factors in risk adjustment for these quality measures, and we will closely review its findings.

We note that measures in the HAC Reduction Program, generally, represent never events, and are often preventable conditions like central line associated bloodstream infections, catheter associated urinary tract infections, and other complications or conditions that arise after a patient was admitted to the hospital for the treatment of another condition. We believe these events should not be influenced by social risk factors; instead, they are risk-adjusted for factors listed in specifications for the AHRQ and CDC. Currently, risk factors such as the patient’s age, gender, comorbidities, and complications are considered in the calculation of the measure rates so that they account for the clinical differences in the patients served by hospitals. Our measures continually undergo maintenance to determine the need for updated specifications, and to monitor for trends and any relevant risk-adjustment changes needed for the measures. We remind readers that, beginning for payments made in FY 2018, we adopted the modified PSI 90: Patient Safety and Adverse Events Composite (NQF #0531); the composite was revised to reflect the relative importance and harm associated with each component indicator, and to provide a more reliable and valid signal of patient safety events (81 FR 57020).

We also adopted a continuous scoring approach in the HAC Reduction Program that brings our scoring domains into alignment each other, essentially eliminates ties in Total HAC Scores, reduces effects on outliers, and enhances the ability to distinguish among hospitals of varying quality (81 FR 57025).

As we continue to consider the analyses and recommendations from these reports and await the results of the NQF trial on risk adjustment for quality measures, we are continuing to work with stakeholders in this process. As we have previously communicated, we are concerned about holding providers to different standards for the outcomes of their patients with social risk factors because we do not want to mask potential disparities or minimize incentives to improve the outcomes for disadvantaged populations. Keeping this concern in mind, while we sought input on this topic previously, we
continue to seek public comment on whether we should account for social risk factors in the HAC Reduction Program, and if so, what method or combination of methods would be most appropriate for accounting for social risk factors. Examples of methods include: Adjustment of the payment adjustment methodology under the HAC Reduction Program; adjustment of provider performance scores (for instance, stratifying providers based on the proportion of their patients who are dual eligible); confidential reporting of stratified measure rates to providers; public reporting of stratified measure rates; risk adjustment of a particular measure as appropriate based on data and evidence; and redesigning payment incentives (for instance, rewarding improvement for providers caring for patients with social risk factors or incentivizing providers to achieve health equity).

We note that in section V.I.9. of the preamble of this proposed rule, we discuss considerations for stratifying hospitals into peer groups for purposes of assessing payment adjustments under the Hospital Readmissions Reduction Program, as required under the 21st Century Cures Act. We refer readers to that section for a detailed discussion of these alternatives; while this discussion and corresponding proposal are specific to the Hospital Readmissions Reduction Program, they reflect the level of analysis we would undertake when evaluating methods and combinations of methods for accounting for social risk factors in CMS’ other value-based purchasing programs, such as the HAC Reduction Program. While we consider whether and to what extent we currently have statutory authority to implement one or more of the above-described methods, we are seeking comments on whether any of these methods should be considered, and if so, which of these methods or combination of methods would best account for social risk factors in the HAC Reduction Program. In addition, we are also seeking public comment on which social risk factors might be most appropriate for stratifying measure scores and/or potential risk adjustment of a particular measure. Examples of social risk factors include, but are not limited to, dual eligibility/low-income subsidy, race and ethnicity, and geographic area of residence. We are seeking comments on which of these factors, including current data sources where this information would be available, could be used alone or in combination, and whether other data should be collected to better capture the effects of social risk. We will take commenters’ input into consideration as we continue to assess the appropriateness and feasibility of accounting for social risk factors in the HAC Reduction Program. We note that any such changes would be proposed through future notice-and-comment rulemaking.

We look forward to working with stakeholders as we consider the issue of accounting for social risk factors and reducing health disparities in CMS programs. Of note, implementing any of the above methods would be taken into consideration in the context of how this and other CMS programs operate (for example, data submission methods, availability of data, statistical considerations relating to reliability of data calculations, among others), so we also welcome comment on operational considerations. CMS is committed to ensuring that its beneficiaries have access to and receive excellent care, and that the quality of care furnished by providers and suppliers is assessed fairly in CMS programs.

6. Request for Comments on Inclusion of Disability and Medical Complexity for CDC NHSN Measures

The intent of the HAC Reduction Program is to encourage all hospitals to reduce the incidence of HACs. We continue to believe that there is room for improvement in the incidence of HACs, regardless of the institution or hospital. The measures adopted in the HAC Reduction Program, which are risk-adjusted to account for the different patient populations that hospitals serve, target important quality improvement areas. In its IMPACT Act report, ASPE suggested payment strategies to improve the HAC Reduction Program. ASPE noted that it is well-proven that higher levels of medical risk are associated with a higher risk for many (although not all) patient safety events, particularly infections. For example, diabetes is associated with roughly 70 percent higher odds of surgical site infections and diabetes, pulmonary disease, renal failure, and exposure to nursing homes are associated with a higher risk of MRSA. Many of the same medical factors also confer a higher risk of C. diff. infection, as well as CAUTI and CLABSI.

ASPE suggested that patient-level clinical data from the CDC healthcare-associated infection (HAI) measures should be examined and considered for additional risk adjustment. ASPE also noted that the clinical risk-adjustment of the patient safety and hospital-acquired infection measures should be improved to ensure the measures adequately adjust for differences in patients’ clinical risk, so that fair comparisons for hospital accountability and performance assessment can be made to hold providers to the same fair standard. ASPE recommended additional analyses for measure developers such as AHRQ and CDC to determine whether adjusting key components of the patient safety or HAI measures (for example frailty, functional limitations, prior hospitalizations or nursing home residence, or other markers of immune system deficiencies or unmeasured medical complexity) may better account for susceptibility to infection and patient safety events.

Based on ASPE’s analysis and considerations, we are requesting stakeholder feedback on risk-adjusting the CDC NHSN measures for disability or medical complexity. Although we are not proposing any specific changes to the measures at this time, we will consider all comments as a guide to potential future action.

7. Maintenance of Technical Specifications for Quality Measures

Technical specifications for Patient Safety and Adverse Events Composite Measure in Domain 1 can be found at AHRQ’s Web site at: http://qualityindicators.ahrq.gov/Modules/PSI_TechSpec.aspx. Technical specifications for the CDC NHSN HAI measures in Domain 2 can be found at CDC’s NHSN Web site at: http://www.cdc.gov/nhsn/acute-care-hospital/index.html. Both Web sites provide measure updates and other information necessary to guide hospitals participating in the collection of HAC Reduction Program data.

In the FY 2015 IPPS/LTCH PPS final rule (79 FR 50100), we described a policy under which we use a subregulatory process to make nonsubstantive updates to measures used for the HAC Reduction Program. We are not proposing any changes to this policy at this time.
8. Extraordinary Circumstance Exception (ECE) Policy for the HAC Reduction Program

Many of our quality reporting and value-based purchasing programs share a common process for requesting an exception from program reporting due to an extraordinary circumstance not within a provider’s control. The Hospital IQR, Hospital OQR, IPFQR, ASCQR, and PCHQR Programs, as well as the Hospital Readmissions Reduction Program, share common processes for Extraordinary Circumstance Exception (ECE) requests. In reviewing the policies for these programs, we recognized that there are five areas in which these programs have variance regarding ECE requests. These are: (1) Allowing the facilities or hospitals to submit a form signed by the facility’s or hospital’s CEO versus CEO or designated personnel; (2) requiring the form be submitted within 30 days following the date that the extraordinary circumstance occurred versus within 90 days following the date the extraordinary circumstance occurred; (3) inconsistency regarding specification of a timeline for us to provide our formal response notifying the facility or hospital of our decision; (4) inconsistency regarding specification of our authority to grant ECEs due to CMS data system issues; and (5) referring to the program as “extraordinary extensions/exemptions” versus as “extraordinary circumstances exceptions.” We believe addressing these five areas, as appropriate, can improve administrative efficiencies for affected facilities or hospitals.

In the FY 2016 IPPS/LTCH PPS final rule (80 FR 49579 through 49581), we adopted an ECE policy for the HAC Reduction Program beginning in FY 2016. This policy was similar to the ECE policy for the Hospital IQR Program, as finalized in the FY 2012 IPPS/LTCH PPS final rule (76 FR 51651), modified in the FY 2014 IPPS/LTCH PPS final rule (79 FR 50836) (designation of a non-CEO hospital contact), and further modified in the FY 2015 IPPS/LTCH PPS final rule (79 FR 50277) (amended 42 CFR 412.140(c)(2) to refer to “extension or exemption” instead of the former “extension or waiver”). In section IX.A.15. of the preamble of this proposed rule, we are proposing to amend the Hospital IQR Program regulations at 42 CFR 412.140(c)(2) to refer to “extraordinary circumstances exceptions” and we will continue to use this nomenclature for the HAC Reduction Program.

We are proposing to modify the ECE policy for the HAC Reduction Program by: (1) Allowing the facility to submit a form signed by the facility’s CEO or designated personnel; (2) specifying that we will strive to provide our formal response notifying the facility of our decision within 90 days of receipt of the facility’s request; and (3) specifying that CMS may grant ECEs due to CMS data system issues which affect data submission. These proposed modifications generally align with policies in the Hospital IQR Program (76 FR 51651 through 51652; 78 FR 50836 through 50837; and 81 FR 57181 through 57182), the Hospital OQR Program (77 FR 68489 and 81 FR 79795), as well as other quality reporting programs. We are proposing that these modifications would apply beginning in FY 2018 as related to extraordinary circumstances that occur on or after October 1, 2017.

We note that there may be circumstances in which it is not feasible for a facility’s CEO to sign the ECE request form. In these circumstances, we believe that facilities affected by such circumstances should be able to submit ECE forms regardless of the CEO’s availability to sign. Therefore, the first proposed modification would allow any hospital to designate an appropriate, non-CEO, contact at its discretion. This individual would be responsible for the submission, and would be the one signing the form. We would accept ECE forms which have been signed by designated personnel.

We also believe that it is important for facilities to receive timely feedback regarding the status of ECE requests. We strive to complete our review of each ECE request as quickly as possible. However, we recognize that the number of requests we receive, and the complexity of the information provided impacts the actual timeframe to make ECE determinations. To improve transparency of our process, we believe it is appropriate to clarify that we will strive to provide our response within 90 days of receipt.

Although we do not anticipate this situation will happen on a regular basis, there may be times where CMS experiences issues with its data systems that directly affects facilities’ abilities to submit data. In these cases, we believe it would be inequitable to require facilities to report. Therefore, we are proposing to allow CMS to grant ECEs to facilities if we determine that a systemic problem with one of our data collection systems directly affected the ability of the facilities to submit data. If we make the determination to grant ECEs, we are proposing to communicate this decision through routine communication channels.

We are inviting public comment on these proposed modifications to the HAC Reduction Program’s ECE policy.

L. Rural Community Hospital Demonstration Program

1. Introduction

The Rural Community Hospital Demonstration was originally authorized for a 3-year period by section 410A of the Medicare Prescription Drug, Improvement, and Modernization Act of 2003 (MMMA) (Pub. L. 108–173), and extended for another 5-year period by sections 3123 and 10313 of the Affordable Care Act (Pub. L. 111–148). Subsequently, section 15003 of the 21st Century Cures Act (Pub. L. 114–255), enacted December 13, 2016, amended section 410A of Public Law 108–173 to require a 10-year extension period (in place of the 5-year extension required by the Affordable Care Act, as further discussed below). Section 15003 also requires that no later than 120 days after enactment of Public Law 114–255 the Secretary issue a solicitation for applications to select additional hospitals to participate in the demonstration program for the second 5 years of the 10-year extension period so long as the maximum number of 30 hospitals stipulated by the Affordable Care Act is not exceeded. In this proposed rule, we provide a summary of the previous legislative provisions and their implementation; a description of the provisions of section 15003 of Public Law 114–255 and our proposals for implementation; and our proposals for budget neutrality, including a discussion of the budget neutrality methodology used in previous final rules, the proposed budget neutrality methodology for the extension period authorized by section 15003 of Public Law 114–255, and the proposed reconciliation of actual and estimated costs of the demonstration for previous years (2011, 2012, and 2013).

2. Background

Section 410A(a) of Public Law 108–173 required the Secretary to establish a demonstration program to test the feasibility and advisability of establishing “rural community” hospitals to furnish covered inpatient hospital services to Medicare beneficiaries. The demonstration pays rural community hospitals under a reasonable cost-based methodology for Medicare payment purposes for covered inpatient hospital services furnished to Medicare beneficiaries. A rural community hospital, as defined in section 410A(f)(1), is a hospital that—
• Is located in a rural area (as defined in section 1886(d)(2)(D) of the Act) or is treated as being located in a rural area under section 1886(d)(6)(C) of the Act;
• Has fewer than 51 beds (excluding beds in a distinct part psychiatric or rehabilitation unit) as reported in its most recent cost report;
• Provides 24-hour emergency care services; and
• Is not designated or eligible for designation as a CAH under section 1820 of the Act.

Section 410A(a)(4) of Public Law 108–173 specified that the Secretary was to select for participation no more than 15 rural community hospitals in rural areas of States that the Secretary identified as having low population densities. Using 2002 data from the U.S. Census Bureau, we identified the 10 States with the lowest population density in which rural community hospitals were to be located in order to participate in the demonstration: Alaska, Idaho, Montana, Nebraska, Nevada, New Mexico, North Dakota, South Dakota, Utah, and Wyoming (Source: U.S. Census Bureau, Statistical Abstract of the United States: 2003).

CMS originally solicited applicants for the demonstration in May 2004; 13 hospitals began participation with cost reporting periods beginning on or after October 1, 2004. In 2005, 4 of these 13 hospitals withdrew from the demonstration program and converted to CAH status. This left 9 hospitals participating at that time. In 2008, we announced a solicitation for up to 6 additional hospitals to participate in the demonstration program. Four additional hospitals were selected to participate under this solicitation. These 4 additional hospitals began under the demonstration payment methodology with the hospitals’ first cost reporting period starting on or after July 1, 2008. At that time, 13 hospitals were participating in the demonstration.

Five hospitals (3 of the hospitals were among the 13 hospitals that were original participants in the demonstration program and 2 of the hospitals were among the 4 hospitals that began the demonstration program in 2008) withdrew from the demonstration program during CYs 2009 and 2010. In CY 2011, one hospital that was among the original set of hospitals that participated in the demonstration withdrew from the demonstration. These actions left 7 of the originally participating hospitals (that is, hospitals that were selected to participate in either 2004 or 2008) participating in the demonstration program as of June 1, 2011.

Sections 3123 and 10313 of the Affordable Care Act (Pub. L. 111–148) amended section 410A of Public Law 108–173, changing the rural community hospital demonstration program in several ways. First, the Secretary was required to conduct the demonstration program for an additional 5-year period, to begin on the date immediately following the last day of the initial 5-year period. Further, the Affordable Care Act required, in the case of a rural community hospital participating in the demonstration program as of the last day of the initial 5-year period, the Secretary to provide for the continued participation of such rural community hospital in the demonstration program during the 5-year extension period, unless the hospital made an election to discontinue participation.

In addition, the Affordable Care Act required that, during the 5-year extension period, the Secretary shall expand the number of States with low population densities determined by the Secretary to 20. Further, the Secretary was required to use the same criteria and data that the Secretary used to determine the States for purposes of the initial 5-year period. The Affordable Care Act also allowed not more than 30 rural community hospitals in such States to participate in the demonstration program during the 5-year extension period.

We published a solicitation for applications for additional participants in the Rural Community Hospital Demonstration program in the Federal Register on August 30, 2010 (75 FR 52960). The 20 States with the lowest population density that were eligible for the demonstration program were: Alaska, Arizona, Arkansas, Colorado, Idaho, Iowa, Kansas, Maine, Minnesota, Mississippi, Montana, Nebraska, Nevada, New Mexico, North Dakota, Oklahoma, Oregon, South Dakota, Utah, and Wyoming (Source: U.S. Census Bureau, Statistical Abstract of the United States: 2003). Sixteen new hospitals began participation in the demonstration with the first cost reporting period beginning on or after April 1, 2011.

In addition to the 7 hospitals that were selected in either 2004 or 2008, the new selection led to a total of 23 hospitals in the demonstration. During CY 2013, one additional hospital among the set selected in 2011 withdrew from the demonstration, which left 22 hospitals participating in the demonstration, effective July 1, 2013, all of which continued their participation through CY 2016. Starting from that date and extending through the end of FY 2015, the 7 “originally participating” hospitals, that is, hospitals that were selected in either 2004 or 2008, ended on a rolling basis their scheduled 5-year periods of performance authorized by the Affordable Care Act (referred to hereafter as “Cohort 1” hospitals).

Likewise, the participation period for the 14 hospitals that entered the demonstration following the mandate of the Affordable Care Act and that were still participating (referred to as “Cohort 2” hospitals) ended their scheduled periods of performance on a rolling basis according to the end dates of the hospitals’ cost report periods, respectively, from April 30, 2016 through December 31, 2016. (One hospital among the Cohort 2 hospitals closed in October 2015.)


As stated earlier, section 15003 of Public Law 114–255 further amended section 410A of Public Law 108–173 to require the Secretary to conduct the Rural Community Hospital Demonstration for a 10-year extension period (in place of the 5-year extension period required by the Affordable Care Act), beginning on the date immediately following the last day of the initial 5-year period under section 410A(a)(5) of Public Law 108–173. Thus, the Secretary is required to conduct the demonstration for an additional 5-year period. Specifically, section 15003 of Public Law 114–255 amended section 410A(g)(4) of Public Law 108–173 to require that, for hospitals participating in the demonstration as of the last day of the initial 5-year period, the Secretary shall provide for continued participation of such rural community hospitals in the demonstration during the 10-year extension period, unless the hospital makes an election, in such form and manner as the Secretary may specify, to discontinue participation. In addition, section 15003 of Public Law 114–255 amended section 410A(g)(4)(B) (and thereby (b)(1)(A)) of Public Law 108–173 to provide that in calculating the amount of payment under the demonstration program to the rural community hospital for covered inpatient hospital services furnished by the hospital during each 5-year period of such 10 year extension period, the amount of payment (for the first cost reporting period) is the reasonable costs of providing such services for discharges occurring in the first cost reporting period beginning on or after the first day of each applicable 5-year period.
period in the 10-year extension period. Furthermore, section 15003 of Public Law 114–255 added subsection (g)(5) to section 410A of Public Law 108–173 to require that, during the second 5 years of the 10-year extension period, the Secretary shall apply the provisions of section 410A(g)(4) of Public Law 108–173 to rural community hospitals that are not described in subsection (g)(4) but that were participating in the demonstration as of December 30, 2014, in a similar manner as such provisions apply to hospitals described in subsection (g)(4). We interpret this as providing for participation in and payment under the demonstration during the second 5 years of the 10 year extension period for hospitals that are not described in section 410A(g)(4) of Public Law 108–173 (as amended) but that were participating in the demonstration as of December 30, 2014, in a similar manner as such extension and payment applies to hospitals described in section 410A(g)(4) of Public Law 108–173 (as amended), unless a hospital makes an election, in such form and manner as the Secretary may specify, to discontinue participation.

In addition, section 15003 of Public Law 114–255 amended section 410A of Public Law 108–173 to add paragraph (g)(6)(A) which requires that, no later than 120 days after enactment of paragraph (g)(6), the Secretary shall issue a solicitation for applications to select additional rural community hospitals located in any State to participate in the demonstration program for the second 5 years of the 10-year extension period, without exceeding the maximum number of hospitals (that is, 30) permitted under section 410A(g)(3) of Public Law 108–173 (which was added by the Affordable Care Act). Section 15003 also amended section 410A of Public Law 108–173 to add paragraph (g)(6)(B) which provides that, in determining which hospitals submitting an application pursuant to this solicitation are to be selected for participation in the demonstration, the Secretary shall give priority to rural community hospitals located in one of the 20 States with the lowest population densities, as determined using the 2015 Statistical Abstract of the United States. In addition, in determining which among the hospitals submitting an application pursuant to this solicitation are to be selected for participation in the demonstration, section 410A(g)(6)(B) specifies that the Secretary may consider closures of hospitals located in rural areas in the State in which an applicant hospital is located during the 5-year period immediately preceding the date of enactment of section 410A(g)(6) of Public Law 108–173, as well as the population density of the State in which the rural community hospital is located.

b. Proposed Terms of Continuation for Previously Participating Hospitals

As discussed earlier, section 15003 of Public Law 114–255 (the 21st Century Cures Act) amended section 410A of Public Law 108–173 to provide for a 10-year extension of the demonstration (in place of the 5-year extension required by the Affordable Care Act) beginning on the date immediately following the last day of the initial 5-year period under section 410A(a)(5) of Public Law 108–173. Thus, section 15003 of Public Law 114–255 requires an additional 5-year extension of the demonstration beyond the extension required by the Affordable Care Act. Given the timing of the enactment of Public Law 114–255, for most of the previously participating hospitals, there is a gap between the end date of each hospital’s participation in the first 5-year extension period and enactment of Public Law 114–255 on December 13, 2016. For these hospitals, this gap is for a period of between 2 to 23 months. Section 15003 of Public Law 114–255 does not address how the second 5 years of the 10-year extension is to be implemented in the event of a gap between the end of the first 5 years of the 10-year extension period for a participating hospital and the enactment of Public Law 114–255 authorizing the second 5 years of the 10-year extension period. Given this gap and the lack of specific direction in the statute regarding how to implement the extension in this situation for these previously participating hospitals, and the mandate under section 15003 of Public Law 114–255 to issue a solicitation for additional participants for the second 5 years of the 10 year extension, we considered how to implement the second 5 years of the 10-year extension period for the reasons discussed below. We believe that it would be reasonable and consistent with the statute to implement the second 5 years of the 10-year extension period in a way that recognizes a gap in participation for the previously participating hospitals between the end of the first 5 years and the start of the second 5 years of the extension period, and that provides for alignment of the periods of performance under the extension among all participating hospitals. Thus, for each previously participating hospital that decides to participate in the second 5 years of the 10-year extension period, we are proposing that the start date for the period of performance under the second 5-year extension period would be the start of the first cost reporting period on or after October 1, 2017 following upon the announcement of the selection of the additional hospitals for the demonstration. In this manner, we are proposing to align the periods of performance for the previously participating hospitals that decide to participate in the second 5-year extension period with the periods of performance for the additional hospitals authorized by section 15003 of Public Law 114–255. An additional reason for our proposal that the second 5-year period of performance start no earlier than October 1, 2017 for any of the hospitals (previously participating or newly selected) is to align the start of the periods of performance with FY 2018 for purposes of estimating the costs of the demonstration and thus determining the budget neutrality offset amount for the demonstration (discussed later in this section) for FY 2018. (The FY 2018 IPPS/LTCH PPS final rule is effective October 1, 2017.)

We believe the approach we are proposing above is consistent with section 410A of Public Law 108–173, as amended by Public Law 114–255. As discussed earlier, the statutory language does not specifically address how the 10-year extension period is to be implemented in the event of a gap between the end of the first 5-year extension of the demonstration for a participating hospital and the enactment of Public Law 114–255 authorizing the second 5-year extension of the demonstration. Furthermore, we believe that the payment methodology set forth in section 410A(b)(1)(A) and (g)(4) of Public Law 108–173, as amended by section 15003 of Public Law 114–255, contemplates that the first 5 years and the second 5 years of the 10-year extension period be treated as separate periods, in that, as discussed above, it provides for payment of reasonable costs for discharges occurring in the first cost reporting period beginning on or after the first day of “each applicable 5-year period in the 10-year extension period.” We believe that our proposed approach, which provides for a gap in participation between the end of the first 5 years and the start of the second 5 years of the 10-year extension period, is reasonable, given that most of the hospitals that participated in the first 5-year extension under the Affordable Care Act had already ended their participation under the demonstration when Public Law 114–255 was enacted, and that all hospitals now have been...
paid under other payment methodologies outside the demonstration for a significant period of time (anywhere from 3 months to more than 2 years as of the publication of this proposed rule).

In addition, we note that certain types of administrative actions are generally required in order to implement a demonstration program that administers Medicare payment according to a methodology that differs from the methodology that would otherwise apply under the statute. These include development of participation agreements, formulating direction to MACs, and the procurement of audit and evaluation contracts. We believe that implementing the second 5-year extension for each participating hospital beginning with the start of its first cost reporting period on or after October 1, 2017, following upon the announcement of the selection of the additional hospitals for the demonstration, gives us the time necessary to implement such administrative actions.

Furthermore, we believe that it is reasonable and preferable to provide, to the extent possible, for alignment of the periods of participation of the previously participating hospitals with any newly selected hospitals during the second 5 years of the 10-year extension period. Under our proposed implementation approach, all previously participating hospitals would begin their periods of performance under the 5-year extension in FY 2018 on the same basis as the newly selected hospitals (the start of the first cost reporting period beginning on or after October 1, 2017, following upon the announcement of the selection of the new hospitals). We believe that aligning the participation periods for all hospitals in this manner would be more conducive to testing the feasibility and advisability of the payment methodology required by section 410A of Public Law 108–173 because, for all hospitals, the demonstration payment methodology would be applicable and its effect evaluated for similar time periods. In addition, we believe our proposed approach would allow for streamlined and administratively feasible budget neutrality calculations for the second 5-year extension period because the costs of the demonstration would be estimated for periods of performance beginning in the same fiscal year.

We are inviting public comments on the proposed approach discussed above for implementing the second 5-year period of the 10-year extension required under section 15003 of Public Law 114–255 for the previously participating hospitals. In addition, we are inviting public comments on alternative approaches under the statute for implementing the extension, particularly with respect to the commencement of the second 5-year period of the extension for previously participating hospitals.

One potential alternative approach that we considered is for each previously participating hospital to begin the second 5 years of the 10-year extension period and the cost-based payment methodology under section 410A of Public Law 108–173 (as amended by section 15003 of Pub. L. 114–255), on the date immediately after the date of the period of performance under the first 5-year extension period ended. For example, for a hospital whose 5-year period of performance authorized by the Affordable Care Act ended June 30, 2015, the extension period under section 15003 of Public Law 114–255 would be effective July 1, 2015, and it would extend through June 30, 2020. Likewise, for a hospital whose 5-year period of performance ended June 30, 2016, the extension period under section 15003 of Public Law 114–255 would be effective July 1, 2016, and it would extend through June 30, 2021. The methodology we considered for calculating the budget neutrality offset amount under this alternative approach is described in section V.I.4.d. of the preamble of this proposed rule. Although we believe that this alternative approach would also be consistent with the language of section 410A of Public Law 108–173 (as amended) and, unlike the proposed approach, would not provide for a gap in the reasonable cost payment methodology between the end of the first and start of the second 5-year periods of the 10-year extension period, for the reasons discussed below, we believe that our proposed approach outlined above would be more appropriate. First, we note that applying the extension in this alternative manner would result in hospitals being paid under the cost-based payment methodology provided for under section 410A of Public Law 108–173 (as amended) for a period of time during which the hospitals were not actively participating in the demonstration. We believe that it would be more appropriate to conduct both the implementation and evaluation of the demonstration for a period of time for which the hospitals have actively agreed to participate.

Furthermore, we note that applying the demonstration payment methodology starting at the end of each previously participating hospital’s participation in the first 5-year extension period under the Affordable Care Act (as far back as cost reporting years beginning in FY 2015), in addition to implementing a new selection of hospitals, is likely to create a situation whereby the periods of participation for demonstration hospitals under the new extension period would be starting across 4 different fiscal years, because hospitals could have periods of performance that start as early as January 1, 2015, and as late as July 1, 2018. We believe that such a structure for the demonstration would not be as conducive to the goal of testing the feasibility and advisability of the cost-based payment methodology under section 410A of Public Law 108–173, as amended. Implementing a payment methodology that is different from that which would otherwise apply under the statute requires coordination among MACs and audit, quality monitoring, and evaluation contractors.

Administering the second 5-year extension period so that the extension begins over a span of time to include several years would add substantial complexity to these contractual arrangements. In addition, methodologies for evaluating the effects of a payment methodology enacted under a demonstration program often involve examination of the experience of nonparticipating providers. Conducting such an analysis over different time periods might reduce the usefulness of such an evaluation approach because metrics assessed in relation to participating hospitals and nonparticipating hospitals, respectively, would not apply to uniform time periods.

Nevertheless, we are seeking public comments on this alternative approach to implementing the extension to the demonstration under section 15003 of Public Law 114–255 and the corresponding alternative budget neutrality calculation described in section V.I.4.d. of the preamble of this proposed rule.

c. Solicitation for Additional Participants

As required under section 15003 of Public Law 114–255, we will issue a solicitation for additional hospitals to participate in the demonstration. We expect that this solicitation will be released in April 2017, and eligible hospitals will have 30 days to submit applications. Among other things, the solicitation will ask hospitals to describe challenges experienced with the current method of Medicare payment, the impact of rural hospital closures within the State or surrounding
area, and a strategy for financial viability and improving the health care of the population.

Section 15003 of Public Law 114–255 provides that, in determining which rural community hospitals that submitted an application pursuant to the solicitation under subparagraph (A) to select for participation in the demonstration program, the Secretary shall give priority to rural community hospitals located in one of the 20 States with the lowest population densities (as determined by the Secretary using the 2015 Statistical Abstract of the United States). We note that the U.S. Census Bureau ceased publishing the Statistical Abstract of the United States in 2011, and that in the years since then, ProQuest, LLC, a private vendor, has produced a volume intended to serve the same function as a comprehensive collection of national statistics, compiling data from different sources including published reports from the Census Bureau. Thus, we are using ProQuest Statistical Abstract of the United States, 2015 in determining which States to give priority in selecting additional participants for the demonstration. We believe that in the absence of a volume produced by the Census Bureau, using this compendium is consistent with the intent of the statute, and is appropriate for the purpose of designating States to which priority is to be given under section 410A(a)(6)(B)(i) of Public Law 108–173.


Consistent with our policy for the previous solicitations, we are choosing the more recent data source to identify the 20 States to which priority is to be given. These States are: Alaska, Arizona, Arkansas, Colorado, Idaho, Iowa, Kansas, Maine, Mississippi, Montana, Nebraska, Nevada, New Mexico, North Dakota, Oklahoma, Oregon, South Dakota, Utah, Vermont, and Wyoming.

We note that section 410A(a)(6)(B)(ii)(I) of Public Law 108–173 as added by section 15003 of Public Law 114–255 also states that, in selecting additional participants, the Secretary may consider the population density of the State in which the rural community hospital is located. As a result, we will consider the population density of the State in which the hospital is located. We believe that this consideration is reasonable given that the demonstration may have differing effects for health care services and populations depending on State population density. In addition, as permitted by section 410A(g)(6)(B)(ii)(I) of Public Law 108–173, in selecting additional participants under this solicitation, we will consider the impact of closures of hospitals located in rural areas in the State in which the hospital is located during the 5-year period immediately preceding the date of enactment of this paragraph. We believe that this consideration is reasonable, given the possibility that enhanced Medicare payment through the demonstration may increase access to health care services for populations thus affected by hospital closures.

Our goal is to finalize this selection by June 2017, in time to include in the FY 2018 IPPS final rule an estimate of the costs of the demonstration during FY 2018 and the resulting budget neutrality offset amount for these newly participating hospitals (referred to as “Cohort 3” hospitals), as well as for those hospitals among the previously participating hospitals that decide to participate in the extension period (Cohorts 1 and 2 hospitals). Upon announcing the selection of new participants, we will confirm the start dates for the periods of performance for these newly selected hospitals. In accordance with our proposed implementation approach discussed in section V.L.3.b. of the preamble of this proposed rule, if the selection is announced by June 2017, we expect that we would determine the periods of performance for all of the participating hospitals to begin with the first cost reporting period on or after October 1, 2017, and we would include an estimate of the costs for the demonstration for FY 2018 for Cohorts 1, 2, and 3 hospitals in the FY 2018 final rule. As previously discussed, under our proposal, the costs of the demonstration during FY 2018 for Cohorts 1 and 2 hospitals would not start earlier than October 1, 2017.

If final selection of the Cohort 3 hospitals does not occur by June 2017, under our proposed approach, we would not be able to include an estimate of the costs of the demonstration or an estimate of the budget neutrality offset amount for either these Cohort 3 hospitals or the previously participating Cohorts 1 and 2 hospitals in the FY 2018 IPPS final rule. Considering that periods of performance for the Cohorts 1 and 2 hospitals would not be determined until after the selection of the Cohort 3 hospitals, we would not know precisely when the periods of performance would begin for the Cohorts 1 and 2 hospitals, or to what extent they would overlap with the 12 months in FY 2018 until the Cohort 3 hospitals are selected. Therefore, if the announcement of the final selection of the Cohort 3 hospitals does not occur by June 2017, we would not be able to include an estimate of the demonstration costs or budget neutrality offset amount for FY 2018 for the Cohorts 1 and 2 hospitals in the FY 2018 IPPS final rule. As a result, if the announcement of the final selection of the Cohort 3 hospitals does not occur by June 2017, we would specify the dates on which all participating hospitals would start in the second 5 years of the 10-year extension period at the time the selection is announced in accordance with our proposal. We are proposing that if the selection of the Cohort 3 hospitals is not announced in June 2017, we would include the estimated costs of the demonstration for all participating hospitals for FY 2018 in the budget neutrality offset amount to be calculated in the FY 2019 IPPS/LTCH PPS proposed and final rules.

According to our proposal, regardless of whether the final selection of the Cohort 3 hospitals occurs by June 2017, no period of performance in the second 5 years of the 10-year extension period for any of the hospitals (Cohorts 1, 2, and 3) would start earlier than October 1, 2017. Our goal is, to the greatest extent possible, to align the start of the periods of performance with FY 2018 for purposes of estimating the costs of the demonstration and thus determining the budget neutrality offset amount for FY 2018. (We refer readers to section V.L.4. of the preamble of this proposed rule for our proposed calculation methodology for the budget neutrality offset amount for FY 2018.)

4. Budget Neutrality Requirement

Section 410A(c)(2) of Public Law 108–173 requires that, in conducting the demonstration program under this section, the Secretary shall ensure that the aggregate payments made by the Secretary do not exceed the amount which the Secretary would have paid if the demonstration program under this section was not implemented. This requirement is commonly referred to as “budget neutrality.” Generally, when
we implement a demonstration program on a budget neutral basis, the demonstration program is budget neutral on its own terms; in other words, the aggregate payments to the participating hospitals do not exceed the amount that would be paid to those same hospitals in the absence of the demonstration program. Typically, this form of budget neutrality is viable when, by changing payments or aligning incentives to improve overall efficiency, or both, a demonstration program may reduce the use of some services or eliminate the need for others, resulting in reduced expenditures for the demonstration program’s participants. These reduced expenditures offset increased payments elsewhere under the demonstration program, thus ensuring that the demonstration program as a whole is budget neutral or yields savings. However, the small scale of this demonstration program, in conjunction with the payment methodology, makes it extremely unlikely that this demonstration program could be viable under the usual form of budget neutrality—that is, cost-based payments to participating small rural hospitals are likely to increase Medicare outlays without producing any offsetting reduction in Medicare expenditures elsewhere. In addition, a rural community hospital’s participation in this demonstration program would be unlikely to yield benefits to the participants if budget neutrality were to be implemented by reducing other payments for these same hospitals. Therefore, in the 12 IPPS final rules spanning the period from FY 2005 through FY 2016, we adjusted the national inpatient PPS rates by an amount sufficient to account for the added costs of this demonstration program, thus applying budget neutrality across the payment system as a whole rather than merely across the participants in the demonstration program. (In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57034), we described a different methodology which we specify below.) As we discussed in the FYs 2005 through 2017 IPPS final rules (69 FR 49183; 70 FR 47462; 71 FR 48100; 72 FR 47392; 73 FR 48670; 74 FR 43922, 75 FR 50343, 76 FR 51698, 77 FR 53449, 78 FR 50740, 77 FR 50145; 80 FR 49585; and 81 FR 57034, respectively), we believe that the language of the statutory budget neutrality requirements permits the agency to implement the budget neutrality provision in this manner.

b. Methodology Used in Previous Final Rules

We generally incorporated two components into the budget neutrality offset amounts identified in the final IPPS rules in previous years. First, we estimated the costs of the demonstration for the upcoming fiscal year, generally determined from historical, “as submitted” cost reports for the hospitals participating in that year. Update factors representing nationwide trends in cost and volume increases were incorporated into these estimates, as specified in the methodology described in the final rule for each fiscal year. Second, as finalized cost reports became available, we determined the amount by which the actual costs of the demonstration for an earlier, given year differed from the estimated costs for the demonstration set forth in the final IPPS rule for the corresponding fiscal year, and we incorporated that amount into the budget neutrality offset amount for the upcoming fiscal year. If the actual costs for the demonstration for the earlier fiscal year exceeded the estimated costs of the demonstration identified in the final rule for that year, this difference was added to the estimated costs of the demonstration for the upcoming fiscal year when determining the budget neutrality adjustment for the upcoming fiscal year. Conversely, if the estimated costs of the demonstration set forth in the final rule for a prior fiscal year exceeded the actual costs of the demonstration for that year, this difference was subtracted from the estimated cost of the demonstration for the upcoming fiscal year when determining the budget neutrality adjustment for the upcoming fiscal year. (We note that we have calculated this difference for FYs 2005 through 2010 between the actual costs of the demonstration as determined from finalized cost reports once available, and estimated costs of the demonstration as identified in the applicable IPPS final rules for these years.)

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57036), we finalized a different methodology as compared to previous years for analyzing the costs attributable to the demonstration for FY 2017. We noted in the FY 2017 IPPS/LTCH PPS final rule that, in accordance with the extension mandated by the Affordable Care Act, the demonstration would have substantially phased out by the beginning of FY 2017. In addition to the 7 originally participating hospitals (Cohort 1 hospitals) having ended their scheduled period of performance in the 5-year extension period prior to the start of FY 2016, we noted that the participation periods for the 14 hospitals that entered the demonstration following the extension mandated by the Affordable Care Act (Cohort 2 hospitals) that were still participating were to end on a rolling basis according to the end dates of the hospitals’ cost report periods, respectively, from April 30, 2016 through December 31, 2016. (As noted earlier, 1 hospital among the Cohort 2 hospitals closed in October 2015.) Of these 14 hospitals, 10 ended participation on or before September 30, 2016, leaving 4 hospitals participating for the last 3 months of FY 2016 (that is, the first 3 months of FY 2017). We stated that, given the small number of participating hospitals and the limited time of participation for such hospitals during FY 2017, a revised methodology was appropriate for determining the costs of the demonstration during this period. We noted that, for the 4 hospitals that would end their participation in the demonstration effective December 31, 2016, the financial experience of the last 3 months of the calendar year (that is, the first 3 months of FY 2017) would be included in the finalized cost reports for FY 2016. We stated that examining the finalized cost reports for FY 2016 for these hospitals would lead to a more accurate and administratively feasible calculation of budget neutrality for the demonstration in FY 2017 than conducting an estimate of the costs of the demonstration for this 3-month period based on “as submitted cost reports” (as would occur according to the budget neutrality methodology used prior to the FY 2017 IPPS/LTCH PPS final rule).

Thus, in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57037), we finalized the proposal to forego the process of estimating the costs attributable to the demonstration for FY 2017, and to instead calculate the costs of the demonstration and the resulting budget neutrality adjustment factor for the demonstration for FY 2017 once the finalized cost reports for cost reporting periods beginning in FY 2016 become available.


For the implementation approach that we are proposing in section V.L.3.b. of the preamble of this proposed rule, we are proposing that a budget neutrality offset methodology similar to previous years (prior to FY 2017) would be applied to the periods of performance under the second 5 years of the 10-year
extension period authorized by section 15003 of Public Law 114–255. With the potential exception of the demonstration costs for FY 2018 as discussed below, for the periods of performance under the second 5 years of the 10-year extension period, an estimate of the costs of the demonstration, generally determined from historical, “as submitted” cost reports for the participating hospitals and the appropriate update factors, would be incorporated into a budget neutrality offset amount to be applied to the national IPPS rates for the upcoming fiscal year. We would implement this adjustment through the corresponding proposed and final IPPS rules. In addition, we would include as a second component to the budget neutrality offset amount, the amount by which the actual costs of the demonstration for an earlier, given year (as determined from finalized cost reports when available) differed from the estimated costs for the demonstration set forth in the final IPPS rule for the corresponding fiscal year. Regarding demonstration costs specifically for FY 2018, as described earlier, we are proposing that if the selection of additional hospitals pursuant to section 410A(g)(6) of Public Law 108–173 (as added by section 15003 of Pub. L. 114–255) is announced by June 2017, we would include in the FY 2018 IPPS/LTCH PPS final rule an estimate of the costs of the demonstration for FY 2018 and the resulting estimated budget neutrality offset amount for the newly selected hospitals (Cohorts 1 and 2 hospitals) and for the previously participating hospitals (Cohorts 1 and 2 hospitals). As discussed earlier, if the final selection of the additional hospitals does not occur by June 2017, we would not be able to include an estimate of the costs of the demonstration for any participating hospitals or an estimated budget neutrality adjustment for FY 2018 in the FY 2018 IPPS/LTCH PPS final rule. In that situation, we are proposing to include the estimated costs of the demonstration for FY 2018 for all participating hospitals (Cohorts 1, 2 and 3 hospitals) in the budget neutrality offset adjustment in the FY 2019 IPPS/LTCH PPS proposed and final rules. The budget neutrality offset adjustment for the FY 2019 IPPS/LTCH PPS proposed and final rules would also include the estimated costs of the demonstration for FY 2019 for all participating hospitals based on historical, “as submitted” cost reports and the appropriate update factors.

As described earlier, if the selection of the newly participating hospitals authorized by section 410A(g)(6) of Public Law 108–173 (as added by section 15003 of Pub. L. 114–255) is announced by June 2017, we are proposing that the periods of performance under the second 5 years of the 10-year extension period for each of the participating hospitals (Cohorts 1, 2, and 3) would start with the hospital’s first cost report year on or after October 1, 2017. Thus, the start dates for the periods of performance for the entire set of participating hospitals would occur during FY 2018. If the selection of the new hospitals is announced by June 2017, under our proposed implementation approach as described in section V.L.3.b. of the preamble of this proposed rule, we would continue to use the general methodology finalized in previous final rules (prior to FY 2017) to calculate the estimated budget neutrality adjustment factor to be applied to the FY 2018 national IPPS rates. (We note that the same general methodology would be used if the announcement of the selection of additional hospitals does not occur by June 2017, and thus the budget neutrality offset amount reflecting the costs of the demonstration for hospitals participating in FY 2018 would be applied to the national IPPS rates for FY 2019.)

Consistent with the approach adopted in the FY 2016 IPPS/LTCH PPS final rule, we are proposing a specific calculation to account for the fact that the periods of performance for the participating hospitals would start at different points of time during FY 2018. That is, we are proposing to prorate estimated reasonable cost amounts and amounts that would be paid without the demonstration for FY 2018 according to the fraction of the number of months that the hospital would be participating out of the 12 months within FY 2018. For example, if a hospital would be starting this second 5-year period of the 10-year extension period on January 1, 2018, we would multiply the estimated cost and payment amounts, derived as described below, by a factor of 0.75. (In this discussion, we are discussing how the overall calculations are conducted, this factor is referred to as “the hospital-specific prorating factor”.) Our proposed methodology for calculating the budget neutrality offset amount proceeds in several steps, as set forth below:

**Step 1:** For each of the participating hospitals, we are proposing to identify the reasonable cost amount calculated under the reasonable cost methodology for covered inpatient hospital services, as indicated on the “as submitted” cost report for the most recent cost reporting period available. (We expect that for most of the hospitals these “as submitted” cost reports will be those with cost report period end dates in CY 2015. In the solicitation for additional participants, we will be requesting applicants to submit cost report information from the most recent year available. For the selected additional hospitals (that is, Cohort 3), we would be using the submitted information for the calculation of the budget neutrality offset amount for FY 2018.) We believe the most recent available cost reports to be an accurate predictor of the costs of the demonstration in FY 2018 because they would give us a recent picture of the participating hospitals’ costs.

Because section 410A of Public Law 108–173 stipulates swing-bed services are to be included among the covered inpatient hospital services for which the demonstration payment methodology applies, we are proposing to include the cost of these services, as reported on the cost reports for the hospitals that provide swing-bed services, in estimating the total reasonable cost amount for covered inpatient hospital services under the demonstration.

Similar to what is stated above, we are proposing to use the most recently available “as submitted” cost reports for this calculation.

For each hospital, we are proposing to sum the two above-referenced amounts, and then multiply this sum by the hospital-specific prorating factor (described above), to obtain an unadjusted hospital-specific amount, calculated for each hospital prior to applying adjustments for increases in cost or volume, as described below. (In the discussion below, we refer to this amount as the “unadjusted hospital-specific amount”.) We are proposing to sum these unadjusted hospital-specific amounts for all participating hospitals to obtain an unadjusted total estimated reasonable cost amount for covered inpatient hospital services (for all participating hospitals) to which update factors representing increases in costs and volume would be applied.

Accordingly, we are proposing to multiply this sum (that is, the unadjusted total estimated reasonable cost amount for covered inpatient hospital services for all participating hospitals) by the FY 2016, FY 2017, and final FY 2018 IPPS market basket percentage increases, which are formulated by the CMS Office of the Actuary. We are proposing to use the market basket percentage increases for these particular years because we expect that most of the “as submitted” cost reports that would be used in determining the unadjusted hospital-specific amounts will end in FY 2015. If a majority of these “as submitted”
cost reports end in FY 2016, we would apply only the FY 2017 and final FY 2018 market basket percentage increases. We recognize that applying the set of FY 2016, FY 2017, and FY 2018 market basket percentage increases to a sum that may include information from “as submitted” cost reports ending in FY 2016 (or, conversely, applying these update factors for FY 2017 and FY 2018 to a sum that may include information from “as submitted” cost reports ending in FY 2015) might appear to lessen the precision of the estimate. However, we believe that the potential margin of error in estimating the total costs for the demonstration hospitals inherent in using a uniform set of update factors is justifiable for purposes of streamlining and applying a consistent calculation method for all participating hospitals. In addition, we note that, as in previous years, we are proposing to reconcile the actual costs of the demonstration as determined from finalized cost reports when available with the estimate of the costs of the demonstration in FY 2018 as included in the budget neutrality offset amount, which would ultimately address any potential error in estimating the costs of the demonstration for FY 2018, thereby enhancing the accuracy of the calculation.

In this proposed rule, the current estimate of the FY 2018 IPPS market basket percentage increase provided by the CMS Office of the Actuary is specified in section V.B.1. of the preamble of this proposed rule. We also are proposing to then multiply the product of the unadjusted general total estimated reasonable cost amount for all participating hospitals and the market basket percentage increases applicable to the years involved by a 3-percent annual volume adjustment for each of FYs 2016 through 2018 (or only FYs 2017 and 2018, in accordance with the discussion above). The result would be the general total estimated FY 2018 reasonable cost amount for covered inpatient hospital services for all participating hospitals.

We are proposing to apply the IPPS market basket percentage increases applicable for FYs 2016 through 2018 (or FYs 2017 and 2018, in accordance with the discussion above) to the applicable general total reasonable cost amount described above to model the estimated FY 2018 reasonable cost amount under the demonstration. We are proposing to use the IPPS market basket percentage increases because we believe that these update factors appropriately indicate the trend of increase in inpatient hospital operating costs under the reasonable cost methodology for the years involved. The 3-percent annual volume adjustment was stipulated by the CMS Office of the Actuary and is being proposed because it is intended to accurately reflect the tendency of hospitals’ inpatient caseloads to increase. We acknowledge the possibility that inpatient caseloads for small hospitals may fluctuate, and thus are proposing to incorporate into the estimate of demonstration costs a factor to allow for a potential increase in inpatient hospital services.

Step 2: For each of the participating hospitals, we are proposing to identify the general estimated amount that would otherwise be paid in FY 2018 under applicable Medicare payment methodologies for covered inpatient hospital services (as indicated on the same set of “as submitted” cost reports as in Step 1) if the demonstration was not implemented. Similarly, as in Step 1, for the hospitals that provide swing-bed services, we are proposing to identify the estimated amount that generally would otherwise be paid for these services using the same “as submitted” cost reports as in Step 1 and include it in estimating the total FY 2018 general amount that would otherwise be paid for covered inpatient hospital services without the demonstration. Similar to Step 1, we are proposing to multiply this sum for each participating hospital by the hospital-specific prorating factor. We are then proposing to add together the resulting amounts for all participating hospitals to obtain an estimate of the amount that would otherwise be paid for covered inpatient hospital services for all participating hospitals without the demonstration, to which update factors representing increases in costs and volume would be applied.

Accordingly, we are proposing to then multiply this amount by the FYs 2016 through 2018 (or only FYs 2017 and 2018, in accordance with the discussion above) IPPS applicable percentage increases, depending on whether the majority of the “as submitted” cost reports end in FY 2015 or FY 2016, as discussed in Step 1. This methodology differs from Step 1, in which we are proposing to apply the market basket percentage increases to the sum of the hospitals’ applicable general total estimated reasonable cost amount for covered inpatient hospital services. We believe that the IPPS applicable percentage increases are appropriate factors to update the estimated amounts that generally would otherwise be paid without the demonstration. This is because any payments would constitute the majority of payments that would otherwise be made without the demonstration and the applicable percentage increase is the factor used under the IPPS to update the inpatient hospital payment rates. Most of the hospitals participating in the demonstration would be paid under the IPPS payment methodology if they were not in the demonstration. Then, for the same reasons discussed in Step 1, we are proposing to multiply the product of the applicable estimated total payments that generally would otherwise be made without the demonstration and the IPPS applicable percentage increases applicable to the years involved by the 3-percent annual volume adjustment for each of FYs 2016 through 2018 (or FYs 2017 and 2018, in accordance with the discussion above). The result would be the general total estimated payment amount that would otherwise be paid without the demonstration for FY 2018 to participating hospitals for covered inpatient hospital services.

Step 3: We are proposing to subtract the amount derived in Step 2 (representing the sum of estimated amounts that generally would otherwise be paid to the participating hospitals for covered inpatient hospital services for FY 2018 if the demonstration were not implemented) from the amount derived in Step 1 (representing the sum of the estimated reasonable cost amounts that generally would be paid under the demonstration to all participating hospitals for covered inpatient hospital services for FY 2018). We are proposing that the resulting difference would be the estimated amount of the costs of the demonstration for FY 2018, which would be incorporated into an adjustment to the national IPPS rates.

Similar to previous years, in order to meet the budget neutrality requirement in section 410A(c)(2) of Public Law 108–173, we are proposing that when finalized cost reports for each of the second 5 years of the 10-year extension period become available, we would determine the difference between the actual costs of the demonstration as determined from these finalized cost reports and the estimated cost indicated in the corresponding fiscal year IPPS final rule, and include that difference either as a positive or negative adjustment in the upcoming year’s final rule.

Specifically for FY 2018, when the finalized cost reports beginning in FY 2018 are available, we would determine the difference between the actual costs of the demonstration as determined from these finalized cost reports and the estimated cost indicated in the FY 2018 LTCH PPS final rule, and include that difference either as a positive or
negative adjustment in the applicable year’s final rule.

Thus, in keeping with the methodologies used in previous final rules, we would continue to use a methodology for calculating the budget neutrality offset amount for the second 5-years of the 10-year extension period consisting of two components: (1) The estimated demonstration costs in the upcoming fiscal year (as described above); and (2) the amount by which the actual demonstration costs corresponding to an earlier, given year (which would be known once finalized cost reports became available for that year) differed from the budget neutrality offset amount finalized in the corresponding year’s IPPS final rule.

We are inviting public comments on the budget neutrality calculation methodology proposed above. In addition, we are inviting comments on other approaches that would be consistent with section 410A(c)(2) of Public Law 108–173, and that would provide a reasonable determination of budget neutrality for the demonstration.

d. Alternative Budget Neutrality Approach Considered

In section V.L.3.b. of the preamble of this proposed rule, we described an alternative approach that we considered for implementing the extension of the demonstration pursuant to section 15003 of Public Law 114–255, and we invited public comments on this alternative approach. Under this alternative approach, for each previously participating hospital that decides to participate in the second 5 years of the 10-year extension period, the cost-based payment methodology under section 410A of Public Law 108–173 (as amended by section 15003 of Pub. L. 114–255) would begin on the date immediately following the end date of its period of performance for the first 5-year extension period.

Under this alternative approach that we considered, depending on which among the Cohorts 1 and 2 hospitals choose to participate in this second 5-year extension period, the demonstration’s cost-based payment methodology would be applied to dates as far back as January 1, 2015 and as late as January 1, 2017. This would require reconciling the reasonable costs associated with furnishing Medicare covered inpatient hospital services as reported on cost reports with the amounts already paid under the other Medicare payment methodologies applied since the end of their periods of performance for the first 5-year extension. Under this alternative approach, any additional amounts associated with the cost-based payment methodology for this period would need to be paid to the hospitals.

Although we considered this alternative implementation approach and budget neutrality methodology, for the reasons discussed in section V.L.3.b. of the preamble to this proposed rule, we are instead proposing the implementation approach (according to which the periods of performance for previously participating hospitals for the second 5-year extension period would begin with the hospital’s first cost reporting period on or after October 1, 2017, following the announcement of the selection of additional hospitals) and budget neutrality methodology described in sections V.L.3.b. and V.L.4.c. of the preamble of this proposed rule. Nevertheless, we are inviting public comments on the budget neutrality methodology that we describe below for the alternative approach.

In general, the methodology that we considered for calculating a budget neutrality offset under this alternative approach would involve the following steps:

• To reflect the costs of the demonstration for fiscal years before FY 2018, for the previously participating hospitals (Cohorts 1 and 2) that decide to participate in the 5-year extension period authorized by section 15003 of Public Law 114–255, when final cost reports become available, we would determine the actual costs of the demonstration for cost report periods beginning on the day after the last day of the hospitals’ periods of performance in the first 5-year extension period and extending through the last day of the cost report periods ending in FY 2018 (or FY 2017 for hospitals with an October 1 cost report start date, as explained below), and incorporate these amounts into the calculation of the budget neutrality offset amount to be included in a future IPPS final rule. Thus, we would determine the actual costs for the previously participating hospitals for the period prior to the start of FY 2018. Similar to our proposed approach for implementation and budget neutrality, as described in sections V.L.3.b. and V.L.4.c. of the preamble of this proposed rule, under the alternative methodology we considered, we would seek to begin our estimation of the costs of the demonstration for all hospitals in the same fiscal year (that is, in FY 2018, with each hospital’s first cost reporting period beginning on or after October 1, 2017). (The methodology for estimating the costs for this alternative implementation approach that we considered is described below.)

Thus, under the alternative approach we considered, for a Cohort 1 hospital whose period of performance in the first extension period ended June 30, 2015, we would determine the actual costs of the demonstration for the cost reporting periods from July 1, 2015 through June 30, 2016, from July 1, 2016 through June 30, 2017, and from July 1, 2017 through June 30, 2018. For a Cohort 2 hospital whose period of performance in the first extension period ended June 30, 2016, under this alternative approach that we considered, we would determine the actual costs of the demonstration for the cost reporting periods from July 1, 2016 through June 30, 2017, and from July 1, 2017 through June 30, 2018. We note that for both of these Cohorts 1 and 2 hospitals, this last cost report period would encompass services occurring since the enactment of Public Law 114–255, which authorizes the second extension period. However, we believe that applying a uniform method for determining costs across a cost report year would be more reasonable from the standpoint of operational feasibility and consistent application of cost determination principles. (We note that, for hospitals (either Cohort 1 or 2) with an October 1 start date, the estimation of costs for FY 2018 would apply for the period starting October 1, 2017, that is, the first day of FY 2018. Therefore, for these hospitals, we would determine actual costs from finalized cost reports when available for the period starting from the day after the last day of the period of performance under the first 5-year extension period and concluding with the last day of FY 2017.) For all hospitals, under this alternative approach that we considered, we would incorporate these amounts into a single amount to be included in the calculation of the budget neutrality offset amount to the national IPPS rates in a future final rule after such finalized cost reports become available.

• To reflect the costs of the demonstration for the upcoming fiscal year (that is, FY 2018) for Cohorts 1 and 2 hospitals that have decided to participate in the second 5-years of the 10-year extension period, we would estimate the costs of the demonstration for FY 2018, based on historical “as submitted” cost reports, applying prorating factors and updates as appropriate, as described below. Similar to the proposed methodology described in section V.L.4.c. of the preamble of this proposed rule for estimating the costs of the demonstration for FY 2018, the alternative methodology we considered for estimating the costs of
the demonstration for FY 2018 would follow 3 steps: 

Step 1: We would determine the total estimated reasonable cost amount for covered inpatient hospital services (as indicated on the “as submitted” cost reports for the most recent cost reporting period available) for all participating hospitals for FY 2018 calculated under the demonstration’s reasonable cost-based payment methodology. These calculations would be identical to those described for our proposed methodology in section V.L.4.c. of the preamble of this proposed rule, with the exception that the formulation of the “hospital-specific prorating factor,” to be applied to each participating hospital’s reasonable cost amounts as derived from its most recently available “as submitted” cost report, would be different. Under this alternative methodology that we considered, for hospitals with a cost report start date other than October 1, the hospital-specific prorating factor would be the ratio of the number of months between the end of the cost reporting period ending in FY 2018, on the basis of which actual costs are determined (as described above), and the end of the fiscal year, out of the total number of months in the fiscal year. Therefore, for a hospital (either Cohort 1 or 2) for which the end of the period on which we would determine actual costs (that is, the end date of the hospital’s cost report year) would be June 30, 2018, there would be 3 months remaining in FY 2018, and the hospital-specific prorating factor would be .25. (Hospitals with an October 1 cost report start date would participate in the demonstration for the full 12 months of FY 2018 and thus would have a hospital-specific prorating factor of 1.0.) We would then follow the same calculations as in our proposed budget neutrality calculation described in section V.L.4.c. of the preamble of this proposed rule, including application of the same update factors to reflect increases in cost and volume. 

Step 2: We would estimate the amount that would otherwise be paid for Medicare covered inpatient hospital services to all participating hospitals in FY 2018 without the demonstration. These calculations would be identical to those described for our proposed methodology in section V.L.4.c. of the preamble of this proposed rule, except for the difference that the hospital-specific prorating factor, to be applied to the estimated amount that the hospital would be paid without the demonstration, as derived from its most recently available “as submitted” cost report, would be formulated in the same manner as described in Step 1 above under the alternative methodology. 

Step 3: We would then subtract the amount derived in Step 2 (representing the estimated amount that would otherwise be paid to the participating hospitals for covered inpatient hospital services for FY 2018 if the demonstration were not implemented) from the amount derived in Step 1 (representing the estimated reasonable cost amounts that generally would be paid under the demonstration to all participating hospitals for covered inpatient hospital services for FY 2018). The resulting difference would be the estimated amount of the costs of the demonstration for FY 2018, which would be incorporated into an adjustment to the national IPPS rates. 

- For the Cohort 3 hospitals, we would follow the identical methodology for estimating the costs of the demonstration for FY 2018 as described for the proposed budget neutrality methodology under the proposed implementation approach. Similar to the description above for the proposed approach, if the selection of additional participants under the solicitation authorized by Public Law 114–255 is announced by June 2017, we would be able to incorporate the estimates of the costs of the demonstration for the Cohort 3 hospitals for FY 2018 within a budget neutrality offset adjustment to be included in the FY 2018 IPPS/LTCH PPS final rule. However, we note that if this selection is not announced by that time, we would not be able to include the estimated costs of the demonstration for FY 2018 for the Cohort 3 hospitals in the budget neutrality offset adjustment for FY 2018, and similar to our proposed methodology in that situation, we would incorporate this estimate in the budget neutrality offset adjustment in the FY 2019 IPPS/LTCH PPS final rule. The budget neutrality offset adjustment for the FY 2019 IPPS proposed and final rules would also include the estimated costs of the demonstration for FY 2019 for these Cohort 3 hospitals based on historical, “as submitted” cost reports and the appropriate update factors. 

- Consistent with our approach in previous final rules, when the finalized cost reports for cost reporting periods beginning in FY 2018 are available, we would determine the difference between the actual costs of the demonstration as determined from these finalized cost reports and the estimated cost indicated in the FY 2018 IPPS/LTCH PPS final rule (as explained above), and include that difference either as a positive or negative adjustment in the upcoming year’s final rule. 

- For future years, we would continue to incorporate the estimated costs of the demonstration for all participating hospitals for the upcoming fiscal year in the budget neutrality offset adjustment in the IPPS final rule of the corresponding fiscal year. For these hospitals, we also would determine the actual costs of the demonstration when finalized cost reports become available, and include the difference between the estimated and actual costs of the demonstration in the calculation of the budget neutrality offset amount to the national IPPS rates in the final rule for a future year. 

We note that, under the alternative approach we considered, although we would not be able to include an estimate of the costs of the demonstration for FY 2018 Cohort 3 hospitals in the budget neutrality offset adjustment in the FY 2018 final rule if we were not able to announce the selection of additional hospitals by June 2017, we could do so for the Cohorts 1 and 2 hospitals. However, we note the overall complexity of the methodology for budget neutrality under this alternative implementation methodology, involving various differing methods for either determining or estimating the costs of the demonstration over several different fiscal years, potentially to be applied to budget neutrality offset adjustment amounts for IPPS final rules for different fiscal years. We believe our proposed implementation approach and budget neutrality calculation (described in sections V.L.3.b. and V.L.4.c. of the preamble of this proposed rule) are more reasonable and appropriate for the reasons discussed previously, and because of the complexity inherent in meeting the budget neutrality requirement, and the administrative burden involved in tracking payments and associated calculations over multiple years under the alternative methodology. 

Nevertheless, we are inviting public comments on the alternative budget neutrality calculation methodology we considered, as discussed above. 

e. Reconciling Actual and Estimated Costs of the Demonstration for Previous Years (2011, 2012, and 2013) 

As described earlier, we have calculated the difference for FY’s 2005 through 2010 between the actual costs of the demonstration, as determined from finalized cost reports once available, and estimated costs of the demonstration as identified in the applicable IPPS final rules for these years. In the FY 2017 IPPS/LTCH PPS
final rule (81 FR 57037), we finalized a proposal to reconcile the budget neutrality offset amounts identified in the IPPS final rules for FYs 2011 through 2016 with the actual costs of the demonstration for those years, considering the fact that the demonstration was scheduled to end December 31, 2016. In that final rule, we stated that we believed it would be appropriate to conduct this analysis for FYs 2011 through 2016 at one time, when all of the finalized cost reports for cost reporting periods beginning in FYs 2011 through 2016 are available. We stated that such an aggregate analysis encompassing the cost experience through the end of the period of performance of the demonstration would represent an administratively streamlined method, allowing for the determination of any appropriate adjustment to the IPPS rates and obviating the need for multiple, fiscal year-specific calculations and regulatory actions. Given the general lag of 3 years in finalizing cost reports, we stated that we expected any such analysis would be conducted in FY 2020.

With the extension of the demonstration for another 5-year period, as authorized by section 15003 of Public Law 114-255, we are proposing to modify the plan outlined in the FY 2017 IPPS/LTCH PPS final rule, and instead return to the general procedure in previous final rules; that is, as finalized cost reports become available, to determine the amount by which the actual costs of the demonstration for an earlier, given year differ from the estimated costs for the demonstration set forth in the IPPS final rule for the corresponding fiscal year, and then incorporate that amount into the budget neutrality offset amount for an upcoming fiscal year. If the actual costs of the demonstration for the earlier fiscal year exceed the estimated costs of the demonstration identified in the final rule for that year, this difference would be added to the estimated costs of the demonstration for the upcoming fiscal year when determining the budget neutrality adjustment for the final rule. Conversely, if the estimated costs of the demonstration set forth in the final rule for a prior fiscal year exceed the actual costs of the demonstration for that year, this difference would be subtracted from the estimated cost of the demonstration for the upcoming fiscal year when determining the budget neutrality adjustment for an upcoming fiscal year. However, given that this adjustment for specific years could be positive or negative, we are proposing to combine this reconciliation for multiple prior years into one adjustment to be applied to the budget neutrality offset amount for a single fiscal year, thus lessening the possibility of both positive and negative adjustments to be applied in consecutive years, and enhancing administrative feasibility. Specifically, we are proposing that when finalized cost reports for FYs 2011, 2012, and 2013 are available, we would include this difference for these years in the budget neutrality offset adjustment to be applied to the national IPPS rates in a future final rule. We expect that this would occur in FY 2019. We also are proposing that when finalized cost reports for FYs 2014 through 2016 are available, we would include the difference between the actual costs as reflected on these cost reports and the amounts included in the budget neutrality offset amounts for these fiscal years in a future final rule. We plan to provide an update in a future final rule regarding the year that we would expect that this analysis would occur.

We are inviting public comments on this proposal.

M. Payment for Services in Inpatient and Outpatient Hospital Settings

1. Adjustment to IPPS Rates Resulting From 2-Midnight Policy for FY 2018

In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50906 through 50954), we adopted the 2-midnight policy, effective for dates of admission on or after October 1, 2013. As discussed in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57058 through 57060), under the 2-midnight policy, an inpatient admission is generally appropriate for Medicare Part A payment if the physician (or other qualified practitioner) admits the patient as an inpatient based upon the reasonable expectation that the patient will need hospital care that crosses at least 2 midnights. In assessing the expected duration of necessary care, the physician (or other qualified practitioner) must take into account outpatient hospital care received prior to inpatient admission. If the patient is expected to need less than 2 midnights of care in the hospital, the services furnished should generally be billed as outpatient services. We note that revisions were made to this policy in the CY 2016 OPPS/ASC final rule with comment period (80 FR 70545). Our actuaries estimated that the 2-midnight policy would increase expenditures by approximately $220 million in FY 2014 due to an expected net increase in inpatient encounters. We used our authority under section 1886(d)(5)(I)(i) of the Act to make a reduction of 0.2 percent to the standardized amount, the Puerto Rico standardized amount, and the hospital-specific payment rates, and we used our authority under section 1886(g) of the Act to make a reduction of 0.2 percent to the national capital Federal rate and the Puerto Rico-specific capital rate, in order to offset this estimated $220 million in additional IPPS expenditures in FY 2014.

For the reasons outlined in the FY 2017 IPPS/LTCH PPS proposed and final rules (81 FR 25136 through 25138 and 57058 through 57060), we used our authority under section 1886(d)(5)(I)(i) and 1886(g) of the Act to prospectively remove, beginning in FY 2017, the 0.2 percent reduction to the rates put in place beginning in FY 2014. The 0.2 percent reduction was implemented by including a factor of 0.998 in the calculation of the FY 2014 standardized amount, hospital-specific payment rates, and the national capital Federal rate, permanently reducing the rates for FY 2014 and future years until the 0.998 is removed. In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57281 and 57294), we permanently removed the 0.998 reduction beginning in FY 2017 by including a factor of (1/0.998) in the calculation of the FY 2017 standardized amount, the hospital-specific payment rates, and the national capital Federal rate.

We also stated in the FY 2017 IPPS/LTCH PPS proposed and final rules that, for the reasons outlined in those rules, we believe it would be appropriate to use our authority under sections 1886(d)(5)(I)(i) and 1886(g) of the Act to temporarily increase the rates, only for FY 2017, to address the effect of the 0.2 percent reduction to the rates in effect for FY 2014, the 0.2 percent reduction to the rates in effect for FY 2015 (recall the 0.998 factor included in the calculation of the FY 2014 rates permanently reduced the rates for FY 2014 and future years until it is removed), and the 0.2 percent reduction to the rates in effect for FY 2016. We stated that we believe the most transparent, expedient, and administratively feasible method to accomplish this was a temporary one-time prospective increase to the FY 2017 rates of 0.6 percent (= 0.2 percent + 0.2 percent + 0.2 percent). Specifically, we finalized our proposal to include a factor of 1.006 in the calculation of the standardized amount, the hospital-specific payment rates, and the national capital Federal rate in FY 2017 and then to remove this temporary one-time prospective increase by including a factor of (1/1.006) in the calculation of the rates for FY 2018. We stated that while we generally did not believe it is appropriate in a prospective
In summary, for the reasons described in the FY 2017 IPPS/LTCH PPS proposed and final rules, we finalized our proposal to include a permanent factor of (1/0.998) and a temporary one-time factor of (1.006) in the calculation of the FY 2017 standardized amount, hospital-specific payment rates, and national capital Federal rate and to include a factor of (1/1.006) in the calculation of the FY 2018 standardized amount, hospital-specific payment rates, and national capital Federal rate to remove the temporary one-time factor of 1.006.

In this FY 2018 IPPS/LTCH PPS proposed rule, we are including a factor of (1/1.006) in the calculation of the FY 2018 standardized amount, hospital-specific payment rates, and national capital Federal rate to remove the temporary one-time factor of 1.006, as explained in detail in section II. of the Addendum to this proposed rule.

We note that, in the FY 2017 IPPS/LTCH PPS final rule, in our response to public comments, we recognized that for closed, converted, or new hospitals, our prospective method generally may have had a differential positive or negative impact compared to hospitals that were IPPS hospitals for all of the FY 2014 through FY 2017 time period. We stated that we generally believe that, given the prospective nature of our method and our goal to adopt a transparent, expedient, and administratively feasible approach, these differential impacts would be an appropriate consequence. However, after consideration of the public comments received, we agreed that we should provide a process to address the situation of closed or converted hospitals. Due to the small number of hospitals impacted, we stated that we will address closed and converted hospitals as part of the cost report settlement process. We stated that these hospitals should identify themselves to their MACs so that the appropriate cost report adjustment can be applied.

2. Eliminating Inappropriate Medicare Payment Differentials for Similar Services in the Inpatient and Outpatient Settings

CMS is committed to eliminating inappropriate Medicare payment differentials for similar services in the inpatient and outpatient settings in order to execute our responsibility to taxpayers to prudently pay for high quality care. MedPAC has previously noted, “The high profitability of one-day stays under the inpatient prospective payment system (IPPS) and the generally lower payment rates for similar care under the outpatient prospective payment system (OPPS) have heightened concern about the appropriateness of inpatient one-day stays” (Medicare and the Health Care Delivery System Report to Congress, June 2015).

In the past, CMS has requested public comment on potential payment policy options to address the issue of payment differentials between services provided in the inpatient and outpatient settings. However, our most recent solicitation occurred in the CY 2016 OPPS/ASC final rulemaking (80 FR 70549). Since that time, both hospitals and CMS have had the opportunity to gain experience under the various policy changes that have occurred with respect to short inpatient hospital stays. In this context, we believe it is an appropriate time to seek public comment on transparent ways to identify and eliminate inappropriate payment differentials for similar services provided in the inpatient and outpatient settings.

N. Provider-Based Status of Indian Health Service and Tribal Facilities and Organizations

Since the beginning of the Medicare program, some providers, which we refer to as “main providers,” have functioned as a single entity while owning and operating multiple departments, locations, and facilities. We have maintained that having clear criteria for provider-based status is important because a provider-based status designation can result in additional Medicare payments under the OPPS for services provided at the provider-based facility, as well as increased beneficiary coinsurance liability for Medicare beneficiaries. The Medicare criteria for provider-based status are set forth in our regulations at 42 CFR 413.65. In the April 7, 2000 OPPS final rule (65 FR 18507), CMS (then HCFA), responded to several commenters who were concerned that the implementation of the proposed provider-based regulations would have the effect of denying Medicare participation as provider-based entities to a number of Indian Health Service (IHS) facilities that were being operated by Indian Tribes under the auspices of the Indian Self-Determination and Education Assistance Act in accordance with applicable regulations and policies of the IHS in consultation with Tribes; or (3) owned by the IHS or Tribes and operated by the Tribe under the Indian Self-Determination and Education Assistance Act in accordance with applicable regulations and policies of the IHS in consultation with Tribes.

In order to qualify for grandfathering under §413.65(m), we required that the services be furnished by the facility or organization on or before April 7, 2000 because of our concern that, without such a date limitation, this provision would create an incentive for IHS or Tribal hospitals to establish new outpatient departments that were not sufficiently integrated with the main provider to support payment under the OPPS for the services that they furnished. Our intent was to implement a policy that both addressed a primary concern (that is, the rapid growth of off-campus provider-based clinics) that necessitated the provider-based regulations and recognized longstanding and complex IHS and Tribal
arrangements. Since we finalized the policy at § 413.65(m), we have issued guidance on circumstances that would and would not result in a facility or organization losing its grandfathered status. In particular, we recognized the special relationship between Tribes and the IHS under the Self-Determination and Education Assistance Act and stated that changes in the status of a hospital or a facility from IHS to Tribal operation, or vice versa, or the realignment of a facility from one IHS or Tribal hospital to another IHS or Tribal hospital, would not be a basis for losing such a grandfathered status, so long as the resulting configuration is one that would have qualified for grandfathering under § 413.65(m) had it been in effect on April 7, 2000.

In the years since we implemented § 413.65(m) and issued the guidance described earlier, we have considered whether it remains necessary to require that facilities and organizations be furnishing the services on or before April 7, 2000 in order to qualify for grandfathering. We have concluded that it does not because IHS policies and procedures (for example, as specified in the Indian Health Manual available on the IHS Web site at: https://ihs.gov/aboutihs/indianhealthmanual/) regarding the planning, operation, and funding of such facilities and organizations are resulting in appropriate Medicare payments to them. Therefore, after further consideration of the position CMS has set out in prior guidance, the special and legally recognized relationship between Tribal Tribes and the U.S. Government, as well as current IHS policies and procedures, we are proposing to remove the date limitation in § 413.65(m) that restricted the grandfathering provision to IHS or Tribal facilities and organizations furnishing services on or before April 7, 2000.

We also are proposing to make a technical change to the billing reference in § 413.65(m) by replacing “were billed” with “are billed using the CCN of the main provider and with the consent of the main provider.” We believe this proposed change will make the regulation text more consistent with our current rules that require these facilities to comply with all applicable Medicare conditions of participation that apply to the main provider. We are not proposing to otherwise change the requirement that the only services furnished at the facility or organization must be hospital outpatient services, or to change the other requirements for grandfathering in paragraphs (m)(1) through (3) of § 413.65. Therefore, under our proposal, a facility or organization operated by the IHS or a Tribe will be considered to be a department of a hospital operated by the IHS or a Tribe if it furnishes only hospital outpatient services that are billed using the CMS Certification Number (CCN) of the main provider with the consent of the main provider, and it also meets one of the conditions in § 413.65(m)(1) through (3).

We welcome public comments on our proposals.

O. Request for Information Regarding Physician-Owned Hospitals

We are seeking public comments on the appropriate role of physician-owned hospitals in the delivery system. We are also seeking public comments on how the current scope of and restrictions on physician-owned hospitals affects healthcare delivery. In particular, we are interested in comments on the impact on Medicare beneficiaries.

VI. Proposed Changes to the IPPS for Capital-Related Costs

A. Overview

Section 1886(g) of the Act requires the Secretary to pay for the capital-related costs of inpatient acute hospital services in accordance with a prospective payment system established by the Secretary. Under the statute, the Secretary has broad authority in establishing and implementing the IPPPS for acute care hospital inpatient capital-related costs. We initially implemented the IPPPS for capital-related costs in the FY 1992 IPPS final rule (56 FR 43358). In that final rule, we established a 10-year transition period to change the payment methodology for Medicare hospital inpatient capital-related costs from a reasonable cost-based payment methodology to a prospective payment methodology (based fully on the Federal rate).

FY 2001 was the last year of the 10-year transition period that was established to phase in the IPPPS for hospital inpatient capital-related costs. For cost reporting periods beginning in FY 2002, capital IPPPS payments are based solely on the Federal rate for almost all acute care hospitals (other than hospitals receiving certain exception payments and certain new hospitals). (We refer readers to the FY 2002 IPPPS final rule (66 FR 39910 through 39914) for additional information on the methodology used to determine capital IPPPS payments to hospitals both during and after the transition period.)

The basic methodology for determining capital prospective payments using the Federal rate is set forth in the regulations at 42 CFR 412.312. For the purpose of calculating capital payments for each discharge, the standard Federal rate is adjusted as follows:

(Standard Federal Rate) × (DRG Weight) × (Geographic Adjustment Factor (GAF)) × (COLA for hospitals located in Alaska and Hawaii) × (1 + Capital DSH Adjustment Factor + Capital IME Adjustment Factor, if applicable).

In addition, under § 412.312(c), hospitals also may receive outlier payments under the capital IPPPS for extraordinarily high-cost cases that qualify under the thresholds established for each fiscal year.

B. Additional Provisions

1. Exception Payments

The regulations at 42 CFR 412.348 provide for certain exception payments under the capital IPPPS. The regular exception payments provided under § 412.348(b) through (e) were available only during the 10-year transition period. For a certain period after the transition period, eligible hospitals may have received additional payments under the special exceptions provisions at § 412.348(g). However, FY 2012 was the final year hospitals could receive special exceptions payments. For additional details regarding these exceptions policies, we refer readers to the FY 2012 IPPPS/LTCH PPS final rule (76 FR 51725).

Under § 412.348(f), a hospital may request an additional payment if the hospital incurs unanticipated capital expenditures in excess of $5 million due to extraordinary circumstances beyond the hospital’s control. Additional information on the exception payment for extraordinary circumstances in § 412.348(f) can be found in the FY 2005 IPPPS final rule (69 FR 49185 and 49186).

2. New Hospitals

Under the capital IPPPS, the regulations at 42 CFR 412.300(b) define a new hospital as a hospital that has operated (under previous or current ownership) for less than 2 years and lists examples of hospitals that are not considered new hospitals. In accordance with § 412.304(c)(2), under the capital IPPPS, a new hospital is paid 85 percent of its allowable Medicare inpatient hospital capital-related costs through its first 2 years of operation, unless the new hospital elects to receive full prospective payment based on 100 percent of the Federal rate. We refer readers to the FY 2012 IPPPS/LTCH PPS final rule (76 FR 51725) for additional information on payments to new hospitals under the capital IPPPS.
3. Payments for Hospitals Located in Puerto Rico

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57061), we revised the regulations at 42 CFR 412.374 relating to the calculation of capital IPPS payments to hospitals located in Puerto Rico beginning in FY 2017 to parallel the change in the statutory calculation of operating IPPS payments to hospitals located in Puerto Rico, for discharges occurring on or after January 1, 2016, made by section 601 of the Consolidated Appropriations Act, 2016 (Pub. L. 114–113). Section 601 of Public Law 114–113 increased the applicable Federal percentage of the operating IPPS payment for hospitals located in Puerto Rico from 75 percent to 100 percent and decreased the applicable Puerto Rico percentage of the operating IPPS payments for hospitals located in Puerto Rico from 25 percent to zero percent, applicable to discharges occurring on or after January 1, 2016. As such, under revised §412.374, for discharges occurring on or after October 1, 2016, capital IPPS payments to hospitals located in Puerto Rico are based on 100 percent of the capital Federal rate.

C. Proposed Annual Update for FY 2018

The proposed annual update to the national capital Federal rate, as provided for at §412.306(c), for FY 2018 is discussed in section III. of the Addendum to this proposed rule.

In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50906 through 50954), we adopted the 2-midnight policy effective for dates of admission on or after October 1, 2013, under which an inpatient admission is generally appropriate for Medicare Part A payment if the physician (or other qualified practitioner) admits the patient as an inpatient based upon the reasonable expectation that the patient will need hospital care that crosses at least 2 midnights. At that time, our actuaries estimated that the 2-midnight policy would increase expenditures by approximately $220 million in FY 2014 due to an expected net increase in inpatient encounters. Using our authority under section 1886(g) of the Act, and consistent with the approach taken for the operating IPPS standardized amount, the Puerto Rico-specific standardized amount and the hospital-specific payment rates, we made a reduction of 0.2 percent (an adjustment factor of 0.998) to the national capital Federal rate and the Puerto Rico-specific capital rate to offset the estimated increase in capital IPPS expenditures associated with the projected increase in inpatient encounters that was expected to result from the new inpatient admission guidelines (78 FR 50746 through 50747). (As explained in section V.B.3. of the FY 2017 IPPS/LTCH PPS final rule, we discontinued use of the Puerto Rico capital rate in the calculation of capital IPPS payments to hospitals located in Puerto Rico beginning in FY 2017.)

For the reasons discussed in the FY 2017 IPPS/LTCH PPS proposed and final rules (81 FR 25136 through 25138 and 57058 through 57060) and consistent with our approach for the operating IPPS rates, we used our authority under section 1886(g) of the Act to permanently remove the 0.2 percent reduction to the national capital Federal rate beginning in FY 2017. Specifically, we made an adjustment of (1/0.998) to the national capital Federal rate to remove the 0.2 percent reduction, consistent with the adjustment to the operating IPPS standardized amount and the hospital-specific payment rates.

In addition, consistent with our approach for the operating IPPS standardized amount and hospital-specific payment rates, and for the reasons discussed in the FY 2017 IPPS/LTCH PPS proposed and final rules, we finalized our proposal to use our authority under section 1886(g) of the Act to adjust the FY 2017 national capital Federal rate to address the effects of the 0.2 percent reduction to the national capital Federal rates in effect for FY 2014, FY 2015, and FY 2016 by making a one-time prospective adjustment of 1.006 in FY 2017 to the national capital Federal rate and, for FY 2018, to remove the effects of this one-time prospective adjustment through an adjustment of (1/1.006) to the national capital Federal rate. Therefore, consistent with our finalized policy, for FY 2018, we are including a factor of (1/1.006) in the calculation of the FY 2018 operating IPPS standardized amount, the hospital-specific payment rates, and the national capital Federal rate to remove the temporary one-time factor of 1.006. (For additional details, we refer readers to section IV.P. of the preamble of the FY 2017 IPPS/LTCH PPS final rule (81 FR 57058 through 57060 and 57062 through 57063) and to section V.M. of the preamble of this proposed rule.)

In section II.D. of the preamble of this proposed rule, we present a discussion of the MS–DRG documentation and coding adjustment, including previously finalized policies and historical adjustments, as well as the adjustment to the standardized amount under section 1886(g) of the Act that we are proposing for FY 2018 in accordance with the amendments made to section 7(b)(1)(B) of Public Law 110–90 by section 414 of the MACRA and section 15005 of the 21st Century Cures Act. Because these provisions require us to make an adjustment only to the operating IPPS standardized amount, we are not proposing a similar adjustment to the national capital Federal rate (or to the hospital-specific rates).

VII. Proposed Changes for Hospitals Excluded From the IPPS

A. Proposed Rate-of-Increase in Payments To Excluded Hospitals for FY 2018

Certain hospitals excluded from a prospective payment system, including children’s hospitals, 11 cancer hospitals, and hospitals located outside the 50 States, the District of Columbia, and Puerto Rico (that is, hospitals located in the U.S. Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa) receive payment for inpatient hospital services they furnish on the basis of reasonable costs, subject to a rate-of-increase ceiling. A per discharge limit (the target amount as defined in §413.40(a) of the regulations) is set for each hospital based on the hospital’s own cost experience in its base year, and updated annually by a rate-of-increase percentage. For each cost reporting period, the updated target amount is multiplied by total Medicare discharges during that period and applies as an aggregate upper limit (the ceiling as defined in §413.40(a)) of Medicare reimbursement for total inpatient operating costs for a hospital’s cost reporting period. In accordance with §403.752(a) of the regulations, religious nonmedical health care institutions (RNHCl)s also are subject to the rate-of-increase limits established under §413.40 of the regulations discussed previously.

As explained in the FY 2006 IPPS final rule (70 FR 47396 through 47398), beginning with FY 2006, we have used the percentage increase in the IPPS operating market basket to update the target amounts for children’s hospitals, cancer hospitals, and RNHCl.s. Consistent with §§412.23(g), 413.40(a)(2)(ii)(A), and 413.40(c)(3)(viii), we also have used the percentage increase in the IPPS operating market basket to update the target amounts for short-term acute care hospitals located in the U.S. Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa. In the FYs 2014 and 2015 IPPS/LTCH PPS final rules (76 FR 50747 through 50748 and 79 FR 50136 through 50157, respectively), we adopted a policy of using the percentage increase in the FY
2010-based IPPS operating market basket to update the target amounts for FY 2014 and subsequent fiscal years for children’s hospitals, cancer hospitals, RNCHCs, and short-term acute care hospitals located in the U.S. Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa. However, as discussed in section IV. of the preamble of this proposed rule, we are proposing to revise and rebase the IPPS operating market basket to a 2014 base year. Therefore, we are proposing to use the percentage increase in the 2014-based IPPS operating market basket to update the target amounts for children’s hospitals, the 11 cancer hospitals, RNCHCs, and short-term acute care hospitals located in the U.S. Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa for FY 2018 and subsequent fiscal years. Accordingly, for FY 2018, the rate-of-increase percentage to be applied to the target amount for these children’s hospitals, cancer hospitals, RNCHCs, and short-term acute care hospitals located in the U.S. Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa would be the FY 2018 percentage increase in the 2014-based IPPS operating market basket. Based on IHS Global Insight, Inc.’s 2016 fourth quarter forecast, for this proposed rule, we estimate that the 2014-based IPPS operating market basket update for FY 2018 would be 2.9 percent (that is, the estimate of the market basket rate-of-increase). We are proposing that if more recent data become available for the final rule, we would use them to calculate the IPPS operating market basket update for FY 2018.

In addition, as discussed in section VIII.J. of the preamble of this proposed rule, as originally enacted section 1886(d)(1)(B)(iv) of the Act established an IPPS-excluded category of hospitals that experience extended average inpatient length-of-stays, which are known as LTCHs under the Medicare program. Historically, section 1886(d)(1)(B)(iv) of the Act consisted of two subclauses (I) and (II) (that is, sections 1886(d)(1)(B)(iv)(I) and (d)(1)(B)(iv)(II) of the Act), and the two categories of hospitals were generally referred to as “subclause (I)” and “subclause (II)” LTCHs. Section 15008 of the 21st Century Cures Act (Pub. L. 114–255) amended section 1886(d)(1)(B) of the Act by redesignating the “subclause (II) LTCH” provision in section 1886(d)(1)(B)(iv)(II) of the Act to section 1886(d)(1)(B)(vi) of the Act. In addition, subsection (b) of section 15008 of Public Law 114–255 specifies that, for cost reporting periods beginning on or after January 1, 2015, hospitals classified under section 1886(d)(1)(B)(vi) of the Act are not subject to section 1886(m) of the Act, which sets forth the LTCH PPS. Section 15008(c) further specifies that, for cost reporting periods beginning on or after January 1, 2015, payment for inpatient operating costs for such hospitals is to be made as described in 42 CFR 412.526(c)(3), and payment for capital costs is to be made as described in 42 CFR 412.526(c)(4). In order to implement these requirements, we are proposing to amend § 412.23 to codify the redesignation of such hospitals from section 1886(d)(1)(B)(iv)(II) of the Act to section 1886(d)(1)(B)(vi) of the Act (which we are now referring to as “long-term care neoplastic disease hospitals”) and the statutory payment requirements for inpatient operating and capital costs. (For additional information on “subclause (II)” LTCHs, including the statutory criteria and the establishment of the payment adjustment under § 412.526, and our proposed changes to § 412.23 to implement the provisions of section 15008 of Public Law 114–255, we refer readers to section VIII.J. of the preamble of this proposed rule.)

Under the redesignation of subclause (II) LTCHs to long-term care neoplastic disease hospitals provided by section 15008 of Public Law 114–255 (described above), the statute specifies that payment for inpatient operating costs shall continue to be made on a reasonable cost basis in the manner provided in § 412.526(c)(3) of the regulations. § 412.526(c)(3) provides that the hospital’s Medicare allowable net inpatient operating costs for that period are paid on a reasonable cost basis, subject to that hospital’s ceiling, as determined under § 412.526(c)(1), for that period. Under section 15008(c)(3), for each cost reporting period, the ceiling was determined by multiplying the updated target amount, as defined in § 412.526(c)(2), for that period by the number of Medicare discharges paid during that period. Section 412.526(c)(2)(i) describes the method for determining the target amount for cost reporting periods beginning during FY 2015. Section 412.526(c)(2)(ii) specifies that, for cost reporting periods beginning during fiscal years after FY 2015, the target amount will equal the hospital’s target amount for the previous cost reporting period updated by the applicable annual rate-of-increase percentage specified in § 413.40(c)(3) for the subject cost reporting period (79 FR 50197).

For FY 2018, in accordance with proposed § 412.23(j)(2) and existing § 412.526(c)(2)(ii) of the regulations, we are proposing that, for cost reporting periods beginning during FY 2018, the update to the target amount for long-term care neoplastic disease hospitals (that is, hospitals described under proposed § 412.23(j)) would be the applicable annual rate-of-increase percentage specified in § 413.40(c)(3) for FY 2018, which would be equal to the percentage increase projected by the hospital market basket index, which, in this proposed rule, is estimated to be the percentage increase in the proposed 2014-based IPPS operating market basket (that is, the estimate of the market basket rate-of-increase).

Accordingly, for this proposed rule, the proposed update to a long-term care neoplastic disease hospital’s target amount for FY 2018 is 2.9 percent, which is based on IHS Global Insight, Inc.’s 2016 fourth quarter forecast. Furthermore, we are proposing that if more recent data become available for the final rule, we would use that updated data to calculate the IPPS operating market basket update for FY 2018.

B. Proposed Changes to Hospital-Within-Hospital Regulations

On September 1, 1994, we published regulations governing hospitals-within-hospitals (HwHs) to address inappropriate Medicare payments to LTCHs that were effectively units of other hospitals (59 FR 45330). There was concern that the LTCH HwH model was being used by some acute care hospitals paid under the IPPS as a way of inappropriately receiving higher payments for a subset of their cases. Moreover, we stated that the IPPS-exclusion of long-term care “units” may be inconsistent with the statutory scheme, which does not provide for the exclusion from the IPPS of long-term care units.

Therefore, we codified the HwH regulations at 42 CFR 412.23 (currently at § 412.22(e)) for an LTCH HwH that is co-located with another hospital. A co-located hospital is a hospital that occupies space in a building also used by another hospital or in one or more separate buildings located on the same campus as buildings used by another hospital. The regulations at § 412.22(e) required that, to be excluded from the IPPS, long-term care HwHs must have a separate governing body, a chief medical officer, medical staff, and a chief executive officer from that of the hospital with which it is co-located. In addition, the long-term care HwH must meet either of these two criteria: The HwH must perform certain specified basic hospital functions on its
own and not receive them from the host hospital or a third entity that controls both hospitals; or the HwH must receive at least 75 percent of its inpatients from sources other than the co-located hospital. A third option was added to the regulations on September 1, 1995 (60 FR 45778) that allowed long-term care HwHs to demonstrate their separateness by showing that the cost of the services that the hospital obtains under contracts or other agreements with the co-located hospital or a third entity that controls both hospitals is no more than 15 percent of the hospital’s total inpatient operating cost. In 1997, we extended application of the HwH regulations at § 412.22 to all classifications of IPPS-excluded hospitals. Therefore, effective for cost reporting periods beginning on or after October 1, 1997, psychiatric, rehabilitation, cancer, and children’s hospitals that are co-located with another hospital also are generally required to meet the “separateness” criteria at § 412.22(e). In addition, a “grandfathering” provision (that is, hospitals that were IPPS-excluded HwHs before October 1, 1995 are not required to comply with the separateness and control regulations so long as they continue to operate under the same terms and conditions) was added to the regulations at § 412.22(f).

We later modified the grandfathering provision to allow for a grandfathered hospital to make specified changes (for example increasing the number of beds) during particular timeframes, which vary depending on the change the hospital had made. Below we discuss our FY 2018 proposals to make several changes to our HwH regulations.

In this proposed rule, we are proposing to revise our HwH regulations so that the separateness and control requirements would only apply to IPPS-excluded HwHs that are co-located with IPPS hospitals. Under this proposal, any hospital that occupies a building also used by another hospital, or in one or more separate buildings located on the same campus as buildings used by an IPPS hospital, must meet the criteria specified in § 412.22(e)(1) through (e)(3) in order to be excluded from the IPPS. While we are not proposing changes to our HwH regulations for co-located IPPS and IPPS-excluded hospitals, we are seeking public comments on the issue of whether the separateness and control requirements are still necessary for IPPS-excluded HwHs that are co-located with IPPS hospitals, which we would consider for potential future rulemaking.

In this proposed rule, we also are proposing to revise the requirements at § 412.22(e)(1)(v), which outlines performance of basic hospital functions, to make them effective for fiscal years prior to FY 2018. We believe that the requirements in paragraph (e)(1)(v)(A) are generally duplicative of CMS’ interpretative guidance that relate to a number of hospital conditions of participation (CoPs) that are in the regulations for example, 42 CFR 482.21 through 482.27, 482.30, 482.42, 482.43, and 482.45). As such, we are proposing to remove the overlap between the HwH regulations and the CoP Interpretative Guidance from the regulations by sunsetting the requirements in paragraph (e)(1)(v)(A) of § 412.22. (The CoP Interpretative Guidance for hospitals can be found in Appendix A of the State Operations Manual (CMS Pub. 100–07).)

In addition, we are proposing to remove the requirements in paragraph (e)(1)(v)(B) of § 412.22 because we believe these payment requirements could be interpreted to conflict with the requirements under the hospital CoPs, which do not provide for a minimum cost threshold regarding the services the HwH obtains from the hospital with which it is occupying space. We do not believe that this proposed revision would result in a practical change to how HwHs are currently operated because the performance of basic hospital functions requirements at § 412.22(e)(1)(v) are currently addressed under CMS’ Interpretative Guidance for the hospital CoPs. In addition, we do not believe, at this time, that there are payment policy concerns that would justify imposition of regulatory requirements on the performance of basic hospital functions for HwHs that are more stringent than what is addressed under the Interpretative Guidance for the hospital CoPs.

We welcome public comment on these proposals.

C. Critical Access Hospitals (CAHs)

1. Background

Section 1820 of the Act provides for the establishment of Medicare Rural Hospital Flexibility Programs (MRHFPs), under which individual States may designate certain facilities as critical access hospitals (CAHs). Facilities that are so designated and meet the CAH conditions of participation under 42 CFR part 485, subpart F, will be certified as CAHs by CMS. Regulations governing payments to CAHs for services to Medicare beneficiaries are located in 42 CFR part 413.

2. Frontier Community Health Integration Project (FCHIP) Demonstration

Section 123 of the Medicare Improvements for Patients and Providers Act of 2008 (Pub. L. 110–275), as amended by section 3126 of the Affordable Care Act, authorizes a demonstration project to allow eligible entities to develop and test new models for the delivery of health care services in eligible counties in order to improve access to and better integrate the delivery of acute care, extended care...
and other health care services to Medicare beneficiaries. The demonstration is titled “Demonstration Project on Community Health Integration Models in Certain Rural Counties,” and is commonly known as the Frontier Community Health Integration Project (FCHIP) demonstration.

The authorizing statute states the eligibility criteria for entities to be able to participate in the demonstration. An eligible entity, as defined in section 123(d)(1)(B) of Public Law 110–275, as amended, is an MRHFP grantee under section 1820(g) of the Act (that is, a CAH); and is located in a State in which at least 65 percent of the counties in the State are counties that have 6 or less residents per square mile.

The authorizing statute stipulates several other requirements for the demonstration. Section 123(d)(2)(B) of Public Law 110–275, as amended, limits participation in the demonstration to eligible entities in not more than 4 States. Section 123(d)(1)(B) of Public Law 110–275 requires the demonstration project to be conducted for a 3-year period. In addition, section 123(g)(1)(B) of Public Law 110–275 requires that the demonstration be budget neutral. Specifically, this provision states that in conducting the demonstration project, the Secretary shall ensure that the aggregate payments made by the Secretary do not exceed the amount which the Secretary estimates would have been paid if the demonstration project under the section were not implemented. Furthermore, section 123(f) of Public Law 110–275 states that the Secretary may waive such requirements of titles XVIII and XIX of the Act as may be necessary and appropriate for the purpose of carrying out the demonstration project, thus allowing the waiver of Medicare payment rules encompassed in the demonstration.

In January 2014, CMS released a request for applications (RFA) for the FCHIP demonstration. Using 2013 data from the U.S. Census Bureau, CMS identified Alaska, Montana, Nevada, North Dakota, and Wyoming as meeting the statutory eligibility requirement for participation in the demonstration. The RFA solicited CAHs in these five States to participate in the demonstration, stating that participation would be limited to CAHs in four of the States. To apply, CAHs were required to meet the eligibility requirements in the authorizing legislation, and, in addition, to describe a proposal to enhance health services that would complement those currently provided by the CAH and better serve the community’s needs. In addition, in the RFA, CMS interpreted the eligible entity definition in the statute as meaning a CAH that receives funding through the MRHFP. The RFA identified four interventions, under which specific waivers of Medicare payment rules would allow for enhanced payment for telehealth, skilled nursing facility/nursing facility beds, ambulance services, and home health services, respectively. These waivers were formulated with the goal of increasing access to care with no net increase in costs. Ten CAHs were selected for participation in the demonstration, which started on August 1, 2016. These CAHs are located in Montana, Nevada and North Dakota, and they are participating in three of the four interventions identified in the FY 2017 IPPS/LTCH PPS final rule. Eight CAHs are participating in the telehealth intervention, three CAHs are participating in the skilled nursing facility/nursing facility bed intervention, and two CAHs are participating in the ambulance services intervention. Each CAH is allowed to participate in more than one of the interventions. None of the selected CAHs are participants in the home health intervention, which was the fourth intervention proposed in the RFA.

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57064 through 57065), we finalized a policy to address the budget neutrality requirement for the demonstration. As explained in the FY 2017 IPPS/LTCH PPS final rule, we based our selection of CAHs for participation with the goal of maintaining the budget neutrality of the demonstration on its own terms (that is, the demonstration will produce savings from reduced transfers and admissions to other health care providers, thus offsetting any increase in payments resulting from the demonstration). However, because of the small size of this demonstration and uncertainty associated with projected Medicare utilization and costs, we adopted a contingency plan to ensure that the budget neutrality requirement in section 123 of Public Law 110–275 is met. If analysis of claims data for Medicare beneficiaries receiving services at each of the participating CAHs, as well as from other data sources, including cost from other data sources, including cost
impact for any national payment system for FY 2018.

3. Physician Certification Requirement for Payment of Inpatient CAH Services Under Medicare Part A

a. Background

For inpatient CAH services to be payable under Medicare Part A, section 1814(a)(8) of the Act requires that a physician certify that the individual may reasonably be expected to be discharged or transferred to a hospital within 96 hours after admission to the CAH. The regulations implementing this statutory requirement are located at 42 CFR 424.15.

We most recently addressed the 96-hour certification requirement in the FY 2015 IPPS/LTCH PPS final rule (79 FR 50165). In that rule, we finalized a policy regarding the timing of this physician certification requirement. We revised the regulations such that all physician certification requirements must be completed, signed, and documented in the medical record no later than 1 day before the date on which the claim for payment for the inpatient CAH service is submitted. This policy change was effective October 1, 2014. Prior to that revision, our policy, which was in effect during FY 2014, had been that the certification began with the order for inpatient admission and was required to be completed, signed, and documented in the medical record prior to discharge.

In addition to this change regarding the timing of the 96-hour certification requirement, we also provided a general review of this certification requirement in the FY 2015 IPPS/LTCH PPS final rule (79 FR 50165). We stated that because the statutory requirement at section 1814(a)(8) of the Act is based on an expectation, if a physician certifies, in good faith, that an individual may reasonably be expected to be discharged or transferred to a hospital within 96 hours after admission to the CAH and then something unforeseen occurs that causes the individual to stay longer at the CAH, Medicare will pay for the costs of treating that patient and there would not be a problem with regard to the CAH designation as long as that individual’s stay does not cause the CAH to exceed its 96-hour annual average CoP requirement under 42 CFR 485.620(b) (which we note is separate and distinct from the 96-hour physician certification requirement). However, if a physician cannot in good faith certify that an individual may reasonably be expected to be discharged or transferred within 96 hours after admission to the CAH, the CAH will not receive Medicare Part A payment for any portion of that individual’s inpatient stay (79 FR 50165). We further noted that time as an outpatient at the CAH is not included in applying the 96-hour requirement, nor does time in a CAH swing bed, which is being used to provide skilled nursing services, count towards the 96-hour requirement. The clock for the 96 hours only begins once the individual is admitted to the CAH as an inpatient.

b. Notice Regarding Changes to Instructions for the Review of the CAH 96-Hour Certification Requirement

Based on feedback from stakeholders, we have reviewed the CAH 96-hour certification requirement to determine if there are ways to reduce its burden on providers. The requirement is statutory and cannot be modified through regulation. However, we do have discretion to determine how CMS will prioritize monitoring and enforcement of the policy. In order to minimize the burden of documentation submission requirements for CAHs with respect to the 96-hour certification requirement, in this proposed rule, we are providing notice that CMS will direct Quality Improvement Organizations (QIOs), Medicare Administrative Contractors (MACs), the Supplemental Medical Review Contractor (SMRC), and Recovery Audit Contractors (RACs) to make the CAH 96-hour certification requirement a low priority for medical record reviews conducted on or after October 1, 2017. This means that, absent concerns of probable fraud, waste, or abuse with respect to the 96-hour certification requirement, these contractors will not conduct medical record reviews. Reviews by other entities, including, but not limited to, Zone Program Integrity Contractors (ZPICs), the Office of Inspector General, and the Department of Justice will continue as appropriate. Quality reviews and automated reviews (for example, those reviews that do not involve medical records) will also continue as appropriate.

In the past, RACs have never performed medical record reviews for CAH claims, and we will not approve medical record review of CAHs for only the 96-hour certification requirement. We are providing notice that, beginning October 1, 2017, CMS will direct the QIOs, MACs, and the SMRC to make medical record review of CAHs for only the 96-hour certification requirement, a low priority. QIOs and MACs may continue to conduct medical record review of CAH claims for the purposes of verifying other requirements, such as beneficiary complaints, quality of care reviews, higher weighted DRG reviews, readmission reviews, and the requirement that procedures be medically necessary.

Under the revised instructions to contractors, CAHs will not receive any medical record requests from MACs, RACs, QIO, or SMRCs related to the 96-hour certification unless CMS or its contractors find evidence of gaming or a failure to comply with CMS’ provider screening and revalidation requirements or if medical review is needed for other issues. If this occurs, the MACs, RACs, QIO, or SMRCs could also review the 96-hour certification requirement. In addition, if data analysis or other information indicates that possible fraud exists, CAHs may also receive medical record requests for the 96-hour certification requirement.

VIII. Proposed Changes to the Long-Term Care Hospital Prospective Payment System (LTCH PPS) for FY 2018

A. Background of the LTCH PPS

1. Legislative and Regulatory Authority

Section 123 of the Medicare, Medicaid, and SCHIP (State Children’s Health Insurance Program) Balanced Budget Refinement Act of 1999 (BBRA) (Pub. L. 106–113) as amended by section 307(b) of the Medicare, Medicaid, and SCHIP Benefits Improvement and Protection Act of 2000 (BIPA) (Pub. L. 106–554) provides for payment for both the operating and capital-related costs of hospital inpatient stays in long-term care hospitals (LTCHs) under Medicare Part A based on prospectively set rates. The Medicare prospective payment system (PPS) for LTCHs applies to hospitals that are described in section 1886(d)(1)(B)(iv) of the Act, effective for cost reporting periods beginning on or after October 1, 2002.

Section 1886(d)(1)(B)(iv)(I) of the Act originally defined an LTCH as a hospital with an average inpatient length of stay (as determined by the Secretary) of greater than 25 days. Section 1886(d)(1)(B)(iv)(II) of the Act also provided an alternative definition of LTCHs: specifically, a hospital that first received payment under section 1886(d) of the Act in 1986 and had an average inpatient length of stay (as determined by the Secretary of Health and Human Services (the Secretary)) of greater than 20 days and had 80 percent or more of its annual Medicare inpatient discharges with a principal diagnosis that reflected a finding of neoplastic disease in the 12-month cost reporting period ending in FY 1997. However, as discussed below, section 15008 of the 21st Century Cures
Act (Pub. L. 114–255) amended section 1886 of the Act to exclude former "subclause II" LTCHs from payment under the LTCH PPS and created a new category of IPPS-excluded hospitals (long-term Care neoplastic disease hospitals) for hospitals that were formally classified as "subclause (II)" LTCHs.

Section 123 of the BBRA requires the PPS for LTCHs to be a "per discharge" system with a diagnosis-related group (DRG) based patient classification system that reflects the differences in patient resources and costs in LTCHs.

Section 307(b)(1) of the BIPA, among other things, mandates that the Secretary shall examine, and may provide for, adjustments to payments under the LTCH PPS, including adjustments to DRG weights, area wage adjustments, geographic reclassification, outliers, updates, and a disproportionate share adjustment.

In the August 30, 2002 Federal Register, we issued a final rule that implemented the LTCH PPS authorized under the BBRA and BIPA (67 FR 55954). For the initial implementation of the LTCH PPS (FYs 2003 through FY 2007), the system used information from LTCH patient records to classify patients into distinct long-term care diagnosis-related groups (LTC–DRGs) based on clinical characteristics and expected resource needs. Beginning in FY 2008, we adopted the Medicare severity long-term care diagnosis-related groups (MS–LTC–DRGs) as the patient classification system used under the LTCH PPS. Payments are calculated for each MS–LTC–DRG and provisions are made for appropriate payment adjustments. Payment rates under the LTCH PPS are updated annually and published in the Federal Register.

The LTCH PPS replaced the reasonable cost-based payment system under the Tax Equity and Fiscal Responsibility Act of 1982 (TEFRA) (Pub. L. 97–248) for payments for inpatient services provided by an LTCH with a cost reporting period beginning on or after October 1, 2002. (The regulations implementing the TEFRA reasonable cost-based payment provisions are located at 42 CFR part 413.) With the implementation of the LTCH PPS for acute care hospitals authorized by the Social Security Amendments of 1983 (Pub. L. 98–21), which added section 1886(d) to the Act, certain hospitals, including LTCHs, were excluded from the PPS for acute care hospitals and were paid their reasonable costs for inpatient services subject to a specific ceiling on payments was determined by multiplying the hospital’s updated target amount by the number of total current year Medicare discharges. (Generally, in this section of the preamble of this proposed rule, when we refer to discharges, we describe Medicare discharges.) The August 30, 2002 final rule further details the payment policy under the TEFRA system (67 FR 55954).

In the August 30, 2002 final rule, we provided for a 5-year transition period from payments under the TEFRA system to payments under the LTCH PPS. During this 5-year transition period, an LTCH’s total payment under the PPS was based on an increasing percentage of the Federal rate with a corresponding decrease in the percentage of the LTCH PPS payment that is based on reasonable cost concepts, unless an LTCH made a one-time election to be paid based on 100 percent of the Federal rate. Beginning with LTCHs’ cost reporting periods beginning on or after October 1, 2006, total LTCH PPS payments are based on 100 percent of the Federal rate.

In addition, in the August 30, 2002 final rule, we presented an in-depth discussion of the LTCH PPS, including the patient classification system, relative weights, payment rates, additional payments, and the budget neutrality requirements mandated by section 123 of the BBRA. The same final rule that established regulations for the LTCH PPS under 42 CFR part 412, subpart O, also contained LTCH provisions related to covered inpatient services, limitation on charges to beneficiaries, medical review requirements, furnishing of inpatient hospital services directly or under arrangement, and reporting and recordkeeping requirements. We refer readers to the August 30, 2002 final rule for a comprehensive discussion of the research and data that supported the establishment of the LTCH PPS (67 FR 55954).

In the FY 2016 IPPS/LTCH PPS final rule (80 FR 49601 through 49623), we implemented the provisions of the Pathway for Sustainable Growth Rate (SGR) Reform Act of 2013 (Pub. L. 113–67), which mandated the application of the “site neutral” payment rate under the LTCH PPS for discharges that do not meet the statutory criteria for exclusion beginning in FY 2016. For cost reporting periods beginning on or after October 1, 2015, discharges that do not meet certain statutory criteria for exclusion are paid based on the site neutral payment rate. Discharges that do not meet the statutory criteria continue to receive payment based on the LTCH PPS standard Federal payment rate. For more information on the statutory requirements of the Pathway for SGR Reform Act of 2013, we refer readers to the FY 2016 IPPS/LTCH PPS final rule (80 FR 49601 through 49623).

Section 231 of Consolidated Appropriations Act, 2016 (Pub. L. 114–113) amended section 1886(m)(6) of the Act by revising subparagraph (A)(i) and adding new subparagraph (E), which established a temporary exception to the site neutral payment rate for certain severe wound care discharges occurring prior to January 1, 2017, from LTCHs identified by the amendment made by section 4417(a) of the Balanced Budget Act of 1997 that are located in a rural area (as defined in section 1886(d)(2)(D) of the Act) or treated as being so located in accordance with section 1886(d)(8)(E) of the Act.

We implemented the provisions of section 231 of Public Law 114–113, and amended our regulations at 42 CFR 412.522 to reflect those policies, in an interim final rule with comment period (IFC) that appeared in the Federal Register on April 21, 2016 (81 FR 23428 through 23438). In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57068), we finalized the provisions of the April 21, 2016 IFC and made limited modifications of those policies set forth in the April 21, 2016 IFC by revising the definitions of a “wound with morbid obesity” and an “infected wound,” and adding additional ICD–10 diagnosis codes to our list of such codes to identify cases that meet the established definition of a “severe wound” for the six severe wound categories other than the categories of a “wound with morbid obesity” and an “infected wound.” The provisions implementing section 231 of Public Law 114–113 were effective for LTCH discharges from qualifying LTCHs for discharges on or after April 21, 2016, through December 31, 2016. For a full discussion of these provisions, we refer readers to the April 21, 2016 IFC (81 FR 23428) and the FY 2017 IPPS/LTCH PPS final rule (81 FR 57068 through 57075).

The 21st Century Cures Act (“the Cures Act”) (Pub. L. 114–255) contains several provisions that affect the LTCH PPS. Section 15004 of Public Law 114–255 contains provisions that change the moratorium on increasing the number of beds in existing LTCHs and LTCH satellite facilities. We discuss our implementation of the provisions of section 15004 in section VIII.H. of the preamble of this proposed rule. The provisions of section 15004 also included a change to the payment methodology for high-cost outlier payments made to LTCHs. We discuss our proposals related to high-cost
outlier payments in section V.D. of the Addendum of this proposed rule. The provisions of section 15006 of Public Law 114–255 extended various moratoria on the implementation of the 25-percent payment adjustment threshold policy. We discuss our proposals related to the provisions of section 15006 in section VIII.G. of the preamble of this proposed rule. The provisions of section 15007 of Public Law 114–255 revised the requirements of the average length-of-stay criterion for LTCH classification. We discuss our proposals related to the provisions of section 15007 in section VIII.I. of the preamble of this proposed rule. The provisions of section 15008 of Public Law 114–255 changed the classification of certain hospitals. We discuss our proposals related to the provisions of section 15008 in section VIII.J. of the preamble of this proposed rule. The provisions of section 15009 of Public Law 114–255 contain a temporary exception to the site neutral payment rate for certain spinal cord hospitals. We discuss our proposals related to the provisions of section 15009 in section VIII.E. of the preamble of this proposed rule. The provisions of section 15010 of Public Law 114–255 contain a temporary exception to the site neutral payment rate for certain wound care discharges from certain LTCHs. We discuss our proposals related to the provisions of section 15010 in section VIII.F. of the preamble of this proposed rule. In addition, we are proposing to amend 42 CFR 412.500 to include Public Law 114–255 as one of the bases and scope of subpart O of part 412.

2. Criteria for Classification as an LTCH

a. Classification as an LTCH

Under the regulations at § 412.23(e)(1), to qualify to be paid under the LTCH PPS, a hospital must have a provider agreement with Medicare. Furthermore, § 412.23(e)(2)(i), which implements section 1886(d)(1)(B)(iv) of the Act, requires that a hospital have an average Medicare inpatient length of stay of greater than 25 days to be paid under the LTCH PPS. Alternatively, existing § 412.23(e)(2)(ii) states that, for cost reporting periods beginning on or after August 5, 1997, a hospital that was first excluded from the PPS in 1986 and can demonstrate that at least 80 percent of its annual Medicare inpatient discharges in the 12-month cost reporting period ending in FY 1997 have a principal diagnosis that reflects a finding of neoplastic disease must have an inpatient length of stay for all patients, including both Medicare and non-Medicare inpatients, of greater than 20 days (referred to as “subclause (II)” LTCHs). Under our proposed changes to § 412.23(e)(2)(ii) of the regulations to implement the provisions of section 15008 of Public Law 114–255, we are proposing to add a sunset date to subclause (II) LTCHs (which have become a new category of IPPS-excluded hospitals known as long-term care neoplastic disease hospitals). Long-term care neoplastic disease hospitals are discussed in greater detail in section VIII.J. of the preamble of this proposed rule. In addition, in section VIII.I. of the preamble of this proposed rule, we discuss the proposed changes to the calculation of the greater than 25-day average length-of-stay requirement provided by the provisions of section 15008 of Pub. L. 114–255.

b. Hospitals Excluded From the LTCH PPS

The following hospitals are paid under special payment provisions, as described in § 412.22(c) and, therefore, are not subject to the LTCH PPS rules:

• Veterans Administration hospitals.
• Hospitals that are reimbursed under State cost control systems approved under 42 CFR part 403.
• Hospitals that are reimbursed in accordance with demonstration projects authorized under section 402(a) of the Social Security Amendments of 1967 (Pub. L. 90–248) (42 U.S.C. 1395b–1), section 222(a) of the Social Security Amendments of 1972 (Pub. L. 92–603) (42 U.S.C. 1395b–1 (note)) (Statewide all-payer systems, subject to the rate-of-increase test at section 1814(b) of the Act), or section 3201 of the Patient Protection and Affordable Care Act (Pub. L. 111–148 (42 U.S.C. 1315a)).
• Nonparticipating hospitals furnishing emergency services to Medicare beneficiaries.

3. Limitation on Charges to Beneficiaries

In the August 30, 2002 final rule, we presented an in-depth discussion of beneficiary liability under the LTCH PPS (67 FR 55974 through 55975). This discussion was further clarified in the FY 2005 LTCH PPS final rule (69 FR 25676). In keeping with those discussions, if the Medicare payment to the LTCH is the full LTC–DRG payment amount, consistent with other established hospital prospective payment systems, § 412.507 currently provides that an LTCH may not bill a Medicare beneficiary for more than the deductible and coinsurance amounts as specified under §§ 409.82, 409.83, and 409.87 and for items and services specified under § 409.30(a). However, under the LTCH PPS, Medicare will only pay for days for which the beneficiary has coverage until the short-stay outlier (SSO) threshold is exceeded. If the Medicare payment was for a SSO case (§ 412.529), and that payment was less than the full LTC–DRG payment amount because the beneficiary had insufficient remaining Medicare days, the LTCH is currently also permitted to charge the beneficiary for services delivered on those uncovered days (§ 412.507). In the FY 2016 IPPS/LTCH PPS final rule (80 FR 49623), we amended our regulations to expressly limit the charges that may be imposed on beneficiaries whose discharges are paid at the site neutral payment rate under the LTCH PPS.

In section VII.G. of the preamble of the FY 2017 IPPS/LTCH PPS final rule (81 FR 57102), we also amended the existing regulations relating to the limitation on charges to expressly address beneficiary charges for LTCH services provided by subclause (II) LTCHs as part of our refinement of the payment adjustment for subclause II LTCHs under § 412.526. We also amended the regulations under § 412.507 to clarify our existing policy that blended payments made to an LTCH during its transitional period (that is, payment for discharges occurring in cost reporting periods beginning in FY 2016 or 2017) are considered to be site neutral payment rate payments.

4. Administrative Simplification Compliance Act (ASCA) and Health Insurance Portability and Accountability Act (HIPAA) Compliance

Claims submitted to Medicare must comply with both the Administrative Simplification Compliance Act (ASCA) (Pub. L. 107–105), and the Health Insurance Portability and Accountability Act of 1996 (HIPAA) (Pub. L. 104–191). Section 3 of the ASCA generally requires that the Medicare Program deny payment under Part A or Part B for any expenses incurred for items or services for which a claim is submitted other than in an electronic form specified by the Secretary. Section 1862(h) of the Act (as added by section 3(a) of the ASCA) provides that the Secretary shall waive such denial in two specific types of cases, and may also waive such denial in such unusual cases as the Secretary finds appropriate (68 FR 48805). Section 3 of the ASCA operates in the context of the HIPAA regulations, which include, among other provisions, the transactions and code sets standards requirements codified under 45 CFR parts 160 and 162 (generally known as the Transactions Rule). The Transactions Rule requires covered
entities, including covered health care providers, to conduct certain electronic health care transactions according to the applicable transactions and code sets standards.

The Department of Health and Human Services (HHS) has a number of initiatives designed to encourage and support the adoption of health information technology (health IT) and promote nationwide health information exchange to improve health care. The Office of the National Coordinator for Health Information Technology (ONC) leads these efforts in collaboration with other agencies, including CMS and the Office of the Assistant Secretary for Planning and Evaluation (ASPE).

Through a number of activities, including several open government initiatives, HHS is promoting the adoption of health IT products, including electronic health record (EHR) technology certified under the ONC Health IT Certification Program (https://www.healthit.gov/policy-researchers-implementers/about-onc-health-it-certification-program) developed to support secure, interoperable, health information exchange. We believe that the use of certified EHRs by LTCHs (and other types of providers that are ineligible for the Medicare and Medicaid EHR Incentive Programs) can effectively and efficiently help providers improve internal care delivery practices, support the exchange of important information across care partners and during transitions of care, and enable the reporting of electronically specified clinical quality measures (eCQMs) (as described elsewhere in this proposed rule).

In 2015, ONC released a document entitled “Connecting Health and Care for the Nation: A Shared Nationwide Interoperability Roadmap” (available at: https://www.healthit.gov/sites/default/files/hie-interoperability/nationwide-interoperability-roadmap-final-version-1.0.pdf). In the near term, the Roadmap focuses on actions that will enable individuals and providers across the care continuum to send, receive, find, and use a common set of electronic clinical information at the nationwide level by the end of 2017. The Roadmap’s goals also align with the Improving Medicare Post-Acute Care Transformation Act of 2014 (Pub. L. 113–183) (IMPACT Act), which requires assessment data to be standardized and interoperable to allow for exchange of the data. Moreover, the vision described in the Roadmap significantly expands the types of health information, information sources, and information users well beyond clinical information derived from EHRs. The Roadmap identifies four critical pathways that health IT stakeholders should focus on now in order to create a foundation for long-term success: (1) Improve technical standards and implementation guidance for priority data domains and associated elements; (2) rapidly shift and align Federal, State, and commercial payment policies from fee-for-service to value-based models to stimulate the demand for interoperability; (3) clarify and align Federal and State privacy and security requirements that enable interoperability; and (4) align and promote the use of consistent policies and business practices that support interoperability and address those that impede interoperability, in coordination with stakeholders.

In support of the goals of the Roadmap, ONC released the 2017 Interoperability Standards Advisory (ISA) (available at: https://www.healthit.gov/standards-advisory), a coordinated catalog of standards and implementation specifications developed and used to meet specific interoperability needs. The ISA is intended to serve as an industry resource to further the use of interoperable electronic health information exchange.

B. Proposed Medicare Severity Long-Term Care Diagnosis-Related Group (MS–LTC–DRG) Classifications and Relative Weights for FY 2018

1. Background

Section 123 of the BBRA required that the Secretary implement a PPS for LTCHs to replace the cost-based payment system under TEFRA. Section 307(b)(1) of the BIPA modified the requirements of section 123 of the BBRA by requiring that the Secretary examine the feasibility and the impact of basing payment under the LTCH PPS on the use of existing (or refined) hospital DRGs that have been modified to account for different resource use of LTCH patients.

When the LTCH PPS was implemented for cost reporting periods beginning on or after October 1, 2002, we adopted the same DRG patient classification system utilized at that time under the IPPS. As a component of the LTCH PPS, we refer to this patient classification system as the “long-term care diagnosis-related groups (LTC–DRGs).” Although the patient classification system used under both the LTCH PPS and the IPPS are the same, the relative weight is different. The established relative weight methodology and data used under the LTCH PPS result in relative weights under the LTCH PPS that reflect the differences in patient resource use of LTCH patients, consistent with section 123(a)(1) of the BBRA (Pub. L. 106–113).

As part of our efforts to better recognize severity of illness among patients, in the FY 2008 IPPS final rule with comment period (72 FR 47130), the MS–DRGs and the Medicare severity long-term care diagnosis-related groups (MS–LTC–DRGs) were adopted under the IPPS and the LTCH PPS, respectively, effective beginning October 1, 2007 (FY 2008). For a full description of the development, implementation, and rationale for the use of the MS–DRGs and MS–LTC–DRGs, we refer readers to the FY 2008 IPPS final rule with comment period (72 FR 47141 through 47175 and 47277 through 47299). (We note that, in that same final rule, we revised the regulations at § 412.503 to specify that for LTCH discharges occurring on or after October 1, 2007, when applying the provisions of 42 CFR part 412, subpart O applicable to LTCHs for policy descriptions and payment calculations, all references to LTC–DRGs would be considered a reference to MS–LTC–DRGs. For the remainder of this section, we present the discussion in terms of the current MS–LTC–DRG patient classification system unless specifically referring to the previous LTC–DRG patient classification system that was in effect before October 1, 2007.)

The MS–DRGs adopted in FY 2008 represent an increase in the number of DRGs by 207 (that is, from 538 to 745) (72 FR 47171). The MS–DRG classifications are updated annually. There are currently 757 MS–DRG groupings. For FY 2018, there would be 754 MS–DRG groupings based on the proposed changes discussed in section II.F. of the preamble of this FY 2018 IPPS/LTCH PPS proposed rule.

Consistent with section 123 of the BBRA, as amended by section 307(b)(1) of the BIPA, and § 412.515 of the regulations, we use information derived from LTCH PPS patient records to classify LTCH discharges into distinct MS–LTC–DRGs based on clinical characteristics and estimated resource needs. We then assign an appropriate weight to the MS–LTC–DRGs to account for the difference in resource use by patients exhibiting the case complexity and multiple medical problems characteristic of LTCHs.

In this section of the proposed rule, we provide a general summary of our existing methodology for determining the proposed FY 2018 MS–LTC–DRG relative weights under the LTCH PPS.
determining the relative weights for the large number of low-volume MS–LTC–DRGs, we group all of the low-volume MS–LTC–DRGs into five quintiles based on average charges per discharge. Then, under our existing methodology, we account for adjustments made to LTCH PPS standard Federal payments for short-stay outlier (SSO) cases (that is, cases where the covered length of stay at the LTCH is less than or equal to five-sixths of the geometric average length of stay for the MS–LTC–DRG), and we make adjustments to account for nonmonotonically increasing weights, when necessary. The methodology is premised on more severe cases under the MS–LTC–DRG system requiring greater expenditure of medical care resources and higher average charges such that, in the severity levels within a base MS–LTC–DRG, the relative weights should increase monotonically with severity from the lowest to highest severity level. (We discuss each of these components of our MS–LTC–DRG relative weight methodology in greater detail in section VIII.B.3.g of the preamble of this proposed rule.)

2. Patient Classifications Into MS–LTC–DRGs
a. Background
The MS–DRGs (used under the IPPS) and the MS–LTC–DRGs (used under the LTCH PPS) are based on the CMS DRG structure. As noted previously in this section, we refer to the DRGs under the LTCH PPS as MS–LTC–DRGs although they are structurally identical to the MS–DRGs used under the IPPS. The MS–DRGs are organized into 25 major diagnostic categories (MDCs), most of which are based on a particular organ system of the body; the remainder involve multiple organ systems (such as MDC 22, Burns). Within most MDCs, cases are then divided into surgical DRGs and medical DRGs. Surgical DRGs are assigned based on a surgical hierarchy that orders operating room (O.R.) procedures or groups of O.R. procedures by resource intensity. The GROPER software program does not recognize all ICD–10–PCS procedure codes as procedures affecting DRG assignment. That is, procedures that are not surgical (for example, EKGs), or minor surgical procedures (for example, a biopsy of skin and subcutaneous tissue [procedure code 0BJH3ZX]) do not affect the MS–LTC–DRG assignment based on their presence on the claim.

Generally, under the LTCH PPS, a Medicare payment is made at a predetermined specific rate for each discharge that varies based on the MS–LTC–DRG to which a beneficiary’s discharge is assigned. Cases are classified into MS–LTC–DRGs for payment based on the following six data elements:
• Principal diagnosis;
• Additional or secondary diagnoses;
• Surgical procedures;
• Age;
• Sex; and
• Discharge status of the patient.

Currently, for claims submitted using version ASC X12 5010 format, up to 25 diagnosis codes and 25 procedure codes are considered for an MS–DRG assignment. This includes one principal diagnosis and up to 24 secondary diagnoses for severity of illness determinations. (For additional information on the processing of up to 25 diagnosis codes and 25 procedure codes on hospital inpatient claims, we refer readers to section II.G.11.c. of the preamble of the FY 2011 IPPS/LTCH PPS final rule (75 FR 50127).)

Under the HIPAA transactions and code sets regulations at 45 CFR parts 160 and 162, covered entities must comply with the adopted transaction standards and operating rules specified in Subparts I through S of Part 162. Among other requirements, by January 1, 2012, covered entities were required to use the ASC X12 Standards for Electronic Data Interchange Technical Report Type 3—Health Care Claim: Institutional (837), May 2006, ASC X12N/005010X223, and Type 1 Errata to Health Care Claim: Institutional (837) ASC X12 Standards for Electronic Data Interchange Technical Report Type 3, Office of the National Coordinator for Health Information Technology, June 2005 for the health care claims or equivalent encounter information transaction (45 CFR 162.1102(c)). HIPAA requires covered entities to use the applicable medical data code set requirements when conducting HIPAA transactions (45 CFR 162.1000).

Currently, upon the discharge of the patient, the LTCH must assign appropriate diagnosis and procedure codes from the most current version of the International Classification of Diseases, 10th Revision, Clinical Modification (ICD–10–CM) for diagnosis coding and the International Classification of Diseases, 10th Revision, Procedure Coding System (ICD–10–PCS) for inpatient hospital procedure coding, both of which were required to be implemented October 1, 2015 (45 CFR 162.1002(c)(2) and (3)). For additional information on the implementation of the ICD–10 coding system, we refer readers to section ILF.1. of the FY 2017 IPPS/LTCH PPS final rule (81 FR 56787 through 56790) and section ILF.1. of the preamble of
this proposed rule. Additional coding instructions and examples are published in the AHA’s Coding Clinic for ICD-10-CM/PCS.

To create the MS–DRGs (and by extension, the MS–LTC–DRGs), base DRGs were subdivided according to the presence of specific secondary diagnoses designated as complications or comorbidities (CCs) into one, two, or three levels of severity, depending on the impact of the CCs on resources used for those cases. Specifically, there are sets of MS–DRGs that are split into 2 or 3 subgroups based on the presence or absence of a CC or a major complication or comorbidity (MCC). We refer readers to section II.D. of the FY 2008 IPPS final rule with comment period for a detailed discussion about the creation of MS–DRGs based on severity of illness levels (72 FR 47141 through 47175).

MACs enter the clinical and demographic information submitted by LTCHs into their claims processing systems and subject this information to a series of automated screening processes called the Medicare Code Editor (MCE). These screens are designed to identify cases that require further review before assignment into a MS–LTC–DRG can be made. During this process, certain cases are selected for further explanation (74 FR 43949).

After screening through the MCE, each claim is classified into the appropriate MS–LTC–DRG by the Medicare LTCH GROUPER software on the basis of diagnosis and procedure codes and other demographic information (age, sex, and discharge status). The GROUPER software used under the LTCH PPS is the same GROUPER software program used under the IPPS. Following the MS–LTC–DRG assignment, the MAC determines the prospective payment amount by using the Medicare PRICER program, which accounts for hospital-specific adjustments. Under the LTCH PPS, we provide an opportunity for LTCHs to review the MS–LTC–DRG assignments made by the MAC and to submit additional information within a specified timeframe as provided in § 412.513(c).

The GROUPER software is used both to classify past cases to measure relative hospital resource consumption to establish the MS–LTC–DRG relative weights and to classify current cases for purposes of determining payment. The records for all Medicare hospital inpatient discharges are maintained in the MedPAR file. The data in this file are used to evaluate possible MS–DRG and MS–LTC–DRG classification changes and to recalibrate the MS–DRG and MS–LTC–DRG relative weights during our annual update under both the IPPS (§ 412.60(e)) and the LTCH PPS (§ 412.517), respectively.

b. Proposed Changes to the MS–LTC–DRGs for FY 2018

As specified by our regulations at § 412.517(a), which require that the MS–LTC–DRG classifications and relative weights be updated annually, and consistent with our historical practice of using the same patient classification system under the LTCH PPS as is used under the IPPS, we are proposing to update the MS–LTC–DRG classifications effective October 1, 2017, through September 30, 2018 (FY 2018), consistent with the proposed changes to specific MS–DRG classifications presented in section II.F. of the preamble of this proposed rule. Accordingly, the proposed MS–LTC–DRGs for FY 2018 presented in this proposed rule are the same as the proposed MS–DRGs that are being proposed for use under the IPPS for FY 2018. In addition, because the proposed MS–LTC–DRGs for FY 2018 are the same as the proposed MS–DRGs for FY 2018, the other proposed changes that affect MS–DRG (and by extension MS–LTC–DRG) assignments under proposed GROUPER Version 35 as discussed in section II.F. of the preamble of this proposed rule, including the proposed changes to the MCE software and the ICD–10–CM/PCS coding system, also would be applicable under the LTCH PPS for FY 2018.

3. Development of the Proposed FY 2018 MS–LTC–DRG Relative Weights

a. General Overview of the Development of the MS–LTC–DRG Relative Weights

One of the primary goals for the implementation of the LTCH PPS is to pay each LTCH an appropriate amount for the efficient delivery of medical care to Medicare patients. The system must be able to account adequately for each LTCH’s casemix in order to ensure both fair distribution of Medicare payments and access to adequate care for those Medicare patients whose care is more costly (67 FR 55984). To accomplish these goals, we have annually adjusted the LTCH PPS standard Federal prospective payment rate by the applicable relative weight in determining payment to LTCHs for each case. In order to make these annual adjustments under the dual rate LTCH PPS payment structure, beginning with FY 2016, we recalibrate the MS–LTC–DRG relative weighting factors annually using data from inpatient LTCH cases (80 FR 49614 through 49617). Under this policy, the resulting MS–LTC–DRG relative weights would continue to be used to adjust the LTCH PPS standard Federal payment rate when calculating the payment for LTCH PPS standard Federal payment rate cases.

The established methodology to develop the MS–LTC–DRG relative weights is generally consistent with the methodology established when the LTCH PPS was implemented in the August 30, 2002 LTCH PPS final rule (67 FR 55909 through 55991). However, there have been some modifications of our historical procedures for assigning relative weights in cases of zero volume and/or nonmonotonicity resulting from the adoption of the MS–LTC–DRGs, along with the change made in conjunction with the implementation of the dual rate LTCH PPS payment structure beginning in FY 2016 to use LTCH claims data from only LTCH PPS standard Federal payment rate cases (or LTCH PPS cases that would have qualified for payment under the LTCH PPS standard Federal payment rate if the dual rate LTCH PPS payment structure had been in effect at the time of the discharge). (For details on the modifications to our historical procedures for assigning relative weights in cases of zero volume and/or nonmonotonicity, we refer readers to the FY 2008 IPPS final rule with comment period (72 FR 47289 through 47295) and the FY 2009 IPPS final rule (73 FR 48542 through 48550.) For details on the change in our historical methodology to use LTCH claims data only from LTCH PPS standard Federal payment rate cases (or cases that would have qualified for such payment had the LTCH PPS dual payment rate structure been in effect at the time) to determine the MS–LTC–DRG relative weights, we refer readers to the FY 2016 IPPS/LTCH PPS final rule (80 FR 49614 through 49617). Under the LTCH PPS, relative weights for each MS–LTC–DRG are a primary element used to account for the variations in cost per discharge and resource utilization among the payment groups (§ 412.515). To ensure that Medicare patients classified to each MS–LTC–DRG have access to an appropriate level of services and to encourage efficiency, we calculate a relative weight for each MS–LTC–DRG that represents the resources needed by an average inpatient LTCH case in that MS–LTC–DRG. For example, cases in an MS–LTC–DRG with a relative weight of 2 would, on average, cost twice as much to treat as cases in an MS–LTC–DRG with a relative weight of 1.
b. Development of the Proposed MS–LTC–DRG Relative Weights for FY 2018

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57078 through 57079), we presented our policies for the development of the MS–LTC–DRG relative weights for FY 2017.

In this FY 2018 IPPS/LTCH PPS proposed rule, we are proposing to continue to use our current methodology to determine the proposed MS–LTC–DRG relative weights for FY 2018, including the continued application of established policies related to: The hospital-specific relative value methodology, the treatment of severity levels in the proposed MS–LTC–DRGs, proposed low-volume and no-volume MS–LTC–DRGs, proposed adjustments for nonmonotonicity, the steps for calculating the proposed MS–LTC–DRG relative weights with a proposed budget neutrality factor, and only using data from applicable LTCH cases (which includes our policy of only using cases that would meet the criteria for exclusion from the site neutral payment rate (or, for discharges occurring prior to the implementation of the dual rate LTCH PPS payment structure, would have met the criteria for exclusion had those criteria been in effect at the time of the discharge)).

In this section, we present our proposed application of our existing methodology for determining the proposed MS–LTC–DRG relative weights for FY 2018, and we discuss the effects of our proposals concerning the data used to determine the proposed FY 2018 MS–LTC–DRG relative weights on the various components of our existing methodology in the discussion that follows.

c. Data

For this proposed rule, consistent with our proposals regarding the calculation of the proposed MS–LTC–DRG relative weights for FY 2018, we obtained total charges from FY 2016 Medicare LTCH claims data from the December 2016 update of the FY 2016 MedPAR file, which are the best available data at this time, and we are proposing to use proposed Version 35 of the GROUPER to classify LTCH cases. Consistent with our historical practice, we are proposing that if more recent data become available, we would use those data and the finalized Version 35 of the GROUPER in establishing the FY 2018 MS–LTC–DRG relative weights in the final rule. To calculate the proposed FY 2018 MS–LTC–DRG relative weights under the dual rate LTCH PPS payment structure, we are proposing to continue to use applicable LTCH data, which includes our policy of only using cases that meet the criteria for exclusion from the site neutral payment rate (or would have met the criteria had they been in effect at the time of the discharge) (80 FR 49624). Specifically, we began by first evaluating the LTCH claims data in the December 2016 update of the FY 2016 MedPAR file to determine which LTCH cases would meet the criteria for exclusion from the site neutral payment rate under § 412.522(b) had the dual rate LTCH PPS payment structure been in effect at the time of discharge. (We note that while the dual rate LTCH PPS payment structure began to be phased in during FY 2016, due to the statutory requirement that individual LTCHs begin to receive payment under the dual rate LTCH PPS payment structure based on their individual cost reporting periods, there are LTCH discharges that occurred in FY 2016 that would not have been paid under that structure.) We identified the FY 2016 LTCH cases that were not assigned to MS–LTC–DRGs 876, 880, 881, 882, 883, 884, 885, 886, 887, 897, 895, 896, 897, 945 and 946, which identify LTCH cases that do not have a principal diagnosis related to a psychiatric diagnosis or to rehabilitation; and that either—

- The admission to the LTCH was “immediately preceded” by discharge from a subsection (d) hospital and the immediately preceding stay in that subsection (d) hospital included at least 3 days in an ICU, as we define under the ICU criterion; or
- The admission to the LTCH was “immediately preceded” by discharge from a subsection (d) hospital and the claim for the LTCH discharge includes the applicable procedure code that indicates at least 96 hours of ventilator services were provided during the LTCH stay, as we define under the ventilator criterion. Claims data from the FY 2016 MedPAR file that reported ICD–10–PCS procedure code 5A1953Z were used to identify cases involving at least 96 hours of ventilator services in accordance with the ventilator criterion.

We note that for purposes of developing the proposed FY 2018 MS–LTC–DRG relative weights using our current methodology, we did not make any proposals regarding the identification of cases that would have been excluded from the site neutral payment rate under the statutory provisions that provided for temporary exception from the site neutral payment rate under the LTCH PPS for certain severe wound care discharges from certain LTCHs or for certain spinal cord specialty hospitals provisions by sections 15009 and 15010 of Pub. L. 114–255, respectively, had our implementation of that law and the dual rate LTCH PPS payment structure been in effect at the time of the discharge. At this time, it is uncertain how many LTCHs and how many cases in the claims data we are using for this proposed rule would have met the criteria to be excluded from the site neutral payment rate under those exceptions (had the dual rate LTCH PPS payment structure been in effect at the time of the discharge). Therefore, for the remainder of this section, when we refer to LTCH claims only from cases that meet the criteria for exclusion from the site neutral payment rate (or would have met the criteria had the applicable statutes been in effect at the time of the discharge), such data do not include any discharges that would have been paid based on the LTCH PPS standard Federal payment rate under the provisions of sections 15009 and 15010 of Pub. L. 114–255, had the exception been in effect at the time of the discharge.

Furthermore, consistent with our historical methodology, we are excluding any claims in the resulting data set that were submitted by LTCHs that are all-inclusive rate providers and LTCHs that are paid in accordance with demonstration projects authorized under section 402(a) of Pub. L. 90–248 or section 222(a) of Pub. L. 92–603. In addition, consistent with our historical practice and our proposals, we are excluding any Medicare Advantage (Part C) claims in the resulting data. Such claims were identified based on the presence of a GHO Paid indicator value of “1” in the MedPAR file. The claims that remained after these three trims (that is, the applicable LTCH data) were then used to calculate the proposed MS–LTC–DRG relative weights for FY 2018.

In summary, in general, we identified the claims data used in the development of the proposed FY 2018 MS–LTC–DRG relative weights in this proposed rule, as we are proposing, by trimming claims data that would have been paid the site neutral rate had the dual payment rate structure been in effect (except for discharges which would have been excluded from the site neutral payment under the temporary exception for certain severe wound care discharges from certain LTCHs and under the temporary exception for certain spinal cord specialty hospitals), as well as the claims data of 9 all-inclusive rate providers reported in the December 2016 update of the FY 2016 MedPAR file and any Medicare Advantage claims data. (We note that there were no data from any LTCHs that are paid in accordance with a demonstration project reported in the December 2016
update of the FY 2016 MedPAR file. However, had there been we would trim the claims data from those LTCHs as well, in accordance with our established policy.) We are proposing to use the remaining data (that is, the applicable LTCH data) to calculate the proposed relative weights for FY 2018.

d. Hospital-Specific Relative Value (HSRV) Methodology

By nature, LTCHs often specialize in certain areas, such as ventilator-dependent patients. Some case types (MS–LTC–DRGs) may be treated, to a large extent, in hospitals that have, from a perspective of charges, relatively high (or low) charges. This nonrandom distribution of cases with relatively high (or low) charges in specific MS–LTC–DRGs has the potential to inappropriately distort the measure of average charges. To account for the fact that cases may not be randomly distributed across LTCHs, consistent with the methodology we have used since the inception of the LTCH PPS, in this FY 2017 IPPS/LTCH PPS proposed rule, we are proposing to continue to use a hospital-specific relative value (HSRV) methodology to calculate the proposed MS–LTC–DRG relative weights for FY 2018. We believe that this method removes this hospital-specific source of bias in measuring LTCH average charges (67 FR 55985).

Specifically, under this methodology, we are proposing to reduce the impact of the variation in charges across providers on any particular MS–LTC–DRG relative weight by converting each LTCH’s charge for an applicable LTCH case to a relative value based on that LTCH’s average charge for such cases.

Under the HSRV methodology, we standardize charges for each LTCH by converting its charges for each applicable LTCH case to hospital-specific relative charge values and then adjusting those values for the LTCH’s case-mix. The adjustment for case-mix is needed to rescale the hospital-specific relative charge values (which, by definition, average 1.0 for each LTCH).

The average relative weight for an LTCH is its case-mix; therefore, it is reasonable to scale each LTCH’s average relative charge value by its case-mix. In this way, each LTCH’s relative charge value is adjusted by its case-mix to an average that reflects the complexity of the applicable LTCH cases it treats relative to the complexity of the applicable LTCH cases treated by all other LTCHs (the average LTCH PPS case-mix of all applicable LTCH cases across all LTCHs).

In accordance with our established methodology, for FY 2018, we are continuing to standardize charges for each applicable LTCH case by first dividing the adjusted charge for the case (adjusted for SSOs under §412.529 as described in section VIII.B.3.g. (Step 3) of the preamble of this proposed rule) by the average adjusted charge for all applicable LTCH cases at the LTCH in which the case was treated. SSO cases are cases with a length of stay that is less than or equal to five-sixths the average length of stay of the MS–LTC–DRG (§ 412.529 and §412.503). The average adjusted charge reflects the average intensity of the health care services delivered by a particular LTCH and the average cost level of that LTCH.

The resulting ratio is multiplied by that LTCH’s case-mix index to determine the standardized charge for the case.

Multiplying the resulting ratio by the LTCH’s case-mix index accounts for the fact that the same relative charges are given greater weight at an LTCH with higher average costs than they would at a LTCH with low average costs, which is needed to adjust each LTCH’s relative charge value to reflect its case-mix relative to the average case-mix for all LTCHs. By standardizing charges in this manner, we count charges for a Medicare patient at an LTCH with high average charges as less resource intensive than they would be at an LTCH with low average charges. For example, a $10,000 charge for a case at an LTCH with an average adjusted charge of $17,500 reflects a higher level of relative resource use than a $10,000 charge for a case at an LTCH with the same case-mix, but an average adjusted charge of $35,000. We believe that the adjusted charge of an individual case more accurately reflects actual resource use for an individual LTCH because the variation in charges due to systematic differences in the markup of charges among LTCHs is taken into account.

e. Treatment of Severity Levels in Developing the Proposed MS–LTC–DRG Relative Weights

For purposes of determining the MS–LTC–DRG relative weights, under our historical methodology, there are three different categories of MS–DRGs based on volume of cases within specific MS–LTC–DRGs: (1) MS–LTC–DRGs with at least 25 applicable LTCH cases; (2) low-volume MS–LTC–DRGs that contain between 1 and 24 applicable LTCH cases; and (3) no-volume MS–LTC–DRGs that are cross-walked to other MS–LTC–DRGs based on the clinical similarities and assigned the relative weight of the cross-walked MS–LTC–DRG (as described in greater detail below). For FY 2018, we are proposing to continue to use applicable LTCH cases to establish the same volume-based categories to calculate the FY 2018 MS–LTC–DRG relative weights.

In determining the proposed FY 2018 MS–LTC–DRG relative weights, when necessary, as is our longstanding practice, we are proposing to make adjustments to account for nonmonotonicity, as discussed in greater detail later in Step 6 of section VIII.B.3.g. of the preamble of this proposed rule. We refer readers to the discussion in the FY 2010 IPPS/RY 2010 LTCH PPS final rule for our rationale for including an adjustment for nonmonotonicity (74 FR 43953 through 43954).

f. Proposed Low-Volume MS–LTC–DRGs

In order to account for proposed MS–LTC–DRGs with low-volume (that is, with fewer than 25 applicable LTCH cases), consistent with our existing methodology, we are proposing to continue to employ the quintile methodology for proposed low-volume MS–LTC–DRGs, such that we group the proposed “low-volume MS–LTC–DRGs” (that is, proposed MS–LTC–DRGs that contain between 1 and 24 applicable LTCH cases) into one of five categories (quintiles) based on average charges (67 FR 55984 through 55995; 72 FR 47283 through 47288; and 81 FR 25148).

In cases where the initial assignment of a proposed low-volume MS–LTC–DRG to a quintile results in nonmonotonicity within a base-DRG, we are proposing to make adjustments to the resulting low-volume proposed MS–LTC–DRGs to preserve monotonicity, as discussed in detail in section VIII.B.3.g. (Step 6) of the preamble of this proposed rule.

In this proposed rule, based on the best available data (that is, the December 2016 update of the FY 2016 MedPAR files), we identified 261 proposed MS–LTC–DRGs that contained between 1 and 24 applicable LTCH cases. This list of proposed MS–LTC–DRGs was then divided into one of the proposed 5 low-volume quintiles, each containing at least 52 proposed MS–LTC–DRGs (261/5 = 52 with a remainder of 1). We assigned the proposed low-volume MS–LTC–DRGs to specific proposed low-volume quintiles by sorting the proposed low-volume MS–LTC–DRGs in ascending order by average charge in accordance with our established methodology. Based on the
data available for the proposed rule, the number of proposed MS–LTC–DRGs with less than 25 applicable LTCH cases was not evenly divisible by 5 and, therefore, we are proposing to employ our historical methodology for determining which of the proposed low-volume quintiles contain the additional proposed low-volume MS–LTC–DRG. Therefore, we are proposing to use our historical methodology for determining which of the low-volume quintiles should contain the additional proposed low-volume MS–LTC–DRG. Specifically for this proposed rule, after organizing the proposed MS–LTC–DRGs by ascending order by average charge, we would assign the first 52 (1st through 52nd) of proposed low-volume MS–LTC–DRGs (with the lowest average charge) into Quintile 1. The 52 proposed MS–LTC–DRGs with the highest average charge would be assigned into Quintile 5. Because the average charge of the 105th proposed low-volume MS–LTC–DRG in the sorted list was closer to the average charge of the 104th proposed low-volume MS–LTC–DRG (assigned to Quintile 2) than to the average charge of the 106th proposed low-volume MS–LTC–DRG (assigned to Quintile 3), we assigned it to Quintile 2 (such that Quintile 2 contains 53 proposed low-volume MS–LTC–DRGs before any adjustments for nonmonotonicity, as discussed below). This results in 4 of the 5 proposed low-volume quintiles containing 52 proposed MS–LTC–DRGs (Quintiles 1, 3, 4, and 5) and 1 proposed low-volume quintile containing 53 proposed MS–LTC–DRGs (Quintile 2), Table 13A, listed in section VI. of the Addendum to this proposed rule and available via the Internet, lists the composition of the proposed low-volume quintiles for MS–LTC–DRGs for FY 2018.

In order to determine the proposed FY 2018 relative weights for the proposed low-volume MS–LTC–DRGs, consistent with our historical practice, we are proposing to use the five low-volume quintiles described previously. We determined a proposed relative weight and (geometric) average length of stay for each of the five proposed low-volume quintiles using the proposed methodology described in section VIII.B.3.g. of the preamble of this proposed rule. We are proposing to assign the same proposed relative weight and average length of stay to each of the proposed low-volume MS–LTC–DRGs that make up an individual low-volume quintile. We note that, as this system is dynamic, it is possible that the number and specific type of MS–LTC–DRGs with a low-volume of applicable LTCH cases will vary in the future. Furthermore, we note that we continue to monitor the volume (that is, the number of applicable LTCH cases) in the low-volume quintiles to ensure that our quintile assignments used in determining the MS–LTC–DRG relative weights result in appropriate payment for LTCH cases grouped to proposed low-volume MS–LTC–DRGs and do not result in an unintended financial incentive for LTCHs to inappropriately admit these types of cases.

g. Steps for Determining the Proposed FY 2018 MS–LTC–DRG Relative Weights

In this proposed rule, we are proposing to continue to use our current methodology to determine the proposed FY 2018 MS–LTC–DRG relative weights.

In summary, to determine the proposed FY 2018 MS–LTC–DRG relative weights, we are proposing to group applicable LTCH cases to the appropriate proposed MS–LTC–DRG, while taking into account the proposed low-volume quintiles (as described above) and cross-walked proposed no-volume MS–LTC–DRGs (as described later in this section). After establishing the appropriate proposed MS–LTC–DRG (or proposed low-volume quintile), we are proposing to calculate the FY 2018 relative weights by first removing cases with a length of stay of 7 days or less and statistical outliers (Steps 1 and 2 below). Next, we are proposing to adjust the number of applicable LTCH cases in each proposed MS–LTC–DRG (or proposed low-volume quintile) for the effect of SSO cases (Step 3 below). After removing applicable LTCH cases with a length of stay of 7 days or less (Step 1 below) and statistical outliers (Step 2 below), which are the SSO-adjusted applicable LTCH cases and corresponding charges (Step 3 below), we are proposing to calculate proposed “relative adjusted weights” for each proposed MS–LTC–DRG (or proposed low-volume quintile) using the HSRV method.

Step 1—Remove cases with a length of stay of 7 days or less.

The first step in our proposed calculation of the proposed FY 2018 MS–LTC–DRG relative weights is to remove cases with a length of stay of 7 days or less. The MS–LTC–DRG relative weights reflect the average of resources used on representative cases of a specific type. Generally, cases with a length of stay of 7 days or less do not belong in an LTCH because these stays do not fully receive or benefit from treatment that is typical in an LTCH stay, and full resources are often not used in the earlier stages of admission to an LTCH. If we were to include stays of 7 days or less in the computation of the FY 2018 MS–LTC–DRG relative weights, the value of many proposed relative weights would decrease and, therefore, payments would decrease to a level that may no longer be appropriate. We do not believe that it would be appropriate to compromise the integrity of the payment determination for those LTCH cases that actually benefit from and receive a full course of treatment at an LTCH by including data from these very short stays. Therefore, consistent with our existing relative weight methodology, in determining the proposed FY 2018 MS–LTC–DRG relative weights, we are proposing to remove LTCH cases with a length of stay of 7 days or less from applicable LTCH cases. (For additional information on what is removed in this step of the relative weight methodology, we refer readers to 67 FR 55989 and 74 FR 43959.)

Step 2—Remove statistical outliers. The next step in our proposed calculation of the proposed FY 2018 MS–LTC–DRG relative weights is to remove statistical outlier cases from the LTCH cases grouped to the LTCH cases with a length of stay of at least 8 days. Consistent with our existing relative weight methodology, we are proposing to continue to define statistical outliers as cases that are outside of 3.0 standard deviations from the mean of the log distribution of both charges per case and the charges per day for each MS–LTC–DRG. These statistical outliers are removed prior to calculating the proposed relative weights because we believe that they may represent aberrations in the data that distort the measure of average resource use. Including those LTCH cases in the calculation of the proposed relative weights could result in an inaccurate relative weight that does not truly reflect relative resource use among those MS–LTC–DRGs. (For additional information on what is removed in this step of the proposed relative weight methodology, we refer readers to 67 FR 55989 and 74 FR 43959.) After removing cases with a length of stay of 7 days or less and statistical outliers, we are left with applicable LTCH cases that have a length of stay greater than or equal to 8 days. In this proposed rule, we refer to these cases as “trimmed applicable LTCH cases.”

Step 3—Adjust charges for the effects of SSOs. As the next step in the proposed calculation of the proposed FY 2018 MS–LTC–DRG relative weights, consistent with our historical approach, we are proposing to adjust each LTCH’s charges per discharge for those
remaining cases (that is, trimmed applicable LTCH cases) for the effects of SSOs (as defined in § 412.529(a) in conjunction with § 412.503). Specifically, we are proposing to make this adjustment by counting an SSO case as a fraction of a discharge based on the ratio of the length of stay of the case to the average length of stay for the MS–LTC–DRG for non-SSO cases. This has the effect of proportionately reducing the impact of the lower charges for the SSO cases in calculating the average charge for the MS–LTC–DRG. This process produces the same result as if the actual charges per discharge of an SSO case were adjusted to what they would have been had the patient’s length of stay been equal to the average length of stay of the MS–LTC–DRG.

Counting SSO cases as full LTCH cases with no adjustment in determining the proposed FY 2018 MS–LTC–DRG relative weights would lower the proposed FY 2018 MS–LTC–DRG relative weight for affected MS–LTC–DRGs because the relatively lower charges of the SSO cases would bring down the average charge for all cases within a MS–LTC–DRG. This would result in an “underpayment” for non-SSO cases and an “overpayment” for SSO cases. Therefore, we are proposing to continue to adjust for SSO cases under § 412.529 in this manner because it would result in more appropriate payments for all LTCH PPS standard Federal payment rate cases. (For additional information on this step of the relative weight methodology, we refer readers to 67 FR 55989 and 74 FR 43959.)

**Step 4**—Calculate the proposed FY 2018 MS–LTC–DRG relative weights on an iterative basis.

Consistent with our historical relative weight methodology, we are proposing to calculate the proposed FY 2018 MS–LTC–DRG relative weights using the HSRV methodology, which is an iterative process. First, for each SSO-adjusted trimmed applicable LTCH case, we calculate a hospital-specific relative charge value by dividing the charge per discharge after adjusting for SSOs of the LTCH case (from Step 3) by the average charge per SSO-adjusted discharge for the LTCH in which the case occurred. The resulting ratio is then multiplied by the LTCH’s case-mix index to produce an adjusted hospital-specific relative charge value for the case. We used an initial case-mix index value of 1.0 for each LTCH. For each proposed MS–LTC–DRG, we calculated the proposed FY 2018 relative weight by dividing the SSO-adjusted average of the hospital-specific relative charge values for applicable LTCH cases for the proposed MS–LTC–DRG (that is, the sum of the hospital-specific relative charge value from above divided by the sum of equivalent cases from Step 3 for each proposed MS–LTC–DRG) by the overall SSO-adjusted average hospital-specific relative charge value across all applicable LTCH cases for all LTCHs (that is, the sum of the hospital-specific relative charge value from above divided by the sum of equivalent applicable LTCH cases from Step 3 for each proposed MS–LTC–DRG). Using these recalculated MS–LTC–DRG relative weights, each LTCH’s average relative weight for all of its SSO-adjusted trimmed applicable LTCH cases (that is, its case-mix) was calculated by dividing the sum of all the LTCH’s MS–LTC–DRG relative weights by its total number of SSO-adjusted trimmed applicable LTCH cases. The LTCH’s hospital-specific relative charge values (from previous) are then multiplied by the hospital-specific case-mix indexes. The hospital-specific case-mix adjusted relative charge values are then used to calculate a new set of proposed MS–LTC–DRG relative weights across all LTCHs. This iterative process continued until there was convergence between the relative weights produced at adjacent steps, for example, when the maximum difference was less than 0.0001.

**Step 5**—Determine a proposed FY 2018 relative weight for MS–LTC–DRGs with no applicable LTCH cases.

Using the trimmed applicable LTCH cases, consistent with our historical methodology, we identified the proposed MS–LTC–DRGs for which there were no claims in the December 2016 update of the FY 2016 MedPAR file and, therefore, for which no charge data was available for these proposed MS–LTC–DRGs. Because patients with a number of the diagnoses under these proposed MS–LTC–DRGs may be treated at LTCHs, consistent with our historical methodology, we are generally proposing to assign a proposed relative weight to each of the proposed no-volume MS–LTC–DRGs based on clinical similarity and relative costliness with the exception of “transplant” proposed MS–LTC–DRGs, “error” proposed MS–LTC–DRGs, and proposed MS–LTC–DRGs that indicate a principal diagnosis related to a psychiatric diagnosis or rehabilitation (referred to as the “psychiatric or rehabilitation” MS–LTC–DRGs), as discussed later in this section of this proposed rule. (For additional information on this step of the proposed relative weight methodology, we refer readers to 67 FR 55991 and 74 FR 43959 through 43960.)

We are proposing to cross-walk each proposed no-volume MS–LTC–DRG to another proposed MS–LTC–DRG for which we calculated a proposed relative weight (determined in accordance with the methodology described above). Then, the “no-volume” proposed MS–LTC–DRG was assigned the same proposed relative weight (and average length of stay) of the proposed MS–LTC–DRG to which it was cross-walked (as described in greater detail in this section of this proposed rule).

Of the 754 proposed MS–LTC–DRGs for FY 2018, we identified 351 MS–LTC–DRGs for which there are no trimmed applicable LTCH cases (the number identified includes the 8 “transplant” MS–LTC–DRGs, the 2 “error” MS–LTC–DRGs, and the 15 “psychiatric or rehabilitation” MS–LTC–DRGs, which are discussed below). We are proposing to assign proposed relative weights to each of the 351 no-volume proposed MS–LTC–DRGs that contained trimmed applicable LTCH cases based on clinical similarity and relative costliness to 1 of the remaining 403 (754 – 351 = 403) proposed MS–LTC–DRGs for which we calculated proposed relative weights based on the trimmed applicable LTCH cases in the FY 2016 MedPAR file data using the steps described previously. (For the remainder of this discussion, we refer to the “cross-walked” proposed MS–LTC–DRGs as the proposed MS–LTC–DRGs to which we cross-walked 1 of the 351 “no-volume” proposed MS–LTC–DRGs.)

Then, we are generally proposing to assign the 351 no-volume proposed MS–LTC–DRGs the proposed relative weight of the cross-walked proposed MS–LTC–DRG. (As explained below in Step 6, when necessary, we made adjustments to account for nonmonotonicity.)

We cross-walked the no-volume proposed MS–LTC–DRG to a proposed MS–LTC–DRG for which we calculated proposed relative weights based on the December 2016 update of the FY 2016 MedPAR file, and to which it is similar clinically in intensity of use of resources and relative costliness as determined by criteria such as care provided during the period of time surrounding surgery, surgical approach (if applicable), length of time of surgical procedure, postoperative care, and length of stay. (For more details on our process for evaluating relative costliness, we refer readers to the FY 2010 IPPS/RY 2010 LTCH PPS final rule (73 FR 48543).) We believe in the rare event that there would be a few LTCH cases grouped to one of the no-volume MS–LTC–DRGs in FY 2017, the relative weights assigned...
based on the cross-walked MS–LTC–DRGs would result in an appropriate LTCH PPS payment because the crosswalks, which are based on clinical similarity and relative costliness, would be expected to generally require equivalent relative resource use.

We then assigned the proposed relative weight of the cross-walked proposed MS–LTC–DRG as the proposed relative weight for the no-volume proposed MS–LTC–DRG such that both of these proposed MS–LTC–DRGs (that is, the no-volume proposed MS–LTC–DRG and the cross-walked proposed MS–LTC–DRG) have the same proposed relative weight (and average length of stay) for FY 2018. We note that, if the proposed cross-walked MS–LTC–DRG had 25 applicable LTCH cases or more, its proposed relative weight (calculated using the methodology described in Steps 1 through 4 above) is assigned to the no-volume proposed MS–LTC–DRG as well. Similarly, if the proposed MS–LTC–DRG to which the no-volume proposed MS–LTC–DRG was cross-walked had 24 or less cases and, therefore, is designated to 1 of the proposed low-volume quintiles for purposes of determining the proposed relative weights, we assigned the proposed relative weight of the applicable proposed low-volume quintile to the no-volume proposed MS–LTC–DRG such that both of these proposed MS–LTC–DRGs (that is, the no-volume proposed MS–LTC–DRG and the cross-walked proposed MS–LTC–DRG) have the same proposed relative weight for FY 2018. (As we noted previously, in the infrequent case where nonmonotonicity involving a no-volume proposed MS–LTC–DRG resulted, additional adjustments as described in Step 6 are required in order to maintain monotonically increasing proposed relative weights.)

For this proposed rule, a list of the no-volume proposed MS–LTC–DRGs and the proposed MS–LTC–DRGs to which each was cross-walked (that is, the proposed cross-walked MS–LTC–DRGs) for FY 2018 is shown in Table 13B, which is listed in section VI. of the Addendum to this proposed rule and is available via the Internet on the CMS Web site.

Example: There were no trimmed applicable LTCH cases in the FY 2016 MedPAR file that we are proposing to use for this proposed rule for proposed MS–LTC–DRG 061 (Acute Ischemic Stroke with Use of Thrombolytic Agent with MCC). We determined that proposed MS–LTC–DRG 070 (Non-specific Cerebrovascular Disorders with MCC) is similar clinically and based on resource use to proposed MS–LTC–DRG 061. Therefore, we assigned the same proposed relative weight (and average length of stay) of proposed MS–LTC–DRG 70 of 0.8890 for FY 2018 to proposed MS–LTC–DRG 061 (we refer readers to Table 11, which is listed in section VI. of the Addendum to this proposed rule and is available via the Internet on the CMS Web site).

Again, we note that, as this system is dynamic, it is entirely possible that the number of MS–LTC–DRGs with no volume will vary in the future. Consistent with our historical practice, we are proposing to use the most recent available claims data to identify the trimmed applicable LTCH cases from which we determined the proposed relative weights in this proposed rule.

For FY 2018, consistent with our historical relative weight methodology, we are proposing to establish a relative weight of 0.0000 for the following transplant MS–LTC–DRGs: Heart Transplant or Implant of Heart Assist System with MCC (MS–LTC–DRG 001); Heart Transplant or Implant of Heart Assist System without MCC (MS–LTC–DRG 002); Liver Transplant with MCC or Intestinal Transplant (MS–LTC–DRG 005); Liver Transplant without MCC (MS–LTC–DRG 006); Lung Transplant (MS–LTC–DRG 007); Simultaneous Pancreas/Kidney Transplant (MS–LTC–DRG 008); Pancreas Transplant (MS–LTC–DRG 010); and Kidney Transplant (MS–LTC–DRG 652). This is because Medicare only covers these procedures if they are performed at a hospital that has been certified for the specific procedures by Medicare and presently no LTCH has been so certified. At the present time, we include these eight transplant proposed MS–LTC–DRGs in the GROUPER program for administrative purposes only. Because we use the same GROUPER program for LTCHs as is used under the IPPS, removing these MS–LTC–DRGs would be administratively burdensome. (For additional information regarding our treatment of transplant MS–LTC–DRGs, we refer readers to the FY 2010 LTCH PPS final rule (74 FR 43964).)

In addition, consistent with our historical policy, we are proposing to establish a relative weight of 0.0000 for the 2 “error” MS–LTC–DRGs (that is, MS–LTC–DRG 998 (Principal Diagnosis Invalid as Discharge Diagnosis) and MS–LTC–DRG 999 (Ungroupable)) because applicable LTCH cases grouped to these MS–LTC–DRGs cannot be properly assigned to an MS–LTC–DRG according to the grouping logic.

In this proposed rule, consistent with our practice in FYs 2016 and 2017, we are proposing to establish a proposed relative weight for FY 2018 equal to the respective FY 2015 relative weight of the MS–LTC–DRGs for the following “psychiatric and rehabilitation” proposed MS–LTC–DRGs: proposed MS–LTC–DRG 876 (O.R. Procedure with Principal Diagnoses of Mental Illness); proposed MS–LTC–DRG 880 (Acute Adjustment Reaction & Psychosocial Dysfunction); proposed MS–LTC–DRG 881 (Depressive Disorders); proposed MS–LTC–DRG 882 (Neuroses Except Depressive); proposed MS–LTC–DRG 883 (Disorders of Personality & Impulse Control); proposed MS–LTC–DRG 884 (Organic Disturbances & Mental Retardation); proposed MS–LTC–DRG 885 (Psychoses); proposed MS–LTC–DRG 886 (Behavioral & Developmental Disorders); proposed MS–LTC–DRG 887 (Other Mental Disorder Diagnoses); proposed MS–LTC–DRG 894 (Alcohol/Drug Abuse or Dependence, Left Ama); proposed MS–LTC–DRG 895 (Alcohol/Drug Abuse or Dependence, with Rehabilitation Therapy); proposed MS–LTC–DRG 896 (Alcohol/Drug Abuse or Dependence, without Rehabilitation Therapy with MCC); proposed MS–LTC–DRG 897 (Alcohol/Drug Abuse or Dependence, without Rehabilitation Therapy without MCC); proposed MS–LTC–DRG 945 (Rehabilitation with CC/MCC); and proposed MS–LTC–DRG 946 (Rehabilitation without CC/MCC). As we discussed when we implemented the dual rate LTCH PPS payment structure, LTCH discharges that are grouped to these 15 “psychiatric and rehabilitation” proposed MS–LTC–DRGs do not meet the criteria for exclusion from the site neutral payment rate. As such, under the criteria for a principal diagnosis relating to a psychiatric diagnosis or to rehabilitation, there are no applicable LTCH cases to use in calculating a dual rate LTCH PPS payment structure, LTCH discharges that are grouped to these 15 “psychiatric and rehabilitation” proposed MS–LTC–DRGs. In other words, any LTCH PPS discharges grouped to any of the 15 “psychiatric and rehabilitation” proposed MS–LTC–DRGs would always be paid at the site neutral payment rate, and, therefore, those proposed MS–LTC–DRGs would never include any LTCH cases that meet the criteria for exclusion from the site neutral payment rate.
neutral payment rate. However, section 1886(m)(6)(B) of the Act establishes a transitional payment method for cases that would be paid at the site neutral payment rate for LTCH discharges occurring in cost reporting periods beginning during FY 2016 or FY 2017. Under the transitional payment method for site neutral payment rate cases, for LTCH discharges occurring in cost reporting periods beginning on or after October 1, 2016, and on or before September 30, 2017, site neutral payment rate cases are paid a blended payment rate, calculated as 50 percent of the applicable site neutral payment rate amount for the discharge and 50 percent of the applicable LTCH PPS standard Federal payment rate. Because the LTCH PPS standard Federal payment rate is based on the relative weight of the MS–LTC–DRG, in order to determine the transitional blended payment for site neutral payment rate cases grouped to one of the “psychiatric or rehabilitation” proposed MS–LTC–DRGs in FY 2018, we assigned a proposed relative weight to these proposed MS–LTC–DRGs for FY 2018 that is the same as the FY 2015 relative weight (which is also the same as the FY 2016 relative weight). We believe that using the respective FY 2015 relative weight for each of the “psychiatric or rehabilitation” proposed MS–LTC–DRGs results in appropriate payments for LTCH cases that are paid at the site neutral payment rate under the transition policy provided by the statute because there are no clinically similar proposed MS–LTC–DRGs for which we were able to determine proposed relative weights based on applicable LTCH cases in the FY 2016 MedPAR file data using the steps described above. Furthermore, we believe that it would be administratively burdensome and introduce unnecessary complexity to the proposed MS–LTC–DRG relative weight calculation to use the LTCH discharges in the MedPAR file data to calculate a proposed relative weight for those 15 “psychiatric or rehabilitation” proposed MS–LTC–DRGs to be used for the sole purpose of determining half of the transitional blended payment for site neutral payment rate cases during the transition period (80 FR 49631 through 49632).

In summary, for FY 2018, we are proposing to establish a proposed relative weight (and average length of stay thresholds) equal to the respective FY 2015 relative weight of the proposed MS–LTC–DRGs for the 15 “psychiatric or rehabilitation” proposed MS–LTC–DRGs listed previously (that is, proposed MS–LTC–DRGs 876, 880, 881, 882, 883, 884, 885, 886, 887, 894, 895, 896, 897, 945, and 946). Table 11, which is listed in section VI. of the Addendum to this proposed rule and is available via the Internet on the CMS Web site, reflects this proposed policy.  

Step 6—Adjust the proposed FY 2018 MS–LTC–DRG relative weights to account for nonmonotonically increasing relative weights. The MS–DRGs contain base DRGs that have been subdivided into one, two, or three severity of illness levels. Where there are three severity levels, the most severe level has at least one secondary diagnosis code that is referred to as an MCC (that is, major complication or comorbidity). The next lower severity level contains cases with at least one secondary diagnosis code that is a CC (that is, complication or comorbidity). Those cases without an MCC or a CC are referred to as “without CC/MCC.” When data do not support the creation of three severity levels, the base MS–DRG is subdivided into either two levels or the base MS–DRG subdivided. The two-level subdivisions may consist of the MS–DRG with CC/MCC and the MS–DRG without CC/MCC. Alternatively, the other type of two-level subdivision may consist of the MS–DRG with MCC and the MS–DRG without MCC.

In those base MS–LTC–DRGs that are split into either two or three severity levels, cases classified into the “without CC/MCC” MS–LTC–DRG are expected to have a lower resource use (and lower costs) than the “with CC/MCC” MS–LTC–DRG (in the case of a two-level split) or both the “with CC” and the “with MCC” MS–LTC–DRGs (in the case of a three-level split). That is, theoretically, cases that are more severe typically require greater expenditure of medical care resources and would result in higher average charges. Therefore, in the three severity levels, relative weights should increase by severity, from lowest to highest. If the relative weights decrease as severity increases (that is, if within a base MS–LTC–DRG, an MS–LTC–DRG with CC has a higher relative weight than one with MCC, or the MS–LTC–DRG “without CC/MCC” has a higher relative weight than either of the others), they are nonmonotonic. We continue to believe that utilizing nonmonotonic relative weights to adjust Medicare payments would result in inappropriate payments because the payment for the cases in the higher severity level in a base MS–LTC–DRG (which are generally expected to have lower resource use and costs) would be lower than the payment for cases in a lower severity level within the same base MS–LTC–DRG (which are generally expected to have lower resource use and costs). Therefore, in determining the proposed FY 2018 MS–LTC–DRG relative weights, consistent with our historical methodology, we are proposing to continue to combine MS–LTC–DRG severity levels within a base MS–LTC–DRG for the purpose of computing a relative weight when necessary to ensure that monotonicity is maintained. For a comprehensive description of our existing methodology to adjust for nonmonotonicity, we refer readers to the FY 2010 IPPS/RY 2010 LTCH PPS final rule (74 FR 43964 through 43966). Any adjustments for nonmonotonicity that were made in determining the proposed FY 2018 MS–LTC–DRG relative weights in this proposed rule by applying this methodology are denoted in Table 11, which is listed in section VI. of the Addendum to this proposed rule and is available via the Internet on the CMS Web site.  

Step 7—Calculate the proposed FY 2018 MS–LTC–DRG reclassification and recalibration budget neutrality factor. In accordance with the regulations at §412.517(b) (in conjunction with §412.503), the annual update to the MS–LTC–DRG classifications and relative weights is done in a budget neutral manner such that estimated aggregate LTCH PPS payments would be unaffected, that is, would be neither greater than nor less than the estimated aggregate LTCH PPS payments that would have been made without the MS–LTC–DRG classification and relative weight changes. (For a detailed discussion on the establishment of the budget neutrality requirement for the annual update of the MS–LTC–DRG classifications and relative weights, we refer readers to the FY 2008 LTCH PPS final rule (72 FR 26681 and 26682)). The MS–LTC–DRG classifications and relative weights are updated annually based on the most recent available LTCH claims data to reflect changes in relative LTCH resource use (§412.517(a) in conjunction with §412.503). To achieve the budget neutrality requirement at §412.517(b), under our established methodology, for each annual update, the MS–LTC–DRG relative weights are uniformly adjusted to ensure that estimated aggregate payments under the LTCH PPS would not be affected (that is, decreased or increased). Consistent with that provision, we are proposing to update the MS–LTC–DRG classifications and relative weights for FY 2018 based on the most recent available LTCH data for applicable LTCH cases, and continue to apply a budget neutrality adjustment in
determining the proposed FY 2018 MS–LTC–DRG relative weights. In this FY 2018 IPPS/LTCH PPS proposed rule, to ensure budget neutrality in the update to the MS–LTC–DRG classifications and relative weights under § 412.517(b), we are proposing to continue to use our established two-step budget neutrality methodology.

To calculate the proposed normalization factor for FY 2018, we grouped applicable LTCH cases using the proposed FY 2018 Version 35 GROUPER, and the recalibrated proposed FY 2018 MS–LTC–DRG relative weights to calculate the average case-mix index (CMI); we grouped the same applicable LTCH cases using the FY 2017 GROUPER Version 34 and MS–LTC–DRG relative weights and calculated the average CMI; and computed the ratio by dividing the average CMI for FY 2017 by the average CMI proposed for FY 2018. That ratio is the proposed normalization factor. Because the calculation of the proposed normalization factor does not involve the proposed relative weights for the proposed MS–LTC–DRGs that contained applicable LTCH cases to calculate the average CMI, any low-volume proposed MS–LTC–DRGs are included in the calculation (and the proposed MS–LTC–DRGs with no applicable LTCH cases are not included in the calculation).

To calculate the proposed budget neutrality adjustment factor, we simulated estimated total FY 2018 LTCH PPS standard Federal payment rate payments for applicable LTCH cases using the proposed FY 2018 normalized relative weights and proposed GROUPER Version 35; simulated estimated total FY 2017 LTCH PPS standard Federal payment rate payments for applicable LTCH cases using the FY 2017 MS–LTC–DRG relative weights and the FY 2017 GROUPER Version 34; and calculated the ratio of these estimated total payments by dividing the simulated estimated total LTCH PPS standard Federal payment rate payments for FY 2017 by the simulated estimated total LTCH PPS standard Federal payment rate payments for FY 2018. The resulting ratio is the proposed budget neutrality adjustment factor. The calculation of the proposed budget neutrality factor involves the proposed relative weights for the LTCH cases used in the payment simulation, which includes any cases grouped to low-volume proposed MS–LTC–DRGs or to proposed MS–LTC–DRGs with no applicable LTCH cases, and generally does not include the MS–LTC–DRG relative weights for cases grouped to a proposed MS–LTC–DRG with no applicable LTCH cases.

(Occasionally, a few LTCH cases (that is, those with a covered length of stay of 7 days or less, which are removed from the proposed relative weight calculation in step 2) that are grouped to a proposed MS–LTC–DRG with no applicable LTCH cases are included in the payment simulations used to calculate the proposed budget neutrality factor. However, the number and payment amount of such cases have a negligible impact on the proposed budget neutrality factor calculation).

In this proposed rule, to ensure budget neutrality in the update to the MS–LTC–DRG classifications and relative weights under § 412.517(b), we are proposing to continue to use our established two-step budget neutrality methodology. Therefore, in this proposed rule, in the first step of our proposed MS–LTC–DRG budget neutrality methodology, for FY 2018, we are proposing to calculate and apply a proposed normalization factor to the recalibrated proposed relative weights (the result of Steps 1 through 6, discussed previously) to ensure that estimated payments are not affected by changes in the composition of case types or the proposed changes to the classification system. That is, the proposed normalization adjustment is intended to ensure that the recalibration of the proposed MS–LTC–DRG relative weights (that is, the process itself) neither increases nor decreases the average case-mix index.

To calculate the proposed normalization factor for FY 2018 (the first step of our proposed budget neutrality methodology), we used the following three steps: (1.a.) used the most recent available applicable LTCH cases from the most recent available data (that is, LTCH discharges from the FY 2016 MedPAR file) and grouped them using the proposed FY 2018 GROUPER (that is, proposed Version 35 for FY 2018) and the recalibrated proposed FY 2018 MS–LTC–DRG relative weights (determined in Steps 1 through 6 above) to calculate the average CMI (the value determined in Step 1.b. grouped the same applicable LTCH cases as are used in Step 1.a.) using the FY 2017 GROUPER (Version 34) and FY 2017 MS–LTC–DRG relative weights and calculated the average case-mix index; and (1.c.) computed the ratio of these average case-mix indexes by dividing the average CMI for FY 2017 (determined in Step 1.b.) by the average case-mix index for FY 2018 (determined in Step 1.a.). As a result, in determining the proposed MS–LTC–DRG relative weight for each case, the recalibrated proposed MS–LTC–DRG relative weight is multiplied by the proposed normalization factor of 1.28875 (determined in Step 1.c.) in the first step of the proposed budget neutrality methodology, which produced “normalized relative weights.”

In the second step of our proposed MS–LTC–DRG budget neutrality methodology, we calculate a second proposed budget neutrality factor consisting of the ratio of estimated aggregate FY 2018 LTCH PPS standard Federal payment rate payments for applicable LTCH cases (the sum of all calculations under Step 1.a. mentioned previously) after reclassification and recalibration to estimated aggregate payments for FY 2018 LTCH PPS standard Federal payment rate payments for applicable LTCH cases before reclassification and recalibration (that is, the sum of all calculations under Step 1.b. mentioned previously).

That is, for this proposed rule, for FY 2018, under the second step of the proposed budget neutrality methodology, we are proposing to determine the proposed budget neutrality factor using the following three steps: (2.a.) simulated estimated total FY 2018 LTCH PPS standard Federal payment rate payments for applicable LTCH cases using the proposed normalized relative weights for FY 2018 and proposed GROUPER Version 35 (as described above); (2.b.) simulated estimated total FY 2017 LTCH PPS standard Federal payment rate payments for applicable LTCH cases using the FY 2017 GROUPER Version 34 and the FY 2017 MS–LTC–DRG relative weights in Table 11 of the FY 2017 IPPS/LTCH PPS final rule available on the Internet, as described in section VI. of the Addendum of that final rule; and (2.c.) calculated the ratio of these estimated total payments by dividing the value determined in Step 2.b. by the value determined in Step 2.a. In determining the proposed FY 2018 MS–LTC–DRG relative weights, each normalized proposed relative weight is then multiplied by a budget neutrality factor of 0.9866449. The Addendum to this proposed rule and is available via the Internet on the CMS Web site, lists the proposed MS–
LTC–DRGs and their respective proposed relative weights, geometric mean length of stay, and five-sixths of the geometric mean length of stay (used to identify SSO cases under § 412.529(a) for FY 2018.

C. Proposed Changes to the LTCH PPS Payment Rates and Other Proposed Changes to the LTCH PPS for FY 2018

1. Overview of Development of the LTCH PPS Standard Federal Payment Rates

The basic methodology for determining LTCH PPS standard Federal payment rates is currently set forth at 42 CFR 412.515 through 412.536. In this section, we discuss the factors that we are proposing to use to update the LTCH PPS standard Federal payment rate for FY 2018, that is, effective for LTCH discharges occurring on or after October 1, 2017 through September 30, 2018. Under the dual rate LTCH PPS payment structure required by statute, beginning with discharges in cost reporting periods beginning in FY 2016, only LTCH discharges that meet the criteria for exclusion from the site neutral payment rate are paid based on the LTCH PPS standard Federal payment rate specified at § 412.523. (For additional details on our finalized policies related to the dual rate LTCH PPS payment structure required by statute, we refer readers to the FY 2016 IPPS/LTCH PPS final rule (80 FR 49601 through 49623).)

Prior to the implementation of the dual payment rate system in FY 2016, all LTCHs were paid similarly to those now exempt from the site neutral payment rate. That legacy payment rate was called the standard Federal rate. For details on the development of the initial standard Federal rate for FY 2003, we refer readers to the August 30, 2002 LTCH PPS final rule (67 FR 56027 through 56037). For subsequent updates to the standard Federal rate (FYs 2003 through 2015), we refer readers to the FY 2016 IPPS/LTCH PPS final rule (80 FR 49601 through 49623).

Prior to the implementation of the dual payment rate system, the standard Federal rate was used to determine the operating and capital-related costs. We adopted the criteria for exclusion from the site neutral payment rate and capital-related costs. We adopted the proposed budget neutrality adjustment stemming from our proposed change to the SSO payment methodology (as discussed in VII.D. of the preamble of this proposed rule).

2. Proposed FY 2018 LTCH PPS Standard Federal Payment Rate Annual Market Basket Update

a. Overview

Historically, the Medicare program has used a market basket to account for input price increases in the services furnished by providers. The market basket used for the LTCH PPS includes both operating and capital related costs of LTCHs because the LTCH PPS uses a single payment rate for both operating and capital-related costs. We adopted the 2013-based LTCH-specific market basket for use under the LTCH PPS beginning in FY 2017 (61 FR 57101 through 57102). For additional details on the history of the development of the market basket used under the LTCH PPS, we refer readers to the FY 2013 IPPS/LTCH PPS final rule (77 FR 53467 through 53476), and for a complete discussion of the LTCH market basket and a description of the methodologies used to determine the operating and capital-related portions of the 2013-based LTCH market basket, we refer readers to section VII.D. of the preamble of the FY 2017 IPPS/LTCH PPS proposed and final rules.

Section 3401(c) of the Affordable Care Act provides for certain adjustments to any annual update to the LTCH PPS standard Federal payment rate and refers to the timeframes associated with such adjustments as a “rate year” (which are discussed in more detail in section VIII.C.2.b. of the preamble of this proposed rule.) We note that because the annual update to the LTCH PPS policies, rates, and factors now occurs on October 1, we adopted the term “fiscal year” (FY) rather than “rate year” (RY) under the LTCH PPS beginning October 1, 2010, to conform with the standard definition of the Federal fiscal year (October 1 through September 30) used by other PPSs, such as the IPPS (75 FR 50396 through 50397). Although the language of sections 3404(a), 3401(c), 10319, and 1105(b) of the Affordable Care Act refers to years 2010 and thereafter under the LTCH PPS as “rate year,” consistent with our change in the terminology used under the LTCH PPS from “rate year” to “fiscal year,” for purposes of clarity, when discussing the annual update for the LTCH PPS standard Federal payment rate, including the provisions of the Affordable Care Act, we use “fiscal year” rather than “rate year” for 2011 and subsequent years.

b. Proposed Annual Update to the LTCH PPS Standard Federal Payment Rate for FY 2018

Section 1886(m)(3)(A) of the Act, provides that beginning in FY 2010, any annual update to the LTCH PPS standard Federal payment rate is reduced by the adjustments specified in clauses (i) and (ii) of subparagraph (A). Clause (i) of section 1886(m)(3)(A) provides for a reduction, for FY 2012 and each subsequent rate year, by the productivity adjustment described in section 1886(b)(3)(B)(xi)(II) of the Act (that is, “the multifactor productivity (MFP) adjustment”). Clause (ii) of section 1886(m)(3)(A) provides for a reduction, for each of FYs 2010 through 2019, by the “other adjustment” described in section 1886(m)(4)(F) of the Act.

Section 411(e) of the Medicare Access and CHIP Reauthorization Act (MACRA) (Pub. L. 114–10), enacted on April 16, 2015, amended section 1886(m)(3) of...
the Act by amending subparagraph (A) to be ‘subject to subparagraph (C)’ and by adding new subparagraph (C), which specifies an additional special rule for FY 2018. Specifically, section 1886(m)(3)(C) of the Act states for FY 2018, the annual update under subparagraph (A) for the fiscal year, after application of clauses (i) and (ii) of subparagraph (A), shall be 1 percent.

That is, the annual update for FY 2018, after applications of the reductions for the MFP adjustment (under clause (i) of section 1886(m)(3)(A)) and the “other adjustment” (under clause (ii) of section 1886(m)(3)(A)) is 1 percent.

Historically, CMS has used an estimated market basket increase to update the LTCH PPS. Under the authority of section 123 of the BBRA as amended by section 307(b) of the Bipartisan Budget Act of 2013, we adopted a newly created 2013-based LTCH-specific market basket for use under the LTCH PPS beginning in FY 2017. The 2013-based LTCH-specific market basket is based solely on the Medicare cost report data submitted by LTCHs and, therefore, specifically reflects the cost structures of only LTCHs. For additional details on the development of the 2013-based LTCH-specific market basket, we refer readers to the FY 2017 IPPS/LTCH PPS final rule (81 FR 57101 through 57102).

For FYs 2010 through 2017, the estimated market basket update under the LTCH PPS was reduced by the MFP adjustment and “other adjustment” as applicable. However, as described above, section 411(e) of the MACRA subsequently amended section 1886(m)(3)(A) of the Act so that, after the adjustments above, the FY 2018 annual update is set at 1 percent.

c. Proposed Adjustment to the LTCH PPS Standard Federal Payment Rate Under the Long-Term Care Hospital Quality Reporting Program (LTCH QRP)

In accordance with section 1886(m)(5) of the Act, as added by section 3004(a) of the Affordable Care Act, the Secretary established the Long-Term Care Hospital Quality Reporting Program (LTCH QRP). Failure to report quality data under the LTCH QRP for FY 2014 and subsequent fiscal years results in a 2.0 percentage point reduction in the annual update as codified under §412.523(c)(4) of the regulations. As previously noted, although the language of section 3004(a) of the Affordable Care Act refers to years 2011 and thereafter under the LTCH QPS as “rate year,” consistent with our change in the terminology used under the LTCH PPS from “fiscal year” to “rate year,” for purposes of clarity, when discussing the annual update for the LTCH PPS, including the provisions of the Affordable Care Act, we use “fiscal year” rather than “rate year” for 2011 and subsequent years.

The LTCH QRP, as required for FY 2014 and subsequent fiscal years by section 1886(m)(5)(A)(i) of the Act, applies a 2.0 percentage point reduction to any update under §412.523(c)(3) for an LTCH that does not submit quality reporting data to the Secretary in accordance with section 1886(m)(3)(C) of the Act with respect to such a year (that is, in the form and manner and at the time specified by the Secretary under the LTCH QRP) (§412.523(c)(4)(i)). Section 1886(m)(5)(A)(ii) of the Act provides that the application of the 2.0 percentage points reduction may result in an annual update that is less than 0.0 for a year, and may result in LTCH PPS payment rates for a year being less than such LTCH PPS payment rates for the preceding year (§412.523(c)(4)(ii)).

Furthermore, section 1886(m)(5)(B) of the Act specifies that the 2.0 percentage points reduction is applied in a noncumulative manner, such that any reduction made under section 1886(m)(5)(A)(ii) of the Act shall apply only with respect to the year involved, and shall not be taken into account in computing the LTCH PPS payment amount for a subsequent year (§412.523(c)(4)(ii)).

We discuss the application of the 2.0 percentage point reduction under §412.523(c)(4)(i) in our discussion of the proposed annual update to the LTCH PPS standard Federal payment rate for FY 2018 in section VIII.C.2.c. of the preamble of this proposed rule. (For additional information on the history of the LTCH QRP, including the statutory authority and the selected measures, we refer readers to section VIII.C.2.c. of the preamble of this proposed rule.)

d. Proposed Annual Update Under the LTCH PPS for FY 2018

Consistent with the amendments to section 1886(m)(3)(C) of the Act provided by section 411 of the MACRA, we are proposing an update to the LTCH PPS standard Federal payment rate of 1 percent for FY 2018.

For FY 2018, section 1886(m)(5) of the Act requires that, for LTCHs that do not submit quality reporting data as required under the LTCH QRP, any annual update to an LTCH PPS standard Federal payment rate, after application of the adjustments required by section 1886(m)(3) of the Act, shall be further reduced by 2.0 percentage points. For LTCHs that fail to submit quality reporting data under the LTCH QRP under §412.523(c)(3)(xiv) in conjunction with §412.523(c)(4), we are proposing to further reduce the proposed annual update to the LTCH PPS standard Federal payment rate by 2.0 percentage points in accordance with section 1886(m)(5) of the Act.

As such, the proposed update to the LTCH PPS standard Federal payment rate for FY 2018 for LTCHs that fail to submit quality reporting data under the LTCH QRP will be the proposed 1-percent annual rate increase for FY 2018 reduced by 2.0 percentage points. For this proposed rule, we are proposing to establish a proposed annual update to the LTCH PPS standard Federal payment rate of ~1 percent (that is, 1 percent minus 2.0 percentage points) for FY 2018 for LTCHs that fail to submit quality reporting data as required under the LTCH QRP. As provided in §412.523(c)(4)(iii) and as noted above, the application of the 2.0 percentage points reduction may result in an annual update that is less than 0.0 for a year, and may result in LTCH PPS payment rates for a year being less than such LTCH PPS payment rates for the preceding year. (We note that, consistent with historical practice, in determining the proposed FY 2018 LTCH PPS standard Federal payment rate, we are also applying a proposed area wage level budget neutrality factor in accordance with §412.523(d)(4) (as discussed in section V.B. of the Addendum to this proposed rule) and a proposed budget neutrality adjustment stemming from our proposed change to the SSO payment methodology (as discussed in VIII.D. of the preamble of this proposed rule).

Absent the special provisions for FY 2018 required by section 1886(m)(3)(C) of the Act, we note the proposed annual market basket update would have been based on the FY 2018 full market basket increase of 2.8 percent (based on IGI’s fourth quarter 2016 forecast of the 2013-based LTCH market basket) reduced by the proposed FY 2018 MFP adjustment of 0.4 percentage point (also based on IGI’s fourth quarter 2016 forecast).

Following application of the productivity adjustment, the adjusted proposed market basket update of 2.4 percent (2.8 percent minus 0.4 percentage point) would have then been further reduced by 0.75 percentage point, as required by sections 1886(m)(3)(A)(ii) and 1886(m)(4)(F) of the Act. This would have resulted in a proposed annual market basket update under to the LTCH PPS standard Federal payment rate for FY 2018 of 1.65 percent (that is, 2.8 percent, less the proposed MFP adjustment of 0.4 percentage point, and less the 0.75 percentage point required under section
In the FY 2003 LTCH PPS final rule (67 FR 55954) that implemented the LTCH PPS, under §412.529, we established a special payment policy for short-stay outlier (SSO) cases; that is, cases with a covered length of stay that is less than or equal to five-sixths of the geometric average length of stay for each LTC–DRG. When we established the SSO policy, we explained that a short-stay outlier case may occur when a beneficiary receives less than the full course of treatment at the LTCH before being discharged (67 FR 55995). Also, in the FY 2003 LTCH PPS final rule, we stated that we had first described the policy in the proposed rule, we based the proposed policy on the belief that many of these patients could have been treated more appropriately in an acute care hospital subject to the acute care hospital inpatient prospective payment system (67 FR 55995). Therefore, under the LTCH PPS, we implemented a special payment adjustment for SSO cases. Under the original SSO policy, for LTCH PPS discharges with a covered length of stay of greater than 25 days. In the FY 2003 LTCH PPS final rule, we implemented a special payment adjustment for SSO cases. Under the original SSO policy, for LTCH PPS discharges with a covered length of stay of greater than 25 days. We believe that the frequency of discharges rises sharply immediately after the SSO threshold is met. This pattern holds true across MS–LTC–DRGs and for every category of LTCHs. We believe that these analyses strongly suggest that LTCHs’ discharge decisions are influenced at least as much by this financial incentive as by clinical considerations. Our own analysis of LTCH claims data showed similar findings. In light of these concerns, in this proposed rule, we are proposing to address this financial incentive and discourage such delay in the discharge of LTCH patients by proposing to revise our SSO policy. We note that, under the dual rate LTCH PPS payment structure, our existing SSO policy only applies to the LTCH PPS standard Federal payment rate. Accordingly, the proposed changes to our SSO policy presented in this section would only apply to LTCH PPS standard Federal payment rate cases (or, for cost reporting periods beginning before October 1, 2017, the LTCH PPS standard Federal payment rate portion of the blended rate payment under §412.522(c)(3)(ii)). Under our proposed policy, the SSO definition would remain unchanged, but the current payment adjustment options would be replaced with a single graduated per diem payment adjustment calculated using a blended payment rate that, as the length of stay increases, consists of a decreasing portion of the payment amount paid at the IPPS per diem amount (referred to as the “IPPS comparable amount”) and an increasing portion paid at 120 percent of the MS–LTC–DRG per diem payment amount (referred to as the “LTCH PPS per diem amount”), with a maximum payment amount set at the full LTCH PPS standard Federal payment rate. Specifically, beginning with discharges occurring on or after October 1, 2017, we are proposing to pay SSO cases solely on the “blended” option in the current SSO payment adjustment formula described at §412.529(c)(2)(iv); that is, a SSO case would be paid based on a blend of the IPPS comparable amount (determined under §412.529(d)(4)(i)) and the MS–LTC–DRG per diem amount (determined under §412.529(d)(1)) in conjunction with §412.503. Under this blended payment method at existing §412.529(c)(2)(iv), as the length of stay of a SSO case increases, the percentage of the per diem payment amounts based on the full MS–LTC–DRG standard Federal payment rate would increase, and the percentage of the payment based on the IPPS comparable amount would decrease. This blended per diem payment rate adjustment would result in paying LTCH cases with a very short length of stay more like an IPPS case, and LTCH cases with relatively longer lengths-of-stay more like a non-short-stay LTCH PPS standard Federal payment rate case. Therefore, as the length of stay of a LTCH PPS standard Federal payment rate case increases, the treatment resources and costs associated with the stay are more comparable to typical LTCH PPS standard Federal payment rate payments and less comparable to payments for the same stay at an acute care hospital under the IPPS. If adopted, this policy would result in payment amounts becoming more commensurate with the LTCH PPS
standard Federal payment rate as the case begins to resemble a more characteristic LTCH PPS standard Federal payment rate case. We believe that, by paying SSO cases on this basis, we would reduce, if not eliminate, the payment “cliffs” (or payment differentials) inherent in our current payment methodology, as well as the financial incentives that appear to have resulted in potentially improper delays in patient discharges other than solely for medical reasons. In addition, we believe that this proposed per diem “blended” approach would provide an appropriate balance between the 1-day marginal payment and the 1-day marginal incurred cost.

Under this proposal, we are proposing to codify the change to the SSO policy described above by revising § 412.529 of the regulation. Specifically, we are proposing to add paragraph (c)(4) to provide that, for discharges occurring on or after October 1, 2017, SSO cases will be paid according to the blended payment option at existing § 412.529(c)(2)(ii) and corresponding changes to § 412.529(c)(3) by sunsetting the previous SSO payment formula as of October 1, 2017.

The goal of this proposed revision to the SSO policy is to remove the incentive to delay patient discharges for payment reasons. In assessing the potential impact of this proposed policy change, we found two different impacts on Medicare LTCH spending: One would increase spending while the other would decrease spending.

First, we expect this proposed SSO payment adjustment methodology would result in increased payments to SSO cases. Based on data and FY 2018 payment estimates used for this proposed rule, we estimate that, under this proposal, Medicare payments to SSO cases would increase approximately 30 percent, or approximately $145 million (without taking into account any assumptions on changes to LTCHs’ discharge practices). These increased payments for SSO cases would produce a somewhat substantial increase in aggregate Medicare spending for LTCH PPS standard Federal payment rate cases (that is, an approximate 4.6-percent increase to current projected LTCH PPS standard Federal payment rate case payments).

At the same time, without the economic incentive to delay discharge until the SSO threshold is met, under our proposal, we expect LTCHs would discharge some patients sooner, even while the length of stay of the patient is still within the SSO period. Therefore, in the absence of the proposed policy, these cases would not have previously been SSO cases. We believe the proposed policy would result in some reduction in Medicare spending due to an expected decrease in Medicare payments for LTCH PPS standard Federal payment rate cases that, under the current SSO policy, were not receiving the SSO payment adjustment (because discharges were delayed until the SSO threshold was met).

However, while we expect this behavior change by LTCHs would reduce Medicare expenditures, we do not believe that the decrease in expenditures from fewer delayed discharge cases would offset the estimated increase in expenditures under the proposed SSO payment adjustment methodology. As such, we project that this proposed change to the payment formula for SSOs would result in a net increase in aggregate Medicare LTCH payments compared to aggregate Medicare payments under the current methodology.

The goal of the proposed policy is to remove the incentive to delay patient discharges for payment reasons, not to increase aggregate Medicare LTCH PPS payments. Therefore, we believe the appropriate policy approach is to propose to implement this proposed change to the SSO payment methodology on a budget neutral basis; that is, to implement the proposed SSO payment adjustment methodology by adjusting the LTCH PPS standard Federal payment rate so that our projection of aggregate FY 2018 payments for LTCH PPS standard Federal payment rate cases made under our proposed SSO payment adjustment methodology would be equal to our projection of aggregate FY 2018 payments paid for LTCH PPS standard Federal payment rate cases under our existing SSO payment adjustment methodology.

We further note that, based on most recent claims data, we believe the effects of a budget neutral approach would primarily occur within each LTCH and, therefore, result in minimal redistribution between different LTCHs. Specifically, FY 2015 claims data show that nearly all LTCHs treated at least one SSO case, and those that did not treat any SSO cases, on average, had very few LTCH PPS standard Federal payment rate cases. In addition, for over 90 percent of all LTCHs, at least 20 percent of their LTCH PPS standard Federal payment rate cases were SSO cases. Therefore, we expect that, for most LTCHs, the increase in payments for their SSO cases under this proposed change to the payment methodology would generally offset any SSO budget neutrality-related decrease in payment to their non-SSO LTCH PPS standard Federal payment rate cases.

In implementing the proposed SSO payment methodology, we are proposing to use a budget neutrality adjustment to offset the projected net increase in Medicare spending, which accounts for both the estimated decrease in Medicare payments resulting from LTCHs no longer holding patients until the SSO threshold is met and the larger estimated increase in spending to SSO cases described earlier. We believe that our proposal to incorporate a projection of the expected decrease in spending resulting from behavior change to not hold patients beyond the SSO threshold appropriately reflects the net impact of the proposed change. Further, this lessens the impact of any budget neutrality adjustment estimated without accounting for these expected behavioral changes—in other words, if the budget neutrality adjustment only adjusted for the increased payments to SSO cases.

To do so, we are proposing to amend § 412.523 by adding a new paragraph (d)(5), which would specify that the LTCH PPS standard Federal payment rate will be adjusted by a one-time, permanent factor that accounts for the projected change in estimated aggregate payments to LTCH PPS standard Federal payment rate cases in FY 2018 due to the change in the payment methodology for SSO cases described at § 412.529(c)(4). (As noted earlier, this budget neutrality adjustment would only affect the LTCH PPS standard Federal payment rate.) This factor would ensure that the proposed change to the SSO payment methodology in FY 2018 does not affect aggregate LTCH PPS payments; that is, this proposed policy change is budget neutral. Specifically, we are proposing to use the following methodology to determine the proposed budget neutrality factor that would be applied to the proposed FY 2018 LTCH PPS standard Federal payment rate using the 2016 LTCH standard Federal payment rate payment rates used for this proposed rule. These estimates are based upon the most recently available data (for example, the December 2016 update of the FY 2016 MedPAR file), and consistent with historical practice, if more recent data become available, we are proposing to use such data for the final rule.

- Step 1—Simulate estimated aggregate FY 2018 LTCH PPS standard Federal payment rate payments using the existing SSO payment methodology at § 412.529(c)(3). (For the remainder of this discussion, we refer to this amount as “estimated FY 2018 payments under the existing SSO payment
Spending that we would expect under different impacts on Medicare LTCH amount reflects the first of the two approximately $145 million. This increase approximately 4.6 percent, or under our SSO policy proposal, FY 2018 absence of any behavioral assumptions, section I.J.1. of the Regulatory Impact proposed rule (which is described in discussion, we refer to this amount as the “estimated FY 2018 payments under the proposed SSO payment methodology”).

Step 2a—Simulate estimated aggregate FY 2018 LTCH PPS standard Federal payment rate payments under the proposed SSO payment methodology without accounting for expected changes in LTCHs’ discharge behavior. (For the remainder of this discussion, we refer to this amount as the “estimated unadjusted FY 2018 payments under the proposed SSO payment methodology.”) We note that this estimate is comprised of estimated unadjusted FY 2018 payments under the proposed SSO payment methodology for non-SSO cases and for SSO cases.) This estimate represents the proposed change in the SSO payment methodology alone in the absence of any behavioral assumptions. (In other words, after applying Step 2b, under our actuarial assumptions, estimated FY 2018 unadjusted payments under the proposed SSO payment methodology for non-SSO cases are projected to be 90 percent of the corresponding estimate for such cases from Step 2a to reflect the expected decrease in non-SSO cases under the proposed changes to the SSO policy.) Based on data used for this proposed rule, we estimate that 10 percent of our estimated unadjusted FY 2018 payments under the proposed SSO payment methodology for non-SSO cases is approximately $272 million. (In Step 2d below, this estimated $272 million is subtracted from our estimated FY 2018 unadjusted payments under the proposed SSO payment methodology to account for the projected decrease in non-SSO cases under the proposed changes to the SSO policy.)

Step 2c—Determine the estimated amount of aggregate FY 2018 LTCH PPS standard Federal payment rate payments that would reflect the projected decrease in non-SSO cases under the proposed changes to the SSO policy. Under our actuarial assumptions (used in Step 2b above and described in detail below), we project SSO cases under the proposed change to the SSO policy to have a length of stay decreased to less than the SSO threshold. (The basis for the decrease in the length of stay is discussed in greater detail below.) This 10 percent of the “aggregate SSO comparable amount” represents our estimate of the aggregate increase in SSO payments under our proposed SSO policy for those cases that are expected to shift to SSO cases from non-SSO cases because we are projecting that 10 percent of non-SSO cases would become SSO cases as a result of our proposal.

Therefore, under this step, we would add an amount equal to 10 percent of the “aggregate SSO comparable amount” to the amount determined in Step 2a. (In other words, under our actuarial assumptions and after applying Step 2c, our estimated unadjusted FY 2018 payments under the proposed SSO payment methodology would be increased to reflect the expected increase in SSO cases.)

To estimate proposed SSO payments based on non-SSO cases under this step, because our proposed payment adjustment for SSO cases depends on the length of stay, these estimated payments depend on where, relative to the SSO threshold, the shifts from non-SSO cases to SSO cases occur. As we discuss in greater detail below, our actuaries estimate the majority of the increase in SSO cases resulting from this proposed policy would occur within 1 to 3 days prior to the SSO threshold. As such, we based our estimated payment amount in this step on our actuarial assumption (discussed in greater detail below) that the length of stay shifts would occur only between 1 and 3 days prior to the SSO threshold. We then performed three payment simulations to estimate proposed SSO payments if all of the non-SSO cases would have a length of stay of 1 day, 2 days, and 3 days prior to the SSO threshold. To determine the estimated SSO payments for the non-SSO cases,
we took an average of those three aggregate estimates: payments where non-SSO cases moved 1 day prior to the SSO threshold; payments where non-SSO cases moved 2 days prior to the SSO threshold; and payments where non-SSO cases moved 3 days prior to the SSO threshold. This amount is the “aggregate SSO comparable amount” described above. Then we took 10 percent of the “aggregate SSO comparable amount” as the estimated increase in aggregate SSO payments expected to result from the expected increase in SSO cases under our proposal. Based on data used for this proposed rule, using the calculation described above, we estimate that 10 percent of the “aggregate SSO comparable amount” is approximately $229 million. (In Step 2d below, this estimated $229 million is added to our estimated FY 2018 unadjusted payments under the proposed SSO payment methodology to account for the projected increase in SSO cases under the proposed changes to the SSO policy.)

Step 2d—Adjust the original estimated unadjusted FY 2018 payments under the proposed SSO payment methodology ($272 million from Step 2a) to account for the projected decrease in non-SSO cases under the proposed changes to the SSO policy (by subtracting the amount determined in Step 2b) and for the projected increase in SSO cases under the proposed changes to the SSO policy (by adding the $229 million from Step 2c). The resulting amounts is the estimated FY 2018 payments under the proposed SSO payment methodology (which is used in Step 3 below). As such, we estimate FY 2018 payments under the proposed SSO payment methodology is $3.279 billion (that is, $3.322 billion from Step 2a minus the $272 million from Step 2b plus the $229 million from Step 2c.) Therefore, we estimate that our proposed change to the SSO payment methodology would result in an increase in payments of approximately $102 million (that is, the $3.177 billion as calculated in Step 1 minus the $3.279 billion as calculated here in Step 2).

Actuarial Assumptions for Shifts in Cases Used under Steps 2b and 2c: Our actuarial assumptions for LTCHs’ discharge behavior under our proposed SSO policy were estimated based on a comparative analysis of distributions of LTCH discharges relative to the SSO thresholds in FY 2003 and FY 2015 using data from FY 2002 (the year before the LTCH PPS was implemented and the first year a SSO payment adjustment) to LTCH discharges in FY 2015 (the most recent complete year of data available at the time the comparative analysis was performed in preparation for this proposed rule). (We note that, for FY 2002, because there was no applicable SSO threshold, we used the SSO thresholds from FY 2003 (LTC–DRG Version 23) based on the billed LTC–DRG (LTC–DRG Version 22) on the FY 2002 claim.)

The FY 2002 distribution shows a nearly continuous distribution of LTCH discharges relative to what would become the SSO threshold in FY 2003, and approximate symmetry before and after the SSO threshold. In other words, for FY 2002, the distribution of discharges just after what would become the FY 2003 threshold looks similar to the distribution of discharges just before that threshold, and there is a corresponding similarity between discharges well after and well before what would become the SSO threshold.

While the FY 2015 distribution of LTCH discharges relative to the SSO threshold shows the same symmetry among discharge patterns before and well after the threshold, there are significantly fewer discharges just before the SSO threshold and significantly more discharges just after the SSO threshold (instead of a symmetry among discharges just before and just after the SSO threshold). For FY 2015, this lack of symmetry is concentrated in the 3 days leading up to the SSO threshold. (We note that, in our analysis of LTCH discharge patterns relative to the applicable SSO threshold, we found similar patterns for FYs 2003 through 2014 as those observed for FY 2015, as well as for FY 2016 LTCH discharges.)

In particular, the FY 2015 LTCH discharges have, as a proportion of total FY 2015 LTCH discharges, approximately 20 percent more discharges occurring just after the SSO threshold when compared to FY 2002 discharges. However, due to other substantial changes in Medicare payments to LTCHs, including the introduction of the LTCH PPS in FY 2003, we do not believe the entire 20-percent shift in discharges is attributable to only the introduction and subsequent revisions to the LTCH PPS SSO payment adjustment. Moreover, this shift is not uniform across all SSO discharges because the majority of shifting past the SSO threshold occurs within 3 days of the SSO threshold. Based on this, our actuaries estimate that the elimination of the payment cliff would result in a 10-percent reduction in non-SSO cases, resulting in an increase payments of approximately $102 million (that is, the $3.177 billion
as calculated in Step 1 minus the $3,279 billion as calculated in Step 2) which reflects the approximate $43 million decrease that accounts for our actuarial assumptions for expected changes in LTCIs’ discharge behavior under the proposed changes to the SSO policy.

For this proposed rule, using the steps in the proposed methodology described earlier, we have determined a proposed budget neutrality factor for the proposed change to the SSO payment methodology of 0.9672. (We are proposing, consistent with historical practice, that if more recent data become available and if finalized, we would use such data to determine a budget neutrality factor for the proposed change to the SSO payment methodology in the final rule.) Accordingly, in section V.A. of the Addendum to this proposed rule, to determine the proposed FY 2018 LTCH PPS standard Federal payment rate, we are proposing to apply a one-time, permanent budget neutrality factor of 0.9672 for the proposed change in the SSO payment methodology. The proposed FY 2018 LTCH PPS standard Federal payment rate shown in Table 1E in section VI. of the Addendum to this proposed rule reflects this proposed adjustment.

E. Temporary Exception to the Site Neutral Payment Rate for Certain Spinal Cord Specialty Hospitals

Section 15009 of Public Law 114–255 added new subparagraph (F) to section 1886(m)(6) of the Act, which provides for a temporary exception to the site neutral payment rate for certain spinal cord specialty hospitals. Under this provision, discharges occurring in cost reporting periods beginning during FY 2018 and FY 2019 for LTCHs that meet the specified statutory criteria are excepted from the site neutral payment rate (that is, all discharges from such LTCHs during this period would be paid at the LTCH PPS standard Federal payment rate). Clauses (i) through (iii) of section 1886(m)(6)(F) of the Act state that, in order for an LTCH to qualify for this temporary exception, the LTCH must: (1) Have been a not-for-profit LTCH on June 1, 2014, as determined by cost report data; (2) of the discharges in the calendar year 2013 from the LTCH for which payment was made under the LTCH PPS, at least 50 percent were classified under MS–LTC–DRGs 28, 29, 52, 57, 551, 573, and 963; and (3) have discharged inpatients (including both individuals entitled to, or enrolled for, Medicare Part A benefits and individuals entitled or enrolled) during FY 2014 who had been admitted from at least 20 of the 50 States, determined by the States of residency of such inpatients and based on such data submitted by the hospital to the Secretary as the Secretary may require. The statute further provides authority for the Secretary to implement the third criterion (set forth at section 1886(m)(6)(F)(iii) of the Act and referred to as the “significant out-of-state admissions criterion”) by program instruction or otherwise, and exempts the policy initiatives from any information collection requirements under the Paperwork Reduction Act (Chapter 35 of Title 44 of the United States Code). Given this express authority, we plan to provide further details regarding the implementation of the significant out-of-state admissions criterion through subregulatory guidance. However, in this proposed rule, we are proposing to codify the requirements of the temporary exception to the site neutral payment rate for certain spinal cord specialty hospitals specified under section 1886(m)(6)(F) of the Act, as added by section 15009 of Public Law 114–255. Specifically, we are proposing to codify the requirements of this provision at new § 412.522(b)(4), by providing for an exception from the site neutral payment rate for discharges occurring in cost reporting periods beginning during FYs 2018 and 2019 for LTCHs that meet the specified statutory criteria. We are seeking public comments on this proposal. Based on information currently available, we believe that two hospitals may qualify for this exception.

F. Temporary Exception to the Site Neutral Payment Rate for Certain Discharges With Severe Wounds From Certain LTCHs

Section 15010 of Public Law 114–255 added a new subparagraph (G) to section 1886(m)(6) of the Act, which creates a temporary exception to the site neutral payment rate for certain severe wound discharges from certain LTCHs during such LTCHs’ cost reporting periods beginning during FY 2018 (that is, for cost reporting periods beginning on or after October 1, 2017 and on or before September 30, 2018). Under the provisions of section 15010 of Public Law 114–255, in order for an LTCH’s discharge to be excluded from the site neutral payment rate under this section during its FY 2018 cost reporting period, the discharge must be: (1) From an LTCH “identified by the last sentence of subsection (d)(1)(B)” of the Act; (2) classified under MS–LTC–DRG 602, 603, 530, or 540; and (3) with respect to an individual treated by an LTCH, for a severe wound. The statute defines a “severe wound,” for the purposes of the exception, as “a wound which is a stage 3 wound, stage 4 wound, unstageable wound, non-healing surgical wound, or fistula as identified in the claim from the long-term care hospital.” The statute further defines a “wound” as “an injury involving division of tissue or rupture of the integument or mucous membrane with exposure to the external environment.”

Much of this language is identical or substantially similar to the language for the previous temporary exception for discharges for the treatment of severe wounds provided for under the amendments made by section 231 of the Consolidated Appropriations Act, 2016 (Pub. L. 114–113), except for three key differences. First, the previous temporary exception for severe wound discharges applied to LTCHs that are grandfathered hospitals-within-hospitals (HwHs) (that is, hospitals that are described under § 412.23(e)(2)(i) that meet the criteria of § 412.22(f)) and are located in a rural area or treated as rural (§ 412.522(b)(2)(ii)); under the new temporary exception for severe wound discharges only requires that LTCHs are grandfathered HwHs (and does not require the LTCH to also be located in a rural area or treated as rural). Second, under this new temporary exception for severe wound discharges, the definition of a “severe wound” includes only five of the eight categories (stage 3 wound, stage 4 wound, unstageable wound, non-healing surgical wound, and fistula) included in the definition of a “severe wound” under the original temporary exception for severe wound discharges (and does not include the categories of infected wound, osteomyelitis, and wound with morbid obesity). Finally, this new temporary exception for severe wound discharges is limited to discharges that meet the definition of a severe wound and are grouped to certain specified MS–LTC–DRGs, while the previous temporary exception for severe wound discharges only required the discharge to meet the definition of a severe wound (and did not include the requirement for the discharge to also be grouped to certain specified MS–LTC–DRGs). Additional details of the new temporary exception for payment for severe wound discharges provided by Public Law 114–255, including further discussion of the likenesses to and differences from the original temporary exception for payment for severe wound discharges provided by Public Law 114–113 are discussed below.

We implemented the original temporary exception for payment for discharges for the treatment of severe wounds that was provided by the
amendments made by section 231 of Public Law 114–113 in an interim final rule with comment period (IFC) that appeared in the Federal Register on April 21, 2016 (81 FR 23428 through 23438) (referred to as the “April 21, 2016 IFC”) and finalized concurrently in the FY 2017 IPPS/LTC PPS final rule (81 FR 57070). Therefore, to the extent applicable, we are implementing this provision in an identical manner to our implementation of the amendments made by section 231 of the Consolidated Appropriations Act, which is codified in the LTCH PPS regulations at § 412.522(b)(2). Specifically, § 412.522(b)(2)(ii)(B)(1) refers to LTCHs “identified by the last sentence of subsection (d)(1)(B)” of the Act as LTCHs “[d]escribed in § 412.23(e)(2)(i) and meets the criteria of § 412.22(f).”

We are proposing to codify the requirements of this “new” temporary exception for severe wounds at new § 412.522(b)(3), by providing for an exception for discharges meeting the statutory criteria that occur in a cost reporting period that begins during FY 2018 for LTCHs described in § 412.23(e)(2)(i) and meets the criteria of § 412.22(f).

Clauses (ii) and (iii) of section 1886(m)(6)(G) of the Act, respectively, as added by section 15010 of Public Law 114–255, includes definitions of “severe wound” and “wound” for purposes of this “new” temporary exception for discharges for the treatment of severe wounds. We are proposing to incorporate the definitions of “wound” and “severe wound” at § 412.522(b)(3)(ii) as they are defined in the statute. We note that the definition of a “wound” in section 15010 is nearly identical to CMS’ definition of “wound” at existing § 412.522(b)(2)(i). We further note that the definition of a “severe wound” is nearly identical to the definition used in section 231 of Public Law 114–113 with the exception that three categories included in the latter (that is, infected wound, osteomyelitis, and wound with morbid obesity) are not included in the definition set forth in section 15010 of Public Law 114–255. The five remaining categories of stage 3 wound, stage 4 wound, unstageable wound, non-healing surgical wound, and fistula are identified by the list of ICD–10–CM codes posted to the CMS Web site at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/LongTermCareHospitalPPS/download.html under the “Severe Wound Diagnosis Codes by Category for Implementation of Section 231 of Public Law 114–113” download file. For more information on our interpretation of these terms, we refer readers to the April 21, 2016 IFC (81 FR 23428 through 23438) and the FY 2017 IPPS/LTC PPS final rule (81 FR 57070).

Therefore, this information on how CMS interpreted the meanings of these categories of a “severe wound” for Public Law 114–113 was available at the time Public Law 114–255 was enacted. As such, we are implementing the “new” temporary exception for discharges for the treatment of severe wounds provided for by section 15010 using the same list of ICD–10–CM codes to identify the five categories of severe wounds enumerated in that section of Public Law 114–255. In addition, as provided by section 1886(m)(6)(G)(i)(III) of the Act as added by section 15010 of Public Law 114–255, we are proposing at new § 412.522(b)(3)(ii) that the patient must be treated for a severe wound that meets the statutory definition of a “severe wound” at proposed § 412.522(b)(3)(i) in order for the LTCH discharge to meet this “new” temporary exception for discharges for the treatment of severe wounds.

We believe that the requirement under the “new” temporary exception for discharges for the treatment of severe wounds set forth under section 1886(m)(6)(G)(i)(II) of the Act as added by section 15010 of Public Law 114–255 for an LTCH discharge be classified under MS–LTCH–DRG 602, 603, 539, or 540 is self-implementing. Accordingly, we are proposing to codify this requirement at new § 412.522(b)(3)(ii)(C) by listing the applicable MS–LTCH–DRGs.

Section 1886(m)(6)(G)(i)(I) of the Act, as added by section 15010 of Public Law 114–255, specifies that, for purposes of this “new” temporary exception for discharges for the treatment of severe wounds, the LTCH discharge must be from an LTCH “identified by the last sentence of subsection (d)(1)(B)”. The phrase “identified by the last sentence of subsection (d)(1)(B) [of the Act]” is equivalent to the phrase “identified by the amendment made by section 4417(a) of the Balanced Budget Act of 1997” used in section 231 of Public Law 114–113, because the amendment made by section 4417(a) of the Balanced Budget Act of 1997 added the last sentence of subsection (d)(1)(B) to the Act. As discussed in the April 21, 2016 IFC (81 FR 23428), the phrase “identified by the amendment made by section 4417(a) of the Balanced Budget Act of 1997” (which as previously discussed is equivalent to “identified by the last sentence of subsection (d)(1)(B) of the Act”) has been interpreted by CMS to mean hospitals-within-hospitals (HwHs) that were participating in Medicare, but excluded from the hospital IPPS on or before September 30, 1995 (that is, hospitals which are described under § 412.23(e)(2)(i) that meet the criteria of § 412.22(f) (81 FR 23430 through 23432).

As further discussed in the April 21, 2016 IFC, § 412.22(f) generally requires that, in order to have grandfathered status, an HwH must continue to operate under the same terms and conditions, including, but not limited to, the number of beds. A limited exception to this general policy allowed eligible hospitals to increase the number of beds between October 1, 1995, and September 30, 2003, without loss of their grandfathered status. A second exception allows grandfathered HwHs to increase square footage or decrease the number of beds for cost reporting periods beginning on or after October 1, 2006, while still retaining grandfathered status. Because this phrase had already been interpreted in this manner, the April 21, 2016 IFC adopted the same meaning of the phrase for purposes of implementing section 231 of Public Law 114–113. For additional information on hospitals “identified by the amendment made by section 4417(a) of the Balanced Budget Act of 1997,” we refer readers to the April 21, 2016 IFC (81 FR 23431 through 23432). Therefore, for the purposes of the new temporary exception for LTCH discharges for the treatment of severe wounds, “identified by the last sentence of subsection (d)(1)(B) of the Act” means HwHs that were participating in Medicare, but excluded from the hospital IPPS on or before September 30, 1995 (that is, hospitals which are described under § 412.23(e)(2)(i) that meet the criteria of § 412.22(f). We finalized this policy without modification in the FY 2017 IPPS/LTC PPS final rule (81 FR 57069). Because we have already finalized our interpretation of this phrase, we believe that the requirement at section 1886(m)(6)(G)(i)(I) of the Act is self-implementing. Accordingly, we are proposing to codify this requirement at new § 412.522(b)(3)(ii)(B). LTCHs that believe they meet the requirements to be a grandfathered HwH must contact their MACs. MACs will verify that the LTCH meets these requirements.

G. Moratorium and Proposed Regulatory Delay of the Full Implementation of the “25-Percent Threshold Policy” Adjustment (§ 412.538)

The “25-percent threshold policy” is a per discharge payment adjustment in the LTCH PPS that is applied to payments for Medicare patient discharges from an LTCH when the number of such patients originating from any single referring hospital is in
excess of the applicable threshold for a given cost reporting period (such threshold is generally set at 25 percent, with exceptions for rural and urban single or MSA-dominant hospitals). If an LTCH exceeds the applicable threshold during a cost reporting period, payment for the discharge that puts the LTCH over its threshold and all discharges subsequent to that discharge in the cost reporting period from the referring hospital are adjusted at cost report settlement (discharges not in excess of the threshold are unaffected by the 25-percent threshold policy). The 25-percent threshold policy was originally established in the FY 2005 IPPS final rule for LTCH HwHs and satellites (69 FR 49191 through 49214). We later expanded the 25-percent threshold policy in the FY 2008 LTCH PPS final rule to include all LTCHs and LTCH satellite facilities (72 FR 26919 through 26944). Several laws have mandated delayed implementation of the policy, including, most recently, section 1206 of the Pathway for Sustainable Growth Rate (SGR) Reform Act (Pub. L. 113–67). Section 1206(b)(1)(B) provides a permanent exemption from the application of the 25-percent threshold policy for LTCHs identified by the amendment made by section 4417(a) of the Balanced Budget Act of 1997 (Pub. L. 105–33). As explained more fully in section VIII.H. of the preamble of this proposed rule, LTCHs “identified by the amendment made by section 4417(a) of the Balanced Budget Act of 1997” are HwHs that were participating in Medicare, but excluded from the hospital IPPS on or before September 30, 1995 (that is, hospitals which are described under §412.23(e)(2)(i)) that meet the criteria of §412.22(f). LTCHs that believe they meet the requirements to be a grandfathered HwH should contact their MACs. MACs will verify that the LTCH meets these requirements. Section 1206(b)(1)(A) of Public Law 113–67 extended prior moratoria on the full implementation of the 25-percent threshold policy until cost reporting periods beginning on or after either July 1, 2016 (for LTCHs subject to 42 CFR 412.534) or October 1, 2016 (for LTCHs subject to 42 CFR 412.536). For more details on the various laws that delayed the full implementation of the 25-percent threshold policy, we refer readers to the FY 2015 IPPS/LTCH PPS final rule (79 FR 50356 through 50357). In the FY 2017 IPPS/LTCH PPS final rule, we consolidated the 25-percent threshold provisions under new sections §412.534 and §412.536 and establishing provisions under new section §412.538.

Section 15006 of Public Law 114–255 further amended section 114(c)(1)(A) of the MMASEA (as amended) by striking “for a 9-year period” and inserting “through June 30, 2016, and for discharges occurring on or after October 1, 2016 and before October 1, 2017”, which provides for an extension of the moratorium on the full implementation of the 25-percent threshold policy. In addition, section 15006(b) of Public Law 114–255 further amended section 114(c)(2) of the MMASEA (as amended) by inserting “or any similar provision,” after “Regulations,” in subparagraphs (A) and (B). (We note that the functional result of the extension of the moratorium under section 15006(a) of Public Law 114–255 only extends to discharges on or after October 1, 2016 and before October 1, 2017.)

To implement the provisions of section 15006 of Public Law 114–255, we are proposing to make conforming amendments to the regulations that currently govern the application of the 25-percent threshold policy. Section 114(c)(1) of the MMASEA, from its inception, precluded CMS from implementing either §§412.534 or 412.536 (as applicable), as well as any similar provision to hospitals described in the provision of the MMASEA. Section 15006 of Public Law 114–255 amended section 114(c)(2) of the MMASEA by adding the words “or any similar provisions” to both (A) and (B). Section 412.538 of the regulations is a similar provision to the provisions of both §§412.534 and 412.536 (we adopted the payment policy under §412.538 to create a consolidated and streamlined 25-percent threshold policy to replace the policies under §§412.534 and 412.536, which were sunset). Therefore, in order to implement the moratorium on the implementation of the 25-percent threshold policy provided under section 15006 of Public Law 114–255, we are proposing to amend §412.538 to account for these statutory changes. We note that, similar to the July 1, 2012 through September 30, 2012 “gap” period discussed in the FY 2013 IPPS/LTCH PPS final rule (77 FR 53484 through 53486), this extension of the moratorium on the full application of the 25-percent threshold policy results in a “gap” period where LTCHs are required to comply with the fully-implemented 25-percent threshold policy for their cost reporting periods beginning on or after July 1, 2016, and before October 1, 2016, for any discharges occurring on or before September 30, 2016. For the same reasons discussed in the FY 2013 IPPS/LTCH PPS final rule (77 FR 53485 through 53486), although those LTCHs with cost reporting periods beginning on or after July 1 and before October 1, 2016 are “technically” subject to the 25-percent threshold policy until October 1, 2016, we believe that very few, if any, LTCHs will actually receive a payment adjustment because these LTCHs would rarely, if ever, admit more than 25 percent of their discharges from any one referring hospital during the limited period of 1 to 3 months (depending on the LTCH’s cost reporting beginning date) that the 25-percent threshold policy was technically in effect.

In addition, we are proposing to adopt a 1-year regulatory moratorium on the implementation of the 25-percent threshold policy; that is, we are proposing to impose a regulatory moratorium on our implementation of §412.538 until October 1, 2018. This proposal is made in response to the further statutory delays and our continued consideration of public comments received in response to our proposal to consolidate and streamline the 25-percent threshold policy in the FY 2017 IPPS/LTCH PPS proposed rule. In response to that proposed rule, several commenters stated that the new site neutral payment rate would alleviate the policy concerns underlying the 25-percent threshold policy. As we stated in more detail in our response to those comments in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57106), we are not convinced that this is the case.

However, given this additional statutory moratorium, we believe that it is appropriate at this time to propose to establish a regulatory moratorium on the implementation of the 25-percent threshold policy until we can examine data under the application of the site neutral payment rate to further evaluate, when more data are available, whether the policy is in fact still necessary. While we are not convinced that the application of the site neutral payment rate removes the need for the 25-percent threshold policy, we believe that evaluating the impact of the application of the site neutral payment rate on LTCH admission practices would be premature at this time. The statute provides that the site neutral payment rate be phased-in, effective with LTCH cost reporting periods beginning on or after October 1, 2015 and before October 1, 2017 (that is, LTCH cost reporting periods beginning in FYs 2016 and 2017). LTCH claims data for discharges that occurred in FY 2016 is currently the best available data, and given that phase-in of the site neutral payment rate is based on LTCHs’ cost reporting period start dates, many LTCH discharges that occurred during FY 2016
were not yet subject to the site neutral payment rate because they occurred in a LTCH cost reporting period that had begun prior to October 1, 2016. Consequently, at this time we only have a partial year of LTCH claims data under the period where the site neutral payment rate was in effect, which may not be fully reflective of any changes in LTCH admission practices under the new dual rate LTCH PPS. Proposing an additional regulatory moratorium on the 25-percent threshold policy through FY 2018 would allow CMS the opportunity to do an analysis of LTCH admission practices under the new dual payment rate LTCH PPS based on more complete data and would avoid creating any additional confusion by having the 25-percent threshold policy become effective for a period of time when future analysis of LTCH claims data may indicate the policy concerns underlying the 25-percent threshold policy have been moderated.

Therefore, in this proposed rule, we are proposing to revise the effective date of § 412.538 so that the 25-percent threshold policy would apply to discharges occurring on or after October 1, 2018. Further, we are proposing that if, in response to public comments, we do not finalize this proposed additional 1-year regulatory moratorium, we would revise § 412.538 so that the 25-percent threshold policy would apply to discharges occurring on or after October 1, 2017, consistent with the provisions of section 15006 of Public Law 114–255. We are seeking public comments on our proposals.

H. Revision to Moratorium on Increasing Beds in Existing LTCH or LTCH Satellite Locations Under the 21st Century Cures Act (Pub. L. 114–255) (§ 412.23)

Section 1206(b)(2) of Public Law 113–67, as amended by section 112(b) of the Protecting Access to Medicare Act of 2014 (PAMA) (Pub. L. 113–93), established “new” statutory moratoria on the establishment of new LTCHs and LTCH satellite facilities and on the increase in the number of hospital beds in existing LTCHs and LTCH satellite facilities, effective April 1, 2014 through September 30, 2017, by amending section 114(d)(1)(I) of the MMSEA (as amended). In addition, the statute also provided an exception under the “new” moratorium under section 114(d)(7) of the MMSEA (as amended) to establish a new LTCH or LTCH satellite facility during the period between April 1, 2014, and September 30, 2017, if a hospital or entity meets criteria, which mirror the expired provisions of section 114(d)(2)(A). For a discussion on our implementation of these moratoria, we refer readers to the FY 2015 IPPS/LTCH PPS final rule (79 FR 50189 through 50193).

Section 15004(a) of Public Law 114–255 further amended section 114(d)(7) of the MMSEA (as amended) by striking “The moratorium under paragraph (1)(A)” and inserting “[a]ny moratorium under paragraph (1)” and specified that such amendment shall take effect as if included in the enactment of section 112 of the PAMA. Under this amendment, all existing LTCHs or LTCH satellite locations are no longer subject to a moratorium on an increase in LTCH beds set forth in paragraph (1)(B) if they meet certain criteria. In order to implement this statutory change, we are proposing to amend § 412.23(e)(7) by revising paragraph (e)(7)(iii) to specify that the moratorium on increasing the number of beds in existing LTCHs and existing LTCH satellites does not apply if one or more or the exceptions described in § 412.23(e)(6)(ii) is met in accordance with the provisions of section 1500(a) of Public Law 114–255. (We note that section 15004(b) of Public Law 114–255 provides for a modification to LTCH high-cost outlier payments. Our proposals to implement this provision are discussed in section V.D. of the Addendum to this proposed rule.) We are seeking public comments on this proposal.

I. Proposed Change to the Average Length of Stay Criterion Under the 21st Century Cures Act (Pub. L. 114–255)

Under the requirements at sections 1886(d)(1)(B)(i)(I) and 1861(ccc) of the Act, in order for a hospital to be classified as an LTCH, the hospital had to maintain an average length of stay of greater than 25 days as calculated by the Secretary. Section 1206(a)(3) of the Pathway for SGR Reform Act of 2013 (Pub. L. 113–67) excluded Medicare Advantage plans’ and site neutral payment rate discharges from this calculation for hospitals that were classified as LTCHs as of December 10, 2013. We implemented this provision in the FY 2016 IPPS/LTCH PPS final rule (80 FR 49638). Section 15007 of Public Law 114–255 amended section 1206(a)(3) of the Pathway for SGR Reform Act by extending the exclusion of Medicare Advantage plans’ and site neutral payment rate discharges from the calculation of the average length of stay to all LTCHs, for discharges occurring in cost reporting periods beginning on or after October 1, 2015. In order to implement this provision, we are proposing to remove the final sentence of our regulations at 42 CFR 412.23(e)(2)(vii), which included site neutral payment rate and Medicare Advantage discharges in the calculation of the average length of stay for LTCHs which were classified as such after December 10, 2013. We are seeking public comments on our proposal.

J. Change in Medicare Classification for Certain Hospitals (§ 412.23)

When enacted, section 1886(d)(1)(B)(iv) of the Act established a category of hospitals that experience extended average inpatient length of stays, which are known as LTCHs under the Medicare program. Clause (iv) of section 1886(d)(1)(B) consisted of two subclauses (I) and (II) that is, section 1886(d)(1)(B)(iv)(I) and section 1886(d)(1)(B)(iv)(II) of the Act which corresponded to two categories of hospitals that were generally referred to as “subclause (I)” and “subclause (II)” LTCHs. “Subclause (I)” LTCHs were required to have an average inpatient length of stay that is greater than 25 days. “Subclause (II)” LTCHs were only required to have an average inpatient length of stay of greater than 20 days. The “subclause (II)” LTCH definition further limited the classification of a “subclause (II)” LTCH by including the requirement that the LTCH must have been first excluded from the IPPS in CY 1986, and treated a Medicare inpatient population in which 80 percent of the discharges in the 12-month reporting period ending in Federal FY 1997 had a principal diagnosis that reflected a finding of neoplastic disease as defined in subsection (I)(i)(iv) section 1886 of the Act. This statutory requirement was implemented under 42 CFR 412.23(e)(2)(ii).

As part of our FY 2015 IPPS/LTCH PPS rulemaking cycle, under the authority provided by section 1206(d)(2) of the Pathway to SGR Reform Act (Pub. L. 113–67), we adopted an adjustment to the LTCH PPS payment for LTCHs classified under section 1886(d)(1)(B)(iv)(II) of the Act (“subclause (II)” LTCHs). Under this payment adjustment, “subclause (II)” LTCHs receive payment under the LTCH PPS that is generally equivalent to an amount determined under the reasonable cost-based payment rules for both operating and capital-related costs under 42 CFR part 413 (that is, an amount generally equivalent to an amount determined under the TEFRA payment system methodology). This payment adjustment for “subclause (II)” LTCHs is specified at § 412.526. For more information on this payment adjustment, we refer readers to the FY 2016 IPPS/LTCH PPS final rule (79 FR 50193 through 50197). As initially adopted, the “TEFRA-like” reasonable
Section 15008 of Public Law 114–225 provides for a change in Medicare classification for “subclause (II)” LTCH by redesignating such hospitals from section 1886(d)(1)(B)(iv)(II) to section 1886(d)(1)(B)(vi) of the Act. In addition, subsection (b) of section 15008 specifies that, for cost reporting periods beginning on or after January 1, 2015, such hospitals classified under section 1886(d)(1)(B)(vi) of the Act are not subject to section 1886(m) of the Act, which sets forth the LTCH PPS. Section 15008 further specifies that, for cost reporting periods beginning on or after January 1, 2015, payment for inpatient operating costs is to be made as described in 42 CFR 412.526(c)(3), including any subsequent modifications, and payment for capital costs is to be made as described in 42 CFR 412.526(c)(4) as in effect on January 1, 2015. (We note that there have been no revisions to the regulations at 42 CFR 512.526, including § 412.526(c)(3) and § 412.526(c)(4), since January 1, 2015.)

In order to implement these requirements, we are proposing to revise § 412.23(e)(2)(ii). We are seeking public comments on our proposal.

IX. Quality Data Reporting Requirements for Specific Providers and Suppliers

We seek to promote higher quality and more efficient healthcare for Medicare beneficiaries. This effort is supported by the adoption of widely agreed-upon quality measures. We have worked with stakeholders to define quality measures for most settings and to measure various aspects of care for most Medicare beneficiaries. These measures assess structural aspects of care, clinical processes, care coordination, and improving patient outcomes (including patient experiences with care).

We have implemented quality reporting programs for multiple care settings, including, for example:

• Hospital inpatient services under the Hospital Inpatient Quality Reporting (IQR) Program (formerly referred to as the Reporting Hospital Quality Data for Annual Payment Update (RHQDAPU) Program);

• Prospective Payment System (PPS)-exempt cancer hospitals under the PPS-Exempt Cancer Hospital Quality Reporting (PCHQR) Program;

• Long-term care hospitals under the Long-Term Care Hospital Quality Reporting Program (LTCH QRP) (also referred to as the LTCHQQR Program);

• Inpatient psychiatric facilities under the Inpatient Psychiatric Facilities Quality Reporting (IPFQR) Program;

• Hospital outpatient services under the Hospital Outpatient Quality Reporting (OQR) Program (formerly referred to as the Hospital Outpatient Quality Data Reporting Program (HOP QDRP));

• Ambulatory surgical centers under the Ambulatory Surgical Center Quality Reporting (ASCQR) Program;

• Inpatient rehabilitation facilities under the Inpatient Rehabilitation Facility Quality Reporting Program (IRF QRP);

• Care furnished by physicians and other eligible professionals under the Physician Quality Reporting System (PQRS). We note that beginning in CY 2018 PQRS will be replaced by the Quality Payment Program (QPP);

• Skilled nursing facilities under the Skilled Nursing Facility Quality Reporting Program (SNF QRP);

• Home health agencies under the Home Health Quality Reporting Program (HH QRP); and

• Hospices under the Hospice Quality Reporting Program (HQRP).

We have also implemented programs which link payment to performance including: The Hospital Readmissions Reduction Program; the Hospital Value-Based Purchasing (VBP) Program (described further below); the Hospital-Acquired Condition (HAC) Reduction Program; the End-Stage Renal Disease Quality Incentive Program (ESRD QIP); and the Quality Payment Program.

In implementing the Hospital IQR Program and other quality reporting programs, we have focused on measures which have high impact and support CMS and HHS priorities for improved quality and efficiency of care for Medicare beneficiaries. Our goal for the future is to align the clinical quality measure requirements of the Hospital IQR Program with various other Medicare and Medicaid programs, including those authorized by the Health Information Technology for Economic and Clinical Health (HITECH) Act, so that the reporting burden for providers will be reduced. As appropriate, we will consider the adoption of clinical quality measures with electronic specifications so the electronic collection of performance information is a seamless component of care delivery. Establishing such a system will require interoperability between electronic health records (EHR) and CMS data collection systems, additional infrastructure development on the part of hospitals and CMS, and adoption of standards for capturing, formatting, and transmitting the data elements that make up the measures. However, once these activities are accomplished, adoption of measures which rely on data obtained directly from EHRs will enable us to expand the Hospital IQR Program measure set with less cost and reporting burden to hospitals. We believe that, in the near future, collection and reporting of data elements through EHRs will greatly simplify and streamline reporting for various CMS quality reporting programs, and hospitals will have decreased burden as they are able to switch primarily to EHR-based data reporting for many measures that are currently manually chart-abstracted and submitted to CMS for the Hospital IQR Program.

We also have implemented a Hospital VBP Program under section 1886(o) of the Act, described in the FY 2013 Hospital Inpatient VBP Program final rule (76 FR 26490 through 26547); the FY 2014 the FY 2014 IPPS/LTCH PPS final rule (78 FR 50676 through 50707); the FY 2015 IPPS/LTCH PPS final rule (79 FR 50048 through 50087); the FY
proposing changes to the Medicare and Medicaid EHR Incentive Programs for eligible hospitals and critical access hospitals (CAHs).

A. Hospital Inpatient Quality Reporting (IQR) Program

1. Background

a. History of the Hospital IQR Program

We seek to promote higher quality and more efficient health care for Medicare beneficiaries. This effort is supported by the adoption of widely-agreed upon quality measures. We have worked with relevant stakeholders to define measures of quality in almost every setting and currently measure some aspect of care for almost all Medicare beneficiaries. These measures assess structural aspects of care, clinical processes, patient experiences with care, and outcomes. We have implemented quality measure reporting programs for multiple settings of care. To measure the quality of hospital inpatient services, we implemented the Hospital Inpatient Quality Reporting (IQR) Program, previously referred to as the Hospital Quality Data for Annual Payment Update (RHQDAPU) Program. We refer readers to the FY 2010 IPPS/LTC PPS final rule (74 FR 43860 through 43861) and the FY 2011 IPPS/LTC PPS final rule (75 FR 50180 through 50181) for detailed discussions of the history of the Hospital IQR Program, including the statutory history, and to the FY 2015 IPPS/LTC PPS final rule (79 FR 50217 through 50249), the FY 2016 IPPS/LTC PPS final rule (80 FR 49660 through 49692), and the FY 2017 IPPS/LTC PPS final rule (81 FR 57148 through 57150) for the measures we have adopted for the Hospital IQR Program measure set through the FY 2019 payment determination and subsequent years.

b. Maintenance of Technical Specifications for Quality Measures

The technical specifications for chart-abstracted clinical process of care measures used in the Hospital IQR Program, or links to Web sites hosting technical specifications, are contained in the CMS/The Joint Commission (TJC) Specifications Manual for National Hospital Inpatient Quality Measures (Specifications Manual). This Specifications Manual is posted on the QualityNet Web site at: http://www.qualitynet.org/. We generally update the Specifications Manual on a semiannual basis and include in the updates detailed instructions and calculation algorithms for hospitals to use when collecting and submitting data on required chart-abstracted clinical process of care measures.

The technical specifications for electronic clinical quality measures (eCQMs) used in the Hospital IQR Program are contained in the CMS Annual Update for Hospital Quality Reporting Programs (Annual Update). This Annual Update is posted on the eCQI Resource Center Web site at: https://ecqi.healthit.gov/. We generally update the measure specifications on an annual basis through the Annual Update, which includes code updates, logic corrections, alignment with current clinical guidelines, and additional guidance for hospitals and EHR vendors to use in order to collect and submit data on eCQMs from hospital EHRs.

In addition, we believe that it is important to have in place a subregulatory process to incorporate nonsubstantive updates to the measure specifications for measures we have adopted for the Hospital IQR Program such that these measures remain up-to-date. We refer readers to the FY 2013 IPPS/LTC PPS final rule (77 FR 53504 through 53505) and the FY 2015 IPPS/LTC PPS final rule (79 FR 50203) for our policy for using a subregulatory process to make nonsubstantive updates to measures used for the Hospital IQR Program.

We recognize that some changes made to measures undergoing maintenance review are substantive in nature and might not be appropriate for adoption using a subregulatory process. We will continue to use rulemaking to adopt substantive updates made to measures we have adopted for the Hospital IQR Program. We refer readers to the FY 2017 IPPS/LTC PPS final rule (81 FR 57111) for additional discussion of the maintenance of technical specifications for quality measures for the Hospital IQR Program. We also refer readers to the FY 2015 IPPS/LTC PPS final rule (79 FR 50202 through 50203) for additional details on the measure maintenance process.

In this proposed rule, we are not proposing any changes to our policies on the measures maintenance process, including the maintenance of nonsubstantive updates to measures used for the Hospital IQR Program.

c. Public Display of Quality Measures

Section 1886(b)(3)(B)(vii) of the Act was amended by the Deficit Reduction Act (DRA) of 2005. Section 5001(a) of the DRA requires that the Secretary establish procedures for making information regarding measures submitted available to the public after ensuring that a hospital has the opportunity to review its data before they are made public. Our current
policy is to report data from the Hospital IQR Program as soon as it is feasible on CMS Web sites such as the Hospital Compare Web site, http://www.medicare.gov/hospitalcompare after a 30-day preview period (78 FR 50776 through 50778).

Information is available to the public on the Hospital Compare Web site. Hospital Compare is an interactive web tool that assists beneficiaries by providing information on hospital quality of care to those who need to select a hospital. The Hospital IQR Program currently includes process of care measures, risk-adjusted outcome measures, the HCAHPS patient experience-of-care survey measure, structural measures, Emergency Department throughput measures, patient safety and adverse event measures, immunization measures, hospital-acquired infection measures, and payment measures, all of which are featured on the Hospital Compare Web site. For more information on measures reported to Hospital Compare, we refer readers to the Web site at: http://www.medicare.gov/hospitalcompare.

Other information that may not be as relevant to or easily understood by beneficiaries and information for which there are unresolved display issues or design considerations are not reported on Hospital Compare and may be made available on other CMS Web sites, such as https://data.medicare.gov.

In this proposed rule, we are not proposing any changes to these policies.

d. Accounting for Social Risk Factors in the Hospital IQR Program

We understand that social risk factors such as income, education, race and ethnicity, employment, disability, community resources, and social support (certain factors of which are also sometimes referred to as socioeconomic status (SES) factors or socio-demographic status (SDS) factors) play a major role in health. One of our core objectives is to improve beneficiary outcomes including reducing health disparities, and we want to ensure that all beneficiaries, including those with social risk factors, receive high quality care. In addition, we seek to ensure that the quality of care furnished by providers and suppliers is assessed as fairly as possible under our programs while ensuring that beneficiaries have adequate access to excellent care.

We have been reviewing reports prepared by the Office of the Assistant Secretary for Planning and Evaluation (ASPE) and the National Academies of Sciences, Engineering, and Medicine on the issue of measuring and accounting for social risk factors in CMS’ value-based purchasing and quality reporting programs, and considering options on how to address the issue in these programs. On December 21, 2016, ASPE submitted a Report to Congress on a study it was required to conduct under section 2(d) of the Improving Medicare Post-Acute Care Transformation (IMPACT) Act of 2014. The study analyzed the effects of certain social risk factors of Medicare beneficiaries on quality measures and measures of resource use used in one or more of nine Medicare value-based purchasing programs. The report also included considerations for strategies to account for social risk factors in these programs. In a January 10, 2017 report released by the National Academies of Sciences, Engineering, and Medicine, that body provided various potential methods for measuring and attributing social risk factors, including stratified public reporting.

As noted in the FY 2017 IPPS/LTCPPS final rule, the NQF has undertaken a 2-year trial period in which new measures, measures undergoing maintenance review, and measures endorsed with the condition that they enter the trial period can be assessed to determine whether risk adjustment for selected social risk factors is appropriate for these measures. This trial entails temporarily allowing inclusion of social risk factors in the risk-adjustment approach for these measures. At the conclusion of the trial, NQF will issue recommendations on the future inclusion of social risk factors in risk adjustment for these quality measures, and we will closely review its findings.

As we continue to consider the analyses and recommendations from these reports and await the results of the NQF trial on risk adjustment for quality measures, we are continuing to work with stakeholders in this process. As we have previously communicated, we are concerned about holding providers to potential disparities or minimize incentives to improve the outcomes for disadvantaged populations. Keeping this concern in mind, while we sought input on this topic previously, we continue to seek public comment on whether we should account for social risk factors in the Hospital IQR Program, and if so, what method or combination of methods would be most appropriate for accounting for social risk factors.

Examples of methods include: Confidential reporting to providers of measure rates stratified by social risk factors; public reporting of stratified measure rates; and potential risk adjustment of a particular measure as appropriate based on data and evidence.

In addition, we are also seeking public comment on which social risk factors might be most appropriate for reporting stratified measure scores and/or potential risk adjustment of a particular measure. Examples of social risk factors include, but are not limited to, dual eligibility/low-income subsidy, race and ethnicity, and geographic area of residence. We are seeking comments on which of these factors, including current data sources where this information would be available, could be used alone or in combination, and whether other data should be collected to better capture the effects of social risk. We will take commenters’ input into consideration as we continue to assess the appropriateness and feasibility of accounting for social risk factors in the Hospital IQR Program. We note that any such changes would be proposed through future notice-and-comment rulemaking.

We refer readers to section IX.A.13 of the preamble of this proposed rule, where we discuss the potential future confidential reporting of stratified measure data for the Hospital 30-day, All-Cause, Risk-Standardized Readmission Rate Following Pneumonia Hospitalization (NQF #0056) and the Hospital 30-day, All-Cause, Risk-Standardized Mortality Rate (RSMR) for Pneumonia measures. Our goal is to provide examples from several domains for the same issue (pneumonia). We want the reader to understand the approaches from as many perspectives as possible. In addition we are seeking comments on options for publicly displaying stratified rates using social risk factors as well as which other social risk factors besides dual eligibility should be used.

We look forward to working with stakeholders as we consider the issue of accounting for social risk factors and reducing health disparities in CMS programs. Of note, implementing any of the above methods would be taken into consideration in the context of how this and other CMS programs operate (for example, data submission methods,
availability of data, statistical considerations relating to reliability of data calculations, among others, so we also welcome comment on operational considerations. CMS is committed to ensuring that its beneficiaries have access to and receive excellent care, and that the quality of care furnished by providers and suppliers is assessed fairly in CMS programs.

2. Retention of Previously Adopted Hospital IQR Program Measures for Subsequent Payment Determinations

We refer readers to the FY 2013 IPPS/LTCH PPS final rule (77 FR 53512 through 53513) for our finalized measure retention policy. Pursuant to this policy, when we adopt measures for the Hospital IQR Program beginning with a particular payment determination, we automatically re-adopt these measures for all subsequent payment determinations unless we propose to remove, suspend, or replace the measures. In this proposed rule, we are not proposing any changes to this policy.

3. Removal and Suspension of Previously Adopted Hospital IQR Program Measures

As discussed above, we generally retain measures from the previous year’s Hospital IQR Program measure set for subsequent years’ measure sets except when we specifically propose to remove, suspend, or replace a measure. We refer readers to the FY 2011 IPPS/LTCH PPS final rule (75 FR 50185) and the FY 2015 IPPS/LTCH PPS final rule (79 FR 50203 through 50204) for more information on the criteria we consider for removing quality measures. We refer readers to the FY 2016 IPPS/LTCH PPS final rule (80 FR 49641 through 49643) for more information on the additional factors we consider in removing quality measures and the factors we consider in order to retain measures. We note in the FY 2015 IPPS/LTCH PPS final rule (79 FR 50203 through 50204), we clarified the criteria for determining when a measure is “topped-out.” In this proposed rule, we are not proposing any changes to these policies.

We refer readers to the FY 2017 IPPS/LTCH PPS final rule (81 FR 57112 through 57120) for the list of 15 measures finalized for removal for the FY 2019 payment determination and subsequent years. In this FY 2018 IPPS/LTCH PPS proposed rule, we are not proposing any measures for removal.

4. Previously Adopted Hospital IQR Program Measures for the FY 2019 Payment Determination and Subsequent Years

The Hospital IQR Program has previously finalized 62 measures for the FY 2019 payment determination and subsequent years as outlined in the table below:

### HOSPITAL IQR PROGRAM MEASURES FOR THE FY 2019 PAYMENT DETERMINATION AND SUBSEQUENT YEARS

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<td>National Healthcare Safety Network (NHSN) Catheter-associated Urinary Tract Infection (CAUTI) Outcome Measure.</td>
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<td>National Healthcare Safety Network (NHSN) Facility-wide Inpatient Hospital-onset Clostridium difficile Infection (CDI) Outcome Measure.</td>
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<td>MRSA Bacteremia</td>
<td>National Healthcare Safety Network (NHSN) Facility-wide Inpatient Hospital-onset Methicillin-resistant Staphylococcus aureus (MRSA) Bacteremia Outcome Measure.</td>
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<td><strong>Claims-Based Patient Safety Measures</strong></td>
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HOSPITAL IQR PROGRAM MEASURES FOR THE FY 2019 PAYMENT DETERMINATION AND SUBSEQUENT YEARS—Continued

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<tr>
<td>READM–30–COPD</td>
<td>Hospital 30-Day, All-Cause, Risk-Standardized Readmission Rate (RSRR) Following Chronic Obstructive Pulmonary Disease (COPD) Hospitalization.</td>
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<tr>
<td>READM–30–HF</td>
<td>Hospital 30-Day, All-Cause, Risk-Standardized Readmission Rate (RSRR) Following Heart Failure (HF) Hospitalization.</td>
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Claims-Based Payment Measures

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<td>THA/TKA Payment</td>
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<td>Cellulitis Payment</td>
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Chart-Abstracted Clinical Process of Care Measures

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<td>Imm-2*</td>
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<td>PC-01*</td>
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<td>Sepsis</td>
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EHR-Based Clinical Process of Care Measures (that is, Electronic Clinical Quality Measures (eCQMs))

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Patient Experience of Care Survey Measures

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<td>Patient Safety Culture (Hospital Survey on Patient Safety Culture)</td>
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</tr>
<tr>
<td>Safe Surgery Checklist Use</td>
<td>N/A</td>
</tr>
</tbody>
</table>

Structural Patient Safety Measures

* Measure listed twice, as both chart-abstracted and electronic clinical quality measure.
+ NQF endorsement has been removed.
5. Considerations in Expanding and Updating Quality Measures

We refer readers to the FY 2013 IPPS/LTCH PPS final rule (77 FR 53510 through 53512) for a discussion of the considerations we use to expand and update quality measures under the Hospital IQR Program. In this proposed rule, we are not proposing any changes to these policies.

6. Refinements to Existing Measures in the Hospital IQR Program for the FY 2020 Payment Determination and Subsequent Years

In this proposed rule, we are proposing refinements to two measures. First, we are proposing refinements to the Hospital Consumer Assessment of Healthcare Providers and Systems (HCAHPS) Survey (NQF #0166) measure for the FY 2020 payment determination and subsequent years. Second, we are proposing refinements to the Stroke 30-Day Mortality Rate (MORT–30–STK) measure for the FY 2023 payment determination and subsequent years. We discuss these refinements in more detail below.

a. Refining the Hospital Consumer Assessment of Healthcare Providers and Systems (HCAHPS) Survey (NQF #0166) Measure for the FY 2020 Payment Determination and Subsequent Years

For the FY 2020 payment determination and subsequent years, we are proposing to refine the existing Hospital Consumer Assessment of Healthcare Providers and Systems (HCAHPS) Survey by refining the current Pain Management questions (HCAHPS Q12, Q13, and Q14) to focus on the hospital’s communications with patients about the patients’ pain during the hospital stay. In accord with this new focus, we are proposing to update the name of the composite measure from “Pain Management” to “Communication About Pain.”

(1) Background

The HCAHPS Survey (NQF #0166) was adopted in the Reporting Hospital Quality Data Annual Payment Update Program in the CY 2007 OPPS/ASC final rule (71 FR 68201), beginning with the FY 2008 payment determination and for subsequent years. This Survey includes three Pain Management questions, Q12, Q13 and Q14. In the FY 2013 IPPS/LTCH PPS final rule (77 FR 53513 through 53516), we added the Care Transition Measure (CTM–3) (NQF #0228) to the existing HCAHPS Survey, NQF #0166. The HCAHPS Survey, combining both NQF #0166 for the original survey and NQF #0228 for the Care Transition Measure adopted into The HCAHPS Survey in 2013, is the first national, standardized, publicly reported survey of patients’ experience of hospital care. The HCAHPS Survey asks discharged patients 32 questions about their recent hospital stay. Survey results have been publicly reported on the Hospital Compare Web site since 2008. We refer readers to the FY 2011 IPPS/LTCH PPS final rule (75 FR 50220), the FY 2012 IPPS/LTCH PPS final rule (76 FR 51641 through 51643), the FY 2013 IPPS/LTCH PPS final rule (77 FR 53537 through 53538), and the FY 2014 IPPS/LTCH PPS final rule (78 FR 50819 through 50820) for details on previously-adopted HCAHPS requirements. We also refer hospitals and HCAHPS Survey vendors to the official HCAHPS Web site at: http://www.hcahpsonline.org for new information and program updates regarding the HCAHPS Survey, its administration, oversight, and data adjustments.

The HCAHPS Survey (OMB control number 0938–0981) is administered to a random sample of adult patients who receive medical, surgical, or maternity care between 48 hours and 6 weeks (42 calendar days) after discharge and is not restricted to Medicare beneficiaries. Hospitals must survey patients throughout each month of the year. The HCAHPS Survey is available in official English, Spanish, Chinese, Russian, Vietnamese, and Portuguese versions. The HCAHPS Survey and its protocols for sampling, data collection and coding, and file submission can be found in the current HCAHPS Quality Assurance Guidelines, which is available on the official HCAHPS Web site at: http://www.hcahpsonline.org/qaguidelines.aspx. AHRQ carried out a rigorous, scientific process to develop and test the HCAHPS instrument. This process entailed multiple steps, including: a public call for measures; literature reviews; cognitive interviews, consumer focus groups; multiple opportunities for additional stakeholder input; a 3-State pilot test; small-scale field tests; and notice and comment rulemaking. We refer readers to the CY 2007 OPPS final rule (71 FR 68201) for a more in-depth discussion about this process. The HCAHPS Survey was endorsed by the NQF on August 5, 2005 (#0166).

The Pain Management questions currently included in the HCAHPS Survey are as follows:

12. During this hospital stay, did you need medicine for pain?

☐ Yes

☐ No → If No, Go to Question 15

13. During this hospital stay, how often was your pain well controlled?

☐ Never

☐ Sometimes

☐ Usually

☐ Always

14. During this hospital stay, how often did the hospital staff do everything they could to help you with your pain?

☐ Never

☐ Sometimes

☐ Usually

☐ Always

In the CY 2017 OPPS/ASC final rule with comment period in the context of the Hospital VBP Program (81 FR 79856), we stated that we received feedback that some stakeholders are concerned about the Pain Management dimension questions being used in a program where there is any link between scoring well on the questions and higher hospital payments (81 FR 79856). The Pain Management dimension used in the Hospital VBP Program is identical in composition to the Pain Management measure used in the Hospital IQR Program, questions Q12, Q13 and Q14 with one difference: The HCAHPS dimension score in the Hospital VBP program is based on the percentage of patients who chose the most positive response option (“top-box” response). For more information about the Hospital VBP Program scoring methodology, we refer readers to the FY 2017 IPPS/LTCH PPS final rule (81 FR 57006).

Some stakeholders believed that the linkage of the Pain Management dimension questions to the Hospital VBP Program payment incentives created pressure on hospital staff to prescribe more opioids in order to achieve higher scores on this dimension (81 FR 79856). We stated that we continue to believe that pain control is an appropriate part of routine patient care that hospitals should manage and is an important concern for patients, their families, and their caregivers (81 FR 79856). Further, we stated that it is important to note that the HCAHPS Survey does not specify any particular type of pain control method (81 FR 79856). We added that appropriate pain management includes communication with patients about pain-related issues, setting expectations about pain, shared decision-making, and proper prescription practices (81 FR 79856). Furthermore, we stated that although we were not aware of any scientific studies that support an association between scores on the Pain Management dimension questions and opioid prescribing practices, we were developing alternative questions for the Pain Management dimension in order to remove any potential ambiguity in the HCAHPS Survey. We noted that we believe that removing the Pain Management dimension from the Hospital VBP Program scoring calculations would address potential confusion about the appropriate use of the Pain Management dimension, and provide us with an opportunity to further refine the pain management questions used in the HCAHPS Survey (81 FR 79859).

In the same final rule, we stated we would follow our standard survey development processes, which included drafting alternative questions, cognitive interviews and focus group evaluation, field testing, statistical analysis, stakeholder input, the Paperwork Reduction Act, and NQF endorsement (81 FR 79856).

In that final rule, numerous commenters supported the development of modified questions regarding pain management for the HCAHPS Survey and some commenters expressed particular support for modified pain management questions that focused on effective communication with patients about pain management-related issues (81 FR 79859 through 79860).
Specifically, a number of commenters recommended modified pain management questions focused on shared decision-making, discussion of treatment options, including non-opioid pain management therapies, patient understanding of pain management options, and patient engagement in their care (81 FR 79860).

Therefore, for the FY 2020 payment determination and subsequent years, we are proposing to update and refine the existing HCAHPS Survey questions (HCAHPS Q12, Q13, and Q14) to focus more directly on communication with patients about their pain during the hospital stay. These proposed revised questions will be used to form the composite measure “Communication about Pain.” The “Communication about Pain” composite measure would be a part of the HCAHPS Survey and would be publicly reported in the Hospital IQR Program. More information about the revised questions/composite measure is included below.

In compliance with section 1890(a)(2) of the Act, measures proposed for the Hospital IQR Program were included in a publicly available document: “List of Measures under Consideration for December 1, 2016” available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment- Instruments/QualityMeasures/Downloads/Measures-under-Consideration-List-for-2016.pdf.

The Measure Applications Partnership (MAP), a multi-stakeholder group convened by the NQF, reviews the measures under consideration for the Hospital IQR Program, among other Federal programs, and provides input on those measures to the Secretary. The MAP’s 2017 recommendations for quality measures under consideration are documented in the following documents: “2016–2017 Process and Approach for MAP Pre Rulemaking Deliberations” available at: http://www.qualityforum.org/WorkArea/linkit.aspx?LinkIdentifier=id&ItemID=84455 and “2016–2017 Spreadsheet of Final Recommendations to HHS and CMS” available at: http://www.qualityforum.org/WorkArea/linkit.aspx?LinkIdentifier=id&ItemID=84452. We considered the input and recommendations provided by the MAP.

The Communication About Pain (MUC16–263) composite measure was reviewed by the MAP in December 2016. The MAP recommended that this composite measure be refined and resubmitted prior to rulemaking. The MAP emphasized the need to include non-pharmacological options used to treat pain. The MAP recommended that the testing results demonstrate reliability and validity for the Hospital IQR Program. The MAP also recommended that the measure be submitted to NQF for review and endorsement.

We plan to resubmit the “Communication About Pain” composite measure to the MAP at the next opportunity. As we discuss in more detail below, the Communication About Pain composite measure underwent field testing in 2016. Results were not yet available for the MAP’s review in December 2016, but are now complete and will be posted on the official HCAHPS On-Line Web site, www.HCAHPSonline.org. We believe the measure is now fully developed and tested and we intend to provide feedback to the MAP Hospital Workgroup for review of testing results.

In early 2016, we empirically tested as part of the field test the reliability and validity of the Communication About Pain questions in a large-scale experiment that involved patients from 50 hospitals across the nation. Our analyses suggest the Communication About Pain composite measure, which includes two substantive items regarding how often staff talked about pain and how often staff discussed how to treat pain while in the hospital (Q13 and Q14), as well as a screener item (Q12), have strong reliability (evidence that scores for hospitals are precisely measured) and validity (evidence that the measure does measure the intended construct of patient experience). These properties of the individual questions used in the proposed Communication About Pain composite measure are as good as or better than the current Pain Management questions. The new questions are not subject to floor or ceiling effects (which would occur if almost all responses were in the lowest or highest response category), have excellent hospital-level reliability (here 0.88 or higher, where 0.70 or higher is the conventional standard) at recommended sample sizes, are not redundant with other current questions, are related in a predictable manner with the standard patient-mix characteristics, positively correlate with the two HCAHPS questions that assess overall patient experience (rating and recommendation) with the hospital, providing evidence of validity and do not vary systematically by survey mode, patient race/ethnicity, or hospital characteristics after adjusting for patient mix. They also have higher internal consistency as a composite measure (Cronbach’s alpha = 0.81), with 0.70 or higher being the conventional standard, further providing evidence of reliability.

As stated above, the MAP recommended the proposed Communication About Pain composite measure be submitted to the NQF for review and endorsement once testing has been completed. The proposed Communication About Pain composite measure is not yet NQF endorsed; however, we intend to submit the measure to the NQF for endorsement when the Person and Family Centered Care Project has a call for measures.

Whenever feasible, we adopt measures that are NQF-endorsed, but note sometimes there are important areas of clinical concern for which NQF endorsed measures do not exist. Section 1886(b)[3][IX][b] of the Act provides that in the case of a specified area or medical topic determined appropriate by the Secretary for which a feasible and practical measure has not been endorsed by the entity with a contract under section 1890(a) of the Act, the Secretary may specify a measure that is not so endorsed as long as due consideration is given to measures that have been endorsed or adopted by a consensus organization identified by the Secretary. (The NQF currently holds this contract.) We considered other existing measures which have been endorsed by NQF and other consensus organizations, but we were unable to identify any NQF-endorsed (or other consensus organization endorsed) measures that were feasible and practical.

While we consider MAP recommendations and NQF endorsement status as part of our decision-making process for which measures to include in the Hospital IQR Program, we believe it is important to adopt this proposed Communication About Pain composite measure because communicating with patients about their pain is an integral part of delivering high quality, person-centered care. In developing the proposed
Communication About Pain composite measure, we followed our standard survey development processes, which included drafting alternative questions, cognitive interviews, focus group evaluation, field testing, statistical analysis, and stakeholder input. We believe the Communication About Pain composite measure has been sufficiently tested, demonstrating high levels of reliability and validity, as noted above.

Further, we have consistently received feedback from some stakeholders expressing concern that the current Pain Management questions encourage overprescribing of opioids as discussed in the CY 2017 OPPS/ASC final rule with comment period (81 FR 79856). As a result, we believe it is important to refine the existing Pain Management measure. We note that if our proposal to revise the current Pain Management measure questions with those in the proposed Communication About Pain composite measure is not finalized, we would continue to use the Pain Management questions as previously finalized.

The Communication About Pain composite measure is discussed below. We are proposing to revise the current Pain Management questions (Q12, Q13, and Q14) in the HCAHPS Survey for the FY 2020 payment determination and subsequent years by adopting the Communication About Pain composite measure in the HCAHPS Survey beginning with the FY 2020 payment determination, which would be applicable to surveys administered to patients beginning with January 1, 2018 discharges and for subsequent years.

In compliance with section 1886(b)(3)(B)(vi) of the Act, we calculate and publicly report HCAHPS measures from four consecutive quarters of data. From that point and forward, the oldest quarter of data is rolled off, the newest quarter is rolled on, and the measure scores are calculated for this unique set of four quarters and are publicly reported on Hospital Compare and available for payment determination. Data submitted for the current Pain Management measure in CY 2017 for the FY 2019 payment determination will be publicly reported on Hospital Compare in October 2018. If our proposal to revise the HCAHPS Pain Management measure with the HCAHPS Communication About Pain composite measure is finalized, we would begin to use the new Pain Management items on the HCAHPS Survey in January of 2018. Once we have collected four consecutive quarters of the HCAHPS Communication About Pain composite measure questions, we will create scores for the Communication About Pain composite measure.

We will be unable to report or use for payment determination either the original or new Pain Management measure unless and until we have collected four quarters of data for the measure. The CY 2017 reporting period/FY 2019 payment determination will be the last period for which we have four quarters of the original Pain Management measure data which, as stated above, will be publicly reported on Hospital Compare in October 2018.

The refined questions that comprise the proposed Communication About Pain composite measure closely mirror the structure and style of the existing Pain Management questions; however, the new questions address how providers communicate with patients about pain while removing any ambiguities in the wording or intent of the questions. This refinement is consistent with the HCAHPS Survey’s original design, development, and NQF endorsement (NQF #0166). Further, we designed the Communication About Pain composite measure to be consistent and compatible with existing HCAHPS questions and HCAHPS sampling and survey administration protocols. The three Communication About Pain composite measure questions are as follows:

www.annemergmed.com/article/S0196-0964(16)30367-5/fulltext

As stated above, in light of the ongoing opioid epidemic, we believe it is important the Communication About Pain composite measure is abundantly clear in its focus on communication about pain between providers and their patients and be applicable to all patients who experienced pain during their hospital stay.

(3) Data Collection

The revised Communication About Pain composite measure questions would be administered and data collected in exactly the same manner as the current Pain Management measure questions; there would be no changes to HCAHPS patient eligibility or exclusion criteria. Detailed information on HCAHPS data collection protocols can be found in the current HCAHPS Quality Assurance Guidelines, located at: http://www.hcahpsonline.org/qaguidelines.aspx. We reiterate that other than the revision of the HCAHPS Pain Management questions, the HCAHPS Survey and its administration and data collection protocols would be unchanged. The survey adjustment and patient-mix adjustment for the new Communication About Pain composite measure would be made available on the official HCAHPS On-Line Web site at: http://www.hcahpsonline.org/modadjustment.aspx.

(4) Public Reporting

The scoring of the new Communication About Pain composite measure would be the same as the current Pain Management measure. Detailed information on how the measure would be scored for purposes of public reporting can be found on the HCAHPS Web site at: http://www.hcahpsonline.org/Files/Calculation%20of%20HCAHPS%20Scores.pdf.

We are inviting public comment on our proposal to revise the current Pain Management questions (Q12, Q13, and Q14) in the HCAHPS Survey for the FY 2020 payment determination and subsequent years by adopting the proposed Communication About Pain composite measure in the HCAHPS Survey beginning with the FY 2020 payment determination and subsequent years, which would be applicable to surveys administered to patients beginning with January 1, 2018 discharges and for subsequent years as discussed above.

b. Refinement of the Hospital 30-Day, All-Cause, Risk-Standardized Mortality Rate (RSMR) Following Acute Ischemic Stroke Hospitalization Measure for the FY 2023 Payment Determination and Subsequent Years

(1) Background

For the FY 2023 payment determination and subsequent years, we are proposing a refinement of the CMS Hospital 30-Day, All-Cause, Risk-Standardized Mortality Rate (RSMR) Following Acute Ischemic Stroke Hospitalization Measure (hereafter referred to as the Stroke 30-Day...
Mortality Rate measure) by changing the measure’s risk adjustment to include stroke severity (Stroke 30-Day Mortality Rate with the refined risk adjustment) obtained from International Classification of Disease, Tenth Edition Clinical Modifier (ICD–10–CM) codes in the administrative claims. The current Stroke 30-Day Mortality Rate measure was finalized in the Hospital IQR Program in the FY 2014 IPPS/LTC PPS final rule (78 FR 50798). The previously adopted measure includes 42 risk variables, but does not include an assessment of stroke severity because, previously, it has not been available in claims data and was not routinely performed by all providers. For more details on the measure as currently adopted and implemented, we refer readers to its measure methodology report and measure risk-adjustment statistical model in the AMI, HF, PN, COPD, and Stroke Mortality Update zip file on our Web site at: http://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment- Instruments/HospitalQualityInitiatives/Metric-Medical-Methodology.html.

In the FY 2017 IPPS/LTC PPS final rule (81 FR 57161), we considered potential inclusion of the National Institutes of Health (NIH) Stroke Scale for the Hospital 30-Day Mortality Following Acute Ischemic Stroke Hospitalization measure beginning as early as the FY 2022 payment determination. Commenters generally supported the inclusion of the NIH Stroke Scale score in the Stroke 30-Day Mortality Rate measure for future inclusion in the Hospital IQR Program. We refer readers to FY 2017 IPPS/LTC PPS final rule (81 FR 57161 through 57163) for a complete discussion of the considered potential measure, public comments, and our responses.

Initial assessment of stroke severity, such as the NIH Stroke Scale score, is one of the strongest predictors of mortality in ischemic stroke patients,108 109 110 and is part of the national guidelines on stroke care.111 This measure refinement was developed in collaboration with the American Heart Association (AHA) and American Stroke Association (ASA). We are seeking to update the current measure to include an assessment of stroke severity, because it has become feasible to do so due to both the increased use of the NIH Stroke Scale related to the AHA/ASA guidelines that recommend administering the NIH Stroke Scale on all stroke patients, as well as due to the recent ability to obtain the scores through claims data by incorporation into ICD–10. The proposed refinement would create a more parsimonious risk model by reducing the total number of risk adjustment variables from 42 to 20 and includes the NIH Stroke Scale112 in the risk-adjustment model as a measure of stroke severity. These refinements result in a modestly higher c-statistic compared with the risk-adjustment model in the current Stroke 30-Day Mortality Rate, which means that the updated measure model better differentiates the risk of mortality among patients.

Mortality following stroke is an important adverse outcome which can be measured reliably and objectively and is influenced by both the severity of the stroke as well as the quality of care provided to patients during their initial hospitalization; therefore, mortality is an appropriate measure of quality of care following stroke hospitalization.113 114 Specifically, post-stroke mortality rates have been shown to be influenced by critical aspects of care such as response to complications, speediness of delivery of care, organization of care, and appropriate imaging.115 116 117 118

We are proposing a refinement to the Stroke 30-Day Mortality Rate for several reasons. First, the proposed, refined measure would allow for more rigorous risk adjustment by incorporating the NIH Stroke Scale, discussed in more detail below, as an assessment of stroke severity.119 Second, the inclusion of the NIH Stroke Scale is aligned with and supportive of clinical guidelines, as use of the NIH Stroke Scale to assess stroke severity when patients first present with acute ischemic stroke is Class I recommended in the AHA and ASA guidelines.120

Third, in October 2016, the ICD–10–CM codes for the NIH Stroke Scale were implemented. As of that date, hospitals can record the NIH Stroke Scale as a representation of stroke severity in Medicare claims by using ICD–10–CM codes, and we can use this information as a variable in the risk-adjustment model for the refined Stroke 30-Day Mortality Rate measure and other claims-based measures with minimal data collection burden for hospitals.121

Fourth, clinicians and stakeholders, including AHA, ASA, and other professional organizations, highlight the importance of including an assessment of stroke severity in risk-adjustment models of stroke mortality.122 In the FY 2014 IPPS/LTC PPS final rule (78 FR 50798 through 50802), commenters emphasized that the medical literature and their own experience suggest that stroke severity is the dominant predictor of mortality in stroke patients; individuals and organizations expressed concern that the measure might be misleading, limited, or inaccurate without adjustment for stroke severity, and four comments suggested risk

112 NIH Stroke Scale. Available at: http://www.nihstrokescale.org/
119 NIH Stroke Scale. Available at: http://www.nihstrokescale.org/
adjusting using the NIH Stroke Scale or a similar index (78 FR 50800).

Members of the Technical Expert Panel convened by the measure developer also suggested risk adjusting for stroke severity. In addition, during the 2012 Neurology Endorsement Maintenance Consensus Development Project, the NQF Neurology Steering Committee specifically identified the lack of the NIH Stroke Scale score in the risk-adjustment model as a concern (78 FR 50800). Therefore, the refined Stroke 30-Day Mortality Rate is responsive to public comments from a broad array of stakeholder groups, including clinical societies and clinical experts, and to feedback received from the Technical Expert Panel convened by the measure developer (81 FR 57162).

Fifth, in addition to a modestly higher c-statistic, which evaluates the measure’s ability to differentiate between patients at different risk of mortality following acute ischemic stroke, the refined Stroke 30-Day Mortality Rate includes a more parsimonious risk model than the stroke mortality measure as previously adopted and specified, with a total of 20 risk adjustment variables including the NIH Stroke Scale, compared to the current use of 42 risk adjustment variables.

In compliance with section 1890A(a)(2) of the Act, the Stroke 30-Day Mortality Rate (MUC15–294) with the refined risk adjustment (using the NIH Stroke Scale) was included on a publicly available document entitled “List of Measures under Consideration for December 2015” (available at: http://www.qualityforum.org/ProjectMaterials.aspx?projectID=75367. Select “2015 Measures Under Consideration List.”). The MAP reviewed and conditionally supported the Stroke 30-Day Mortality Rate (MUC15–294) with the refined risk adjustment pending NQF review and endorsement, and asked that we consider a phased approach in regards to implementation, to avoid multiple versions of the same measure. The MAP also noted that outcomes other than mortality may be more meaningful for stroke patients and to consider cognitive or functional outcomes such as impaired capacity. We considered the input and recommendations provided by the MAP and note that the NIH Stroke Scale incorporates cognitive functions in assessing severity.

To avoid implementing multiple versions of the same measure, we intend for the Hospital IQR Program FY 2023 payment determination measure set either to include the 30-day stroke mortality measure as currently implemented or this modified version that includes the NIH stroke severity scale in the measures risk-adjustment model.

The Stroke 30-Day Mortality Rate with the refined risk adjustment was submitted to NQF for endorsement in the neurology project on January 15, 2016, and did not obtain endorsement. NQF endorsement was not granted primarily due to the inability to test the validity of NIH Stroke Scale data elements derived from Medicare claims prior to implementation of the new ICD–10–CM codes in October 2016. The NQF Consensus Standards Advisory Committee (CSAC) supported the concern of the NQF committee regarding our inability to test the measure using ICD–10–CM codes since the codes were not implemented until October 2016. While we provided risk-standardized mortality rates using data from Medicare administrative claims data from the Get with the Guidelines-Stroke Registry, the Committee noted that we could not validate the National Institutes of Health Stroke Scale (NIH Stroke Scale) against ICD–10–CM codes at the time the measure was considered for endorsement. The CSAC also acknowledged that the primary reason for upholding the Committee’s decision was based on the lack of testing using ICD–10–CM codes. This measure went through the same rigorous development process as the other publicly reported outcomes measures and involved extensive input by stakeholders and clinical experts. It follows the same scientific approach to evaluate hospital performance as other Hospital IQR Program outcome measures.

When the NQF committee considered the scientific acceptability of the Stroke 30-Day Mortality Rate measure, 19 of 22 members voted that the measure met the NQF’s evidence criterion, 19 members voted that the measure met the high or moderate standard for the Performance Gap, 18 members voted that the measure met high or moderate standard for the Performance Gap, 18 members voted that the measure met high or moderate standard for the Performance Gap, 18 members voted that the measure met high or moderate standard for feasibility, and 18 members voted that the measure met the moderate standard for Use and Usability. We tested and validated the measure using NIH Stroke Scale data derived from medical record review done by the Get With The Guidelines (GWTG)-Stroke registry data supplied by AHA/ASA. The NQF committee ultimately determined that the validity testing was not sufficient for endorsement.

However, we believe that the inclusion of the NIH Stroke Scale score in the measure’s risk-adjustment model improves upon the Stroke 30-Day Mortality Rate measure which is currently publicly reported on Hospital Compare and has been implemented in the Hospital IQR Program since FY 2016 (78 FR 50802). This is supported by the improved risk-adjustment model performance. For example, the c-statistic, which is a measure of the ability to discriminate between patients at low and high risk of mortality following ischemic stroke, associated with the new, modified risk-adjustment model was 0.81 in the measure development sample, compared with a c-statistic of 0.75 in the most recent measurement period for the Stroke 30-Day Mortality Rate measure that is currently implemented in the Hospital IQR Program.

The new refined Stroke 30-Day Mortality Rate measure also has increased face validity which is supported by the comments received from stakeholders. For example, we received comments that the more rigorous risk adjustment facilitated by the NIH Stroke Scale would help ensure that the measure accurately risk adjusts for different hospital populations without unfairly penalizing high-performance providers, and that the NIH Stroke Scale is well validated, highly reliable, widely used, and a strong predictor of mortality and short- and long-term functional outcomes. However, we were not able to test the ICD–10 CM codes for NIH Stroke Scale score in claims during measure development because those codes were


124 The memo regarding the CSAC’s decision is available at: http://www.qualityforum.org/WorkArea/linkit.aspx?LinkIdentifier=id&ItemID=83217.

125 The memo regarding the CSAC’s decision is available at: http://www.qualityforum.org/WorkArea/linkit.aspx?LinkIdentifier=id&ItemID=83217.


not available for hospitals to use in their claims until October 2016. Therefore, we are proposing this measure now to inform hospitals that they should begin to include the NIH stroke severity scale codes in the claims they submit for patients with a discharge diagnosis of ischemic stroke. Once hospitals have submitted these data, it will be possible for us to examine the completeness of these data in reevaluation of the new refined Stroke 30-Day Mortality Rate measure before the proposed measure dry run and before the proposed implementation in the Hospital IQR Program. Once that testing is complete we will submit the retested measure to the NQF for endorsement prior to implementation.

Section 1886(b)(3)(B)(IX)(bb) of the Act provides that in the case of a specified area or medical topic determined appropriate by the Secretary for which a feasible and practical measure has not been endorsed by the entity with a contract under section 1890(a) of the Act, the Secretary may specify a measure that is not so endorsed as long as due consideration is given to measures that have been endorsed or adopted by a consensus organization identified by the Secretary. Although the proposed measure and the existing Stroke 30-Day mortality measure are not currently NQF-endorsed, we considered other available measures which have been endorsed or adopted by the NQF, and were unable to identify any other NQF-endorsed measures that assess stroke mortality with a standard period of follow-up. We also are not aware of any other 30-day stroke mortality measures that have been endorsed or adopted by a consensus organization.

However, we are proposing this measure now because we believe that the modifications to the measure’s risk-adjustment model represent a substantial improvement over the Stroke 30-Day Mortality Rate measure that is currently publicly reported and implemented in the Hospital IQR Program and which does not include an assessment of stroke severity in the risk-adjustment model. In addition, by announcing our intention to include the Refined 30-Day Stroke Mortality Rate measure in the Hospital IQR Program in advance of implementation for FY 2023 payment determination and subsequent years, and by describing the proposed additional testing, dry run, and our intent to re-submit the measure to NQF once the NIH Stroke Scale data become available in claims, we are providing information that hospitals require to plan and begin to alter clinical workflows and billing processes in order to capture the NIH Stroke Scale score and include it in Medicare claims. Further, this notice will allow hospitals to complete collecting NIH Stroke Scale data over the three-year time period needed for measure calculation and implementation prior to any payment adjustment. The measure, as refined, is described in more detail below.

(2) Overview of Refined Measure

The measure cohort is aligned with the currently adopted Stroke 30-Day Mortality Rate measure. In addition, the data sources (Medicare Fee-For-Service (FFS) claims), three-year reporting period, inclusion and exclusion criteria, as well as the assessment of the outcome of mortality (assessed using Medicare enrollment data) would all align with the currently adopted measure (78 FR 50798). Only the measures’ risk-adjustment models differ, as described in detail below. For the new refined Stroke 30-Day Mortality Rate measure, we are proposing that the first measurement period would include discharges between July 1, 2018 and June 30, 2021 for public reporting in FY 2022 and for the FY 2023 payment determination.

(3) Risk Adjustment

The Stroke 30-Day Mortality Rate measure that is currently adopted in the Hospital IQR Program adjusts for differences in patients’ level of risk for death in one hospital relative to patients receiving care in another hospital but not for stroke severity. For details about the risk-adjustment model for the currently adopted measure, we refer readers to the Technical Report (78 FR 50798). However, in developing the proposed, refined Stroke 30-Day Mortality Rate measure, we re-selected risk variables, resulting in a final model with 20 risk-adjustment variables, including the NIH Stroke Scale risk variable as an assessment of stroke severity. The NIH Stroke Scale is a 15-item neurologic examination used to include a quantitative measure of stroke-related neurologic deficit. The NIH Stroke Scale evaluates the effect of acute ischemic stroke on a patient’s level of consciousness, language, neglect, visual-field loss, extra-ocular movement, motor strength, ataxia (the loss of full control of bodily movements), dysarthria (difficult or unclear articulation of speech), and sensory loss. The NIH Stroke Scale was designed to be a simple, valid, and reliable assessment tool that can be administered at the bedside consistently by neurologists, physicians, nurses, or therapists, and is Class I recommended in the AHA/ASA guidelines. The NIH Stroke Scale is a publicly available standardized tool, the results of which should be assessed by a clinician when first examining a patient presenting to the hospital with a stroke and then documented in the patient’s medical record. Once this information has been documented by a clinician, it can then be recorded in the claim for that hospital admission using ICD-10-CM codes through the hospital’s normal coding practices.

We sought to develop a risk-adjustment model that included the NIH Stroke Scale variable and other key variables which we believe are clinically relevant and demonstrate a strong statistical association with 30-day mortality. To select candidate variables, we considered those 42 risk-adjustment variables in the currently adopted measure, plus the NIH Stroke Scale as a candidate variable. We then performed a bootstrapping simulation method for variable selection. This bootstrapping simulation method is a means of creating multiple samples to determine which risk variables are most important to include in a model. We selected the best model using the logistic regression model with the stepwise selection method based on 1,000 bootstrapping samples for each copy of the multiple imputed (MI) data. Variable selection rate for all the variables selected into the best model was calculated for each copy of the MI data, and variables were included into the final model if the minimum variable selection rate among the 5 copies of MI was 90 percent or more. This method resulted in 20 risk-adjustment variables that were included more than 90 percent of the time for all the copies of the imputed data were retained in the final model, including the NIH Stroke Scale. For more details on the risk-adjustment variable selection process, we refer readers to the measure methodology report and measure risk-adjustment statistical information on AMI, HF, PN, COPD, and Stroke Mortality Update zip file on our Web site at: http://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInits/Measure-Methodology.html.

Refining the risk adjustment model of the Stroke 30-Day Mortality Rate created...
a modestly higher c-statistic with fewer risk variables, meaning that the proposed, refined measure’s risk-adjustment model better distinguishes among patients with a low risk and high risk of mortality following ischemic stroke compared with the Stroke 30-Day Mortality Rate measure that is currently implemented in the Hospital IQR Program. Including the NIH Stroke Scale in the risk-adjustment model allows the measure to more accurately account for patients’ status upon arrival at the hospital, which is responsive to clinical guidelines and feedback from the medical community and other stakeholders, as discussed above.

In order to use the NIH Stroke Scale data in the proposed, refined Stroke 30-Day Mortality Rate measure, many hospitals that have not routinely captured these data on patients with ischemic stroke will need to implement new workflows to ensure that their clinicians measure and record stroke severity. In addition, hospital coders will need to include the appropriate ICD–10 code for the clinician’s documented NIH Stroke Scale score in the Medicare claim. By proposing this measure, we are providing hospitals the information and advanced notice that they would be required to submit this information in their Medicare claims for payment. In order to have information on the severity of patients’ ischemic stroke included in the calculation Stroke 30-Day Mortality Rate, some hospitals that do not currently capture or record the NIH Stroke Scale would have to create workflows and processes to do this. This additional work, however, is consistent with current clinical guidelines for the care of ischemic stroke patients, and are consistent with the standard of care. Implementation of the proposed, refined Stroke 30-Day Mortality Rate with the refined risk adjustment would require hospitals to document in the medical record the first NIH Stroke Scale on every eligible patient who is admitted for treatment of acute ischemic stroke and provide that information among the ICD–10–CM code recorded on the claim. The new ICD–10–CM code representing the NIH Stroke Scale will be included in the risk adjustment model for the Stroke 30-Day Mortality Rate measure.

Because many hospitals would have to create new clinical workflows to assess and document the NIH Stroke Scale in patients’ medical records as well as include the appropriate ICD–10 CM code for the documented NIH Stroke Scale score in the claim they submit, we would provide hospitals with dry-run results of this proposed, refined measure in their confidential hospital-specific feedback reports prior to implementation of the proposed, refined measure for the FY 2023 payment determination. For example, we anticipate using claims data, which would include ICD–10 CM codes for the NIH Stroke Scale, for discharges occurring between October 1, 2017 through June 1, 2020, to calculate measure results for the dry-run anticipated in CY 2021. The data in the confidential hospital-specific feedback reports would not be publicly reported.

We are inviting public comment on our proposal to adopt a refinement of the Stroke 30-Day Mortality Rate in the Hospital IQR Program for the FY 2023 payment determination and subsequent years as discussed above.

c. Summary of Previously Adopted Hospital IQR Program Measures for the FY 2020 Payment Determination and Subsequent Years

The table below outlines the Hospital IQR Program measure set (including previously adopted measures and proposed refinements from this proposed rule) for the FY 2020 payment determination and subsequent years. The proposed, refined measures, as discussed above, are denoted with a superscript as defined in the legend below the table.
## PREVIOUSLY ADOPTED HOSPITAL IQR PROGRAM MEASURES FOR THE FY 2020 PAYMENT DETERMINATION AND SUBSEQUENT YEARS—Continued

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<thead>
<tr>
<th>Short name</th>
<th>Measure name</th>
<th>NQF #</th>
</tr>
</thead>
<tbody>
<tr>
<td>PSI 90</td>
<td>Patient Safety for Selected Indicators Composite Measure, Modified PSI 90 (Updated Title: Patient Safety and Adverse Events Composite).</td>
<td>0531</td>
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### Claims-Based Mortality Outcome Measures

<table>
<thead>
<tr>
<th>Short name</th>
<th>Measure name</th>
<th>NQF #</th>
</tr>
</thead>
<tbody>
<tr>
<td>MORT–30–AMI</td>
<td>Hospital 30-Day, All-Cause, Risk-Standardized Mortality Rate (RSMR) Following Acute Myocardial Infarction (AMI) Hospitalization.</td>
<td>0230</td>
</tr>
<tr>
<td>MORT–30–CABG</td>
<td>Hospital 30-Day, All-Cause, Risk-Standardized Mortality Rate (RSMR) Following Coronary Artery Bypass Graft (CABG) Surgery.</td>
<td>2558</td>
</tr>
<tr>
<td>MORT–30–COPD</td>
<td>Hospital 30-Day, All-Cause, Risk-Standardized Mortality Rate (RSMR) Following Chronic Obstructive Pulmonary Disease (COPD) Hospitalization.</td>
<td>1893</td>
</tr>
<tr>
<td>MORT–30–HF</td>
<td>Hospital 30-Day, All-Cause, Risk-Standardized Mortality Rate (RSMR) Following Heart Failure (HF) Hospitalization.</td>
<td>0229</td>
</tr>
<tr>
<td>MORT–30–PN</td>
<td>Hospital 30-Day, All-Cause, Risk-Standardized Mortality Rate Following Pneumonia Hospitalization.</td>
<td>0468</td>
</tr>
<tr>
<td>MORT–30–STK</td>
<td>Hospital 30-Day, All-Cause, Risk-Standardized Mortality Rate Following Acute Ischemic Stroke ***.</td>
<td>N/A</td>
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### Claims-Based Coordination of Care Measures

<table>
<thead>
<tr>
<th>Short name</th>
<th>Measure name</th>
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</tr>
</thead>
<tbody>
<tr>
<td>READM–30–AMI</td>
<td>Hospital 30-Day All-Cause Risk-Standardized Readmission Rate (RSRR) Following Acute Myocardial Infarction (AMI) Hospitalization.</td>
<td>0505</td>
</tr>
<tr>
<td>READM–30–CABG</td>
<td>Hospital 30-Day, All-Cause, Unplanned, Risk-Standardized Readmission Rate (RSRR) Following Coronary Artery Bypass Graft (CABG) Surgery.</td>
<td>2515</td>
</tr>
<tr>
<td>READM–30–COPD</td>
<td>Hospital 30-Day, All-Cause, Risk-Standardized Readmission Rate (RSRR) Following Chronic Obstructive Pulmonary Disease (COPD) Hospitalization.</td>
<td>1891</td>
</tr>
<tr>
<td>READM–30–HF</td>
<td>Hospital 30-Day, All-Cause, Risk-Standardized Readmission Rate (RSRR) Following Heart Failure (HF) Hospitalization.</td>
<td>0330</td>
</tr>
<tr>
<td>READM–30–HWR</td>
<td>Hospital-Wide All-Cause Unplanned Readmission Measure (HWR)</td>
<td>1789</td>
</tr>
<tr>
<td>READM–30–PN</td>
<td>Hospital 30-Day, All-Cause, Risk-Standardized Readmission Rate (RSRR) Following Pneumonia Hospitalization.</td>
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<tr>
<td>READM–30–STK</td>
<td>30-Day Risk Standardized Readmission Rate Following Stroke Hospitalization</td>
<td>N/A</td>
</tr>
<tr>
<td>READM–30–THA/TKA</td>
<td>Hospital-Level 30-Day, All-Cause Risk-Standardized Readmission Rate (RSRR) Following Elective Primary Total Hip Arthroplasty (THA) and/or Total Knee Arthroplasty (TKA).</td>
<td>1551</td>
</tr>
<tr>
<td>AMI Excess Days</td>
<td>Excess Days in Acute Care after Hospitalization for Acute Myocardial Infarction</td>
<td>2881</td>
</tr>
<tr>
<td>HF Excess Days</td>
<td>Excess Days in Acute Care after Hospitalization for Heart Failure</td>
<td>2880</td>
</tr>
<tr>
<td>PN Excess Days</td>
<td>Excess Days in Acute Care after Hospitalization for Pneumonia</td>
<td>2882</td>
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### Claims-Based Payment Measures

<table>
<thead>
<tr>
<th>Short name</th>
<th>Measure name</th>
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</thead>
<tbody>
<tr>
<td>AMI Payment</td>
<td>Hospital-Level, Risk-Standardized Payment Associated with a 30-Day Episode-of-Care for Acute Myocardial Infarction (AMI).</td>
<td>2431</td>
</tr>
<tr>
<td>HF Payment</td>
<td>Hospital-Level, Risk-Standardized Payment Associated with a 30-Day Episode-of-Care For Heart Failure (HF).</td>
<td>2436</td>
</tr>
<tr>
<td>PN Payment</td>
<td>Hospital-Level, Risk-Standardized Payment Associated with a 30-Day Episode-of-Care For Pneumonia.</td>
<td>2579</td>
</tr>
<tr>
<td>THA/TKA Payment</td>
<td>Hospital-Level, Risk-Standardized Payment Associated with an Episode-of-Care for Primary Elective Total Hip Arthroplasty and/or Total Knee Arthroplasty.</td>
<td>N/A</td>
</tr>
<tr>
<td>MSPB</td>
<td>Payment-Standardized Medicare Spending Per Beneficiary (MSPB)</td>
<td>2158</td>
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<tr>
<td>Cellulitis Payment</td>
<td>Cellulitis Clinical Episode-Based Payment Measure</td>
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</tr>
<tr>
<td>GI Payment</td>
<td>Gastrointestinal Hemorrhage Clinical Episode-Based Payment Measure</td>
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</tr>
<tr>
<td>Kidney/UTI Payment</td>
<td>Kidney/Urinary Tract Infection Clinical Episode-Based Payment Measure</td>
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</tr>
<tr>
<td>AA Payment</td>
<td>Aortic Aneurysm Procedure Clinical Episode-Based Payment Measurement</td>
<td>N/A</td>
</tr>
<tr>
<td>Chole and CDE Payment</td>
<td>Cholecystectomy and Common Duct Exploration Clinical Episode-Based Payment Measurement</td>
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</tr>
<tr>
<td>SFusion Payment</td>
<td>Spinal Fusion Clinical Episode-Based Payment Measure</td>
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</table>

### Chart-Abstracted Clinical Process of Care Measures

<table>
<thead>
<tr>
<th>Short name</th>
<th>Measure name</th>
<th>NQF #</th>
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<tbody>
<tr>
<td>ED–1 *</td>
<td>Median Time from ED Arrival to ED Departure for Admitted ED Patients</td>
<td>0495</td>
</tr>
<tr>
<td>ED–2 *</td>
<td>Admit Decision Time to ED Departure Time for Admitted Patients</td>
<td>0497</td>
</tr>
<tr>
<td>Imm–2</td>
<td>Influenza Immunization</td>
<td>1659</td>
</tr>
<tr>
<td>PC–01 *</td>
<td>Elective Delivery</td>
<td>0469</td>
</tr>
<tr>
<td>Sepsis</td>
<td>Severe Sepsis and Septic Shock: Management Bundle (Composite Measure)</td>
<td>0500</td>
</tr>
<tr>
<td>VTE–6</td>
<td>Incidence of Potentially Preventable Venous Thromboembolism</td>
<td>+</td>
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</table>

### EHR-Based Clinical Process of Care Measures (that is, Electronic Clinical Quality Measures (eCQMs))

<table>
<thead>
<tr>
<th>Short name</th>
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<tbody>
<tr>
<td>AMI–8a</td>
<td>Primary PCI Received Within 90 Minutes of Hospital Arrival</td>
<td>+</td>
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<tr>
<td>CAC–3</td>
<td>Home Management Plan of Care Document Given to Patient/Caregiver</td>
<td>+</td>
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</table>
### PREVIOUSLY ADOPTED HOSPITAL IQR PROGRAM MEASURES FOR THE FY 2020 PAYMENT DETERMINATION AND SUBSEQUENT YEARS—Continued

<table>
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<tr>
<th>Short name</th>
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<tbody>
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<td>ED–1*</td>
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</tr>
<tr>
<td>ED–2*</td>
<td>Admit Decision Time to ED Departure Time for Admitted Patients</td>
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</tr>
<tr>
<td>EHD1-1a</td>
<td>Hearing Screening Prior to Hospital Discharge</td>
<td>1354</td>
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<tr>
<td>PC–01*</td>
<td>Elective Delivery</td>
<td>0469</td>
</tr>
<tr>
<td>PC–05</td>
<td>Exclusive Breast Milk Feeding</td>
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<tr>
<td>STK–02</td>
<td>Discharged on Antithrombotic Therapy</td>
<td>0435</td>
</tr>
<tr>
<td>STK–03</td>
<td>Anticoagulation Therapy for Atrial Fibrillation/Flutter</td>
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<tr>
<td>STK–05</td>
<td>Antithrombotic Therapy by the End of Hospital Day Two</td>
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<tr>
<td>STK–06</td>
<td>Discharged on Statin Medication</td>
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<tr>
<td>STK–08</td>
<td>Stroke Education</td>
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<tr>
<td>STK–10</td>
<td>Assessed for Rehabilitation</td>
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<td>VTE–1</td>
<td>Venous Thromboembolism Prophylaxis</td>
<td>0371</td>
</tr>
<tr>
<td>VTE–2</td>
<td>Intensive Care Unit Venous Thromboembolism Prophylaxis</td>
<td>0372</td>
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</table>

### Patient Experience of Care Survey Measures

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<thead>
<tr>
<th>Measure name</th>
<th>Measure name</th>
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</tr>
</thead>
<tbody>
<tr>
<td>HCAHPS</td>
<td>Hospital Consumer Assessment of Healthcare Providers and Systems (including Care Transition Measure (CTM-3) and Communication About Pain composite measure)</td>
<td>0166</td>
</tr>
</tbody>
</table>

### Structural Patient Safety Measures

<table>
<thead>
<tr>
<th>Measure name</th>
<th>Measure name</th>
<th>NQF #</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient Safety Culture</td>
<td>Hospital Survey on Patient Safety Culture</td>
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</tr>
<tr>
<td>Safe Surgery Checklist Use</td>
<td>Safe Surgery Checklist Use</td>
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</tr>
</tbody>
</table>

* Measure listed twice, as both chart-abstracted and electronic clinical quality measure.

** Proposed measure refinement of the HCAHPS measure’s Pain Management questions for the FY 2020 payment determination and for subsequent years, as described in section IX.A.6.a. of the preamble of this proposed rule.

*** Proposed measure refinement of the Hospital 30-Day, All-Cause, Risk-Standardized Mortality Rate Following Acute Ischemic Stroke, for the FY 2023 payment determination and for subsequent years, as described in section IX.A.6.b. of the preamble of this proposed rule.

* NQF endorsement has been removed.

7. Proposed Voluntary Hybrid Hospital-Wide Readmission Measure With Claims and Electronic Health Record Data (NQF #2879)

a. Background

In the FY 2016 IPPS/LTCH PPS final rule (80 FR 49698), we stated that we are considering the use of a set of core clinical data elements extracted from hospital EHRs for each hospitalized Medicare FFS beneficiary over the age of 65 years. The core clinical data elements are data which are routinely collected on hospitalized adults, extraction from hospital EHRs is feasible, and can be utilized as part of specific quality outcome measures. One way in which we envisioned using core clinical data elements in conjunction with other sources of data, such as administrative claims, is to calculate “hybrid” outcome measures, which are quality measures that utilize more than one source of data. For more detail about core clinical data elements, we refer readers to our discussion in the FY 2016 IPPS/LTCH PPS final rule (80 FR 49698 through 49704). In addition, we note an important distinguishing factor about core clinical data elements and the hybrid measures: hybrid measure results must be calculated by CMS to determine hospitals’ risk-adjusted rates relative to national rates used in public reporting. With a hybrid measure, hospitals can submit data extracted from the EHR, and we can perform the measure calculations. This was the approach that was finalized for the calculation of the Hybrid Hospital 30-Day, All-Cause Risk-Standardized Mortality Rate (RSMR) Following Acute Myocardial Infarction (NQF #2473), which was incorporated into the Advancing Care Coordination Through Episode Payment Models as a voluntary measure for patients admitted for AMI in the AMI Model (82 FR 354 through 356).

In the FY 2016 IPPS/LTCH PPS final rule, we stated that we developed two hybrid measures: (1) Hospital 30-Day Risk-Standardized Acute Myocardial Infarction (AMI) Mortality eMeasure (NQF #2473) (now called the Hybrid Hospital 30-Day All Cause Risk-Standardized Mortality Rate Following Acute Myocardial Infarction (AMI) (NQF #2473)); and (2) a hybrid hospital-wide 30-day readmission measure now called the Hybrid Hospital-Wide Readmission Measure with Claims and Electronic Health Record Data (NQF #2879). Although the Hybrid Hospital-Wide Readmission Measure with Claims and Electronic Health Record Data (NQF #2879) (hereinafter referred to as Hybrid HWR measure) had not been endorsed when the MAP considered the measure, it encouraged further development (80 FR 49698), and the measure has since been endorsed by the NQF.

In the FY 2016 IPPS/LTCH PPS final rule (80 FR 49702), commenters noted either outright or conditional support for the development of hybrid measures, and for the collection of additional administrative linkage variables to merge data from EHRs with claims. A few commenters noted collection of the core clinical data elements would not impose additional burden on hospitals (80 FR 49702). A few commenters recommended the hybrid measures should go through NQF review or be endorsed by NQF prior to inclusion in a quality reporting program, which we have done, as the Hybrid HWR measure was endorsed by NQF on December 9, 2016. Other commenters recommended that before we require the submission of the core clinical data elements, we should conduct further testing and analysis to ensure the accuracy and completeness of the data being submitted; specifically, one commenter suggested a testing period (80 FR 49703). We conducted further testing, 

which is further described below. We refer readers to the FY 2016 IPPS/LTCH PPS final rule (80 FR 49702 through 49704) for a full discussion of all public comments and our responses related to core clinical data elements.

Since the FY 2016 IPPS/LTCH PPS final rule, in keeping with our goal to move toward greater use of data from EHRs for quality measurement, and in response to stakeholder feedback to include clinical data in outcome measures (80 FR 49702 through 49703), we have further developed the proposed voluntary Hybrid HWR measure. This measure would incorporate a combination of claims data and EHR data submitted by hospitals, and because of these combined data sources, it is referred to as a hybrid measure. The Hybrid HWR measure cohort and outcome are identical to those in the Hospital-Wide All-Cause Unplanned Readmission measure (NQF #1789), which was adopted into the Hospital IQR Program for the FY 2015 payment determination and subsequent years (77 FR 53521).

The Hybrid HWR measure was presented on the List of Measures under Consideration for December 1, 2014. The MAP encouraged further development of the Hybrid HWR measure in December 2014. The Hybrid HWR measure (NQF #2879) was endorsed by NQF on December 9, 2016. This measure aligns with the National Quality Strategy (NQS) priorities of making care safer by reducing harm caused in the delivery of care and promoting effective communication and coordination of care.

Measure development followed the same scientific approach and rigorous process as other Hospital IQR Program outcome measures. To align the core clinical data elements with other measures that utilize EHR data, we developed and tested a Measure Authoring Tool and identified value sets for extraction of the core clinical data elements. As stated in the FY 2016 IPPS/LTCH PPS final rule, the core clinical data elements used existing value sets where possible in an effort to harmonize with other measures and reporting requirements and we completed testing of the electronic specifications for the core clinical data elements used in the Hybrid HWR measure (80 FR 49703). The electronic specifications were tested in four separate health systems that used three separate EHR systems. During Hybrid HWR measure development and testing we demonstrated that the core clinical data elements were feasibly extracted from hospital EHRs for nearly all adult patients admitted. We also demonstrated that the use of the core clinical data elements to risk-adjust the Hybrid HWR measure improves the discrimination of the measure, or the ability to distinguish patients with a low risk of readmission from those at high risk of readmission, as assessed by the c-statistic. In addition, inclusion of clinical information from patient EHRs is responsive to stakeholders who find it preferable to use clinical information that is available to the clinical care team at the time treatment is rendered to account for patients’ severity of illness rather than relying solely on data from claims (80 FR 49702). The Hybrid HWR measure is now fully developed and tested and NQF-endorsed (NQF #2879).

b. Proposal for Voluntary Reporting of Electronic Health Record Data for the Hybrid HWR Measure (NQF #2879)

In accordance with, and to the extent permitted by, the HIPAA Privacy Rule and other applicable law, we are proposing the Hybrid HWR measure as a voluntary measure for the reporting of data on discharges over a 6-month period in the first two quarters of CY 2018 (January 1, 2018 through June 30, 2018). A hospital’s annual payment determination would not be affected by this voluntary measure. As we stated when we adopted the Hospital-Wide All-Cause Unplanned Readmission measure (NQF #1789) that is currently used in the Hospital IQR Program, a hospital’s readmission rate is affected by complex and critical aspects of care such as communication between providers or between providers and patients; prevention of, and response to, complications; patient safety; and coordinated transitions to the outpatient environment, such that a hospital-wide, all-condition readmission measure could portray a broader sense of the quality of care in hospitals and promote hospital quality improvement (77 FR 53522). We believe this would also be the case with using the Hybrid HWR measure (NQF #2879) that is being proposed for voluntary data collection in this proposed rule.

Hospitals that voluntarily submit data for this measure would receive confidential hospital-specific reports that detail submission results from the performance reporting period, as well as the Hybrid HWR measure results assessed from merged files created by our merging of the EHR data elements submitted by each participating hospital with claims data from the same set of index admission. We note that in this proposal we are only seeking to collect data for the Hybrid HWR measure that are in accordance with the measure’s electronic specifications, available on the CMS Web site at: http://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment- Instru...HospitalQualityInits/Measure-Methodology.html. Hospitals that volunteer to submit data would also increase their familiarity with submitting data for hybrid quality measures from their EHR systems. Participating hospitals would receive information and instruction on the use of the electronic specifications for this measure, would have an opportunity to test extraction and submission of data to CMS, and would receive reports from CMS, downloadable from QualityNet, with details on the success of their submission, such as the completeness and accuracy of the data. This would allow us to refine this measure if necessary to provide meaningful information on outcomes for hospitalizations for Medicare FFPS beneficiaries with the intent to propose this as a required measure in future rulemaking. For example, we would consider feedback from hospitals when making refinements to improve the utility of the measure specifications. In addition, we would examine the completeness and accuracy of the data received to determine the adequacy for calculation of the measure’s risk adjustment model and measure results. EHR data or measure results for this proposed voluntary Hybrid HWR measure would not be publicly reported. However, if we propose to require mandatory reporting of the Hybrid HWR measure in future rulemaking, such a proposal would include public reporting of the measure results. Consistent with estimates for previous voluntary measure reporting, such as the Hospital IQR Program eCQM voluntary reporting (79 FR 50346), we


believe up to approximately 100 hospitals would voluntarily submit data for the Hybrid HWR measure. Details about the measure and our proposal for voluntary reporting certain data elements for this measure are discussed below.

c. Data Sources

We are proposing to use two sources of data for the calculation of the proposed voluntary Hybrid HWR measure: Medicare Part A claims and core clinical data elements for Medicare FFS beneficiaries who are 65 years or older, comprising the measure cohort. Claims data would be used to identify index admissions included in the measure cohort, to create a risk-adjustment model, and to assess the 30-day unplanned readmission outcome. This data would be merged with core clinical data elements from each participant hospital’s EHRs collected at presentation (discussed in more detail below) and used for risk-adjustment of patients’ severity of illness (for Medicare Fee-For-Service beneficiaries who are 65 years or older), in addition to data from claims. Medicare enrollment data, from the Medicare Enrollment Database, are used to confirm Medicare enrollment for at least 30 days post hospital discharge for the unplanned readmission outcome assessment.

For this proposed voluntary Hybrid HWR measure, in accordance with, and to the extent permitted by, the HIPAA Privacy Rule and other applicable law, the eHR data submission process would align as much as possible with existing electronic Clinical Quality Measure (eCQM) standards and data reporting procedures for hospitals, as further discussed below. The electronic specifications for the proposed voluntary Hybrid HWR measure, which include the electronic specifications for extraction of the core clinical data elements from hospital EHRs (the Measure Authoring Tool output and value sets) for all included data elements, are available on the CMS Web site at: http://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInitiatives/Measure-Methodology.html.

d. Outcome

As stated above, the proposed voluntary Hybrid HWR measure outcome is aligned with the currently adopted, publicly reported, Hospital-Wide All-Cause Unplanned Readmission measure (77 FR 53521). This algorithm identifies admissions that are typically planned and may occur within 30 days of discharge from the hospital. The algorithm was most recently refined in the FY 2015 IPPS/LTC PPS final rule (79 FR 50211 through 50216) for the previously adopted, claims-based measure. That same algorithm is used for this proposed voluntary Hybrid HWR measure.

A complete description of the CMS Planned Readmission Algorithm, which includes lists of planned diagnoses and procedures, can be found on the CMS Web site at: http://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInitiatives/Measure-Methodology.html.

e. Cohort

As noted above, the proposed voluntary Hybrid HWR measure cohort is aligned with the currently adopted, Hospital-Wide All-Cause Unplanned Readmission measure. The measure cohort consists of Medicare FFS beneficiaries, aged 65 years or older, discharged from non-Federal acute care hospitals. Hospitals would only submit data for this cohort, and the measure would only be calculated for this cohort.

The proposed voluntary Hybrid HWR measure includes admissions for nearly all Medicare FFS beneficiaries over the age of 65 years who are discharged alive from acute care non-federal hospitals. However, during measure calculation, a small number of these admissions are excluded under the measure specifications. Excluded admissions include those for principal discharge diagnoses indicating some psychiatric disorders. These exclusions are only a small proportion of all index admissions and are identified during the measure calculation process.

f. Inclusion and Exclusion Criteria

The proposed voluntary Hybrid HWR measure inclusion and exclusion criteria are also aligned with the currently adopted Hospital-Wide All-Cause Unplanned Readmission measure. For both measures, the index admission is the hospitalization to which the readmission outcome is attributed. Both the claims-based, Hospital-Wide All-Cause Unplanned Readmission measure and the proposed voluntary Hybrid HWR measure include the following index admissions for patients:

• Enrolled in Medicare FFS Part A for the 12 months prior to the date of admission and during the index admission.
• Aged 65 or older.
• Discharged alive from a non-Federal acute care hospital.
• Not transferred to another acute care facility.

This measure excludes the following index admissions for patients:

• Admitted to prospective payment system (PPS)-exempt cancer hospitals.
• Without at least 30 days of post-discharge enrollment in Medicare FFS.
• Discharged against medical advice.
• Admitted for primary psychiatric diagnoses.
• Admitted for rehabilitation.
• Admitted for medical treatment of cancer.

For both measures, each index admission is assigned to one of five mutually exclusive specialty cohorts: Medicine; surgery/gynecology; cardiorespiratory; cardiovascular; and neurology. The cohorts reflect how care is organized within hospitals. To assign admissions to cohorts, admissions are first screened for the presence of an eligible Agency for Healthcare Research and Quality (AHRQ) Clinical Classifications Software (CCS) surgical procedure category. Admissions with an eligible surgical procedure category are assigned...
to the surgical cohort, regardless of the principal discharge diagnosis code of the admission. All remaining admissions are assigned to cohorts based on the AHRQ CCS diagnosis category of the principal discharge diagnosis.

g. Risk-Adjustment

The proposed voluntary Hybrid HWR measure adjusts both for case mix differences (clinical status of the patient, accounted for by adjusting for age and comorbidities, and the core clinical data elements from patients’ EHRs) and service-mix differences (the types of conditions and procedures cared for and procedures conducted by the hospital, accounted for by adjusting for the discharge condition category). Patient comorbidities are based on the index admission, the admission included in the measure cohort, and a full year of prior history. The core clinical data elements are derived from information captured in the EHR during the index admission only, and are listed below.

<table>
<thead>
<tr>
<th>Data elements</th>
<th>Units of measurement</th>
<th>Time window for first captured values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart Rate</td>
<td>Beats per minute</td>
<td>0–2</td>
</tr>
<tr>
<td>Systolic Blood Pressure</td>
<td>mmHg</td>
<td>0–2</td>
</tr>
<tr>
<td>Respiratory Rate</td>
<td>Breath per minute</td>
<td>0–2</td>
</tr>
<tr>
<td>Temperature</td>
<td>Degrees Fahrenheit</td>
<td>0–2</td>
</tr>
<tr>
<td>Oxygen Saturation</td>
<td></td>
<td>0–2</td>
</tr>
<tr>
<td>Weight</td>
<td>Pounds</td>
<td>0–24</td>
</tr>
<tr>
<td>Hematocrit</td>
<td>% red blood cells</td>
<td>0–24</td>
</tr>
<tr>
<td>White Blood Cell Count</td>
<td>Cells/mL</td>
<td>0–24</td>
</tr>
<tr>
<td>Potassium</td>
<td>mEq/L</td>
<td>0–24</td>
</tr>
<tr>
<td>Sodium</td>
<td>mEq/L</td>
<td>0–24</td>
</tr>
<tr>
<td>Bicarbonate</td>
<td>mmol/L</td>
<td>0–24</td>
</tr>
<tr>
<td>Creatinine</td>
<td>mg/dl</td>
<td>0–24</td>
</tr>
<tr>
<td>Glucose</td>
<td></td>
<td>0–24</td>
</tr>
</tbody>
</table>

The risk-adjustment variables included in the development and testing of the proposed voluntary Hybrid HWR measure are derived from both claims and clinical EHR data. The variables are: (1) 13 core clinical data elements derived from hospital EHRs; (2) the Clinical Classification Software (CCS) categories for the principal discharge diagnosis associated with each index admission derived from ICD–10 codes in administrative claims data; and (3) comorbid conditions of each patient identified from inpatient claims in the 12 months prior to and including the index admission derived from ICD–10 codes and grouped into the CMS condition categories (CC).

All 13 core clinical data elements were shown to be statistically significant predictors of readmission in one or more risk-adjustment models of the five specialty cohort groups used to calculate the proposed voluntary Hybrid HWR measure. The proposed voluntary Hybrid HWR measure specialty cohort groups are further defined in section IX.A.7.e. of the preamble of this proposed rule, below. The testing results demonstrate that the core clinical data elements enhanced the discrimination (assessed using the c-statistic) when used in combination with administrative claims data. For additional details regarding the risk-adjustment model, we refer readers to the proposed voluntary Hybrid HWR Measure technical report, which is posted on the CMS Web site at: http://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInits/Measure-Methodology.html. This approach is analogous to a ratio of the hospital’s case mix and service mix, with its observed case mix and service mix, and the denominator is the number of readmissions expected based on the nation’s performance with that hospital’s case mix and service mix. This approach is analogous to a ratio of “observed” to “expected” used in other types of statistical analyses.

The methods used for calculation of the proposed voluntary Hybrid HWR measure align with the methods used to calculate the currently adopted, Hospital-Wide All-Cause Unplanned Readmission measure. Index admissions are assigned to one of five mutually exclusive specialty cohort groups consisting of related conditions or procedures. The five specialty cohort groups are: Surgery/gynecology; general medicine; cardiorespiratory; cardiovascular; and neurology. For each specialty cohort group, the standardized readmission ratio (SRR) is calculated as the ratio of the number of “expected” readmissions at a given hospital. For each hospital, the numerator of the ratio is the number of readmissions within 30 days predicted based on the hospital’s performance with its observed case mix and service mix, and the denominator is the number of readmissions expected based on the nation’s performance with that hospital’s case mix and service mix. The approach is analogous to a ratio of “observed” to “expected” used in other types of statistical analyses.


The composite SRR is multiplied by the national observed readmission rate to produce the RSRR. For additional details regarding the measure specifications to calculate the RSRR, we refer readers to the 2017 All-Cause Hospital-Wide Measure Updates and Specifications Report, which is posted on the CMS Web site at: http://cmsgov/Medicare/Quality-Initiatives-Patient-Assessment- Instruments/HospitalQualityInitis/Measure-Methodology.html.

i. Data Submission and Reporting Requirements

We are proposing that hospitals use QRDA I files for each Medicare Fee-For-Service beneficiary who is 65 years and older. Submission of data using QRDA I files is the current EHR data and measuring reporting standard adopted for electronic clinical quality measures (eCQMs) implemented in the Hospital IQR Program. This standard would be used for reporting the core clinical data elements to the CMS data receiving system. We refer readers to the FY 2016 IPPS/LTCH PPS final rule (80 FR 49706) where we have previously discussed QRDA I standards for use in the Hospital IQR Program. We also refer readers to section IX.A.10.e. of the preamble of this proposed rule for additional proposals related to data submission and reporting requirements for the Hybrid HWR measure.

We also are proposing to use the following criteria to determine if a hospital has successfully submitted voluntary Hybrid HWR measure data:

- Submission of only the first-captured values, which are data collected routinely on each Medicare FFS beneficiary who is 65 years or older upon presentation to the hospital, for each of the 13 core clinical data elements used in risk adjustment to assess the patient’s severity of illness.
- Hospitals would be expected to successfully submit data values from hospital EHRs for vital signs (heart rate, respiratory rate, temperature, systolic blood pressure, oxygen saturation, weight), and six linking variables required to merge with the CMS claims data (CCN, HIC Number or Medicare Beneficiary Identifier, date of birth, sex, admission date, and discharge date). When we tested the electronic specifications for extraction of the core clinical data elements in hospital systems, we also tested the use of these linking variables to merge data from claims and from hospitals’ EHRs from several health systems, and achieved match rates over 90 percent accounting for missing or erroneous data. In order to calculate results for the Hybrid HWR measure, hospitals would need to submit these data on more than 95 percent of on all Medicare FFS patients who are 65 years and older discharged from the hospital.
- Participating hospitals would be requested to submit values for laboratory test results (hematocrit, white blood cell count, sodium, potassium, bicarbonate, creatinine, and glucose) for Medicare FFS beneficiaries, 65 years or older, included in the measure cohort. In order to calculate measure results for the Hybrid HWR measure, hospitals would need to submit these data elements on more than 80 percent of these beneficiaries. However, for the proposed voluntary measure for the CY 2018 reporting period (January 1, 2018 through June 30, 2018) we would request the data elements on at least 50 percent of these patients discharged over the same time period. Data reporting to the CMS data receiving system would occur in the fall of 2018.
- The measurement period would include data collected continuously over a 6-month period in the first two quarters of CY 2018 (January 1, 2018 through June 30, 2018). However, for hospitals that choose to report this measure, we would request submission of these data elements on at least 50 percent of these patients. As we noted above, in our proposal for voluntary data collection of the Hybrid HWR measure, we are only seeking to collect data for this measure on applicable Medicare FFS beneficiaries in accordance with the measure’s electronic specifications, available on the CMS Web site at: http://cmsgov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInitis/Measure-Methodology.html.

j. Confidential Hospital-Specific Reports

Hospitals that voluntarily submit data for this measure would receive confidential hospital-specific reports that detail submission results from the reporting period, including detailed information about the completeness and accuracy of the EHR data they submit, as well as the Hybrid HWR measure results assessed from merged files created by our merging of the EHR data elements submitted by each participating hospital with claims data from the same set of index admission. We would calculate and provide each participating hospital with a risk-standardized readmission rate for the voluntary Hybrid HWR measure. This would provide each hospital with an indication of their performance relative to the other hospitals that participate in the voluntary measure. In addition, we would create a hospital-specific report for each participating hospital which would include detailed information about their Medicare FFS beneficiaries who are 65 and older who had an unplanned readmission within 30 days of hospital discharge, including the patients’ clinical risk factors from claims and EHR data. This information would allow hospitals to identify the factors that increase patients’ risk of readmission and would inform quality improvement strategies to reduce unplanned readmissions. In addition, the reports would include the match rate between the hospital’s submitted EHR data and corresponding claims data, as well as the proportion of patient data submitted relative to all qualifying admissions for each of the 13 core clinical data elements.

We note that we are considering proposing the Hybrid HWR (NQF #2879) measure as a required measure as early as the CY 2021 reporting period/FY 2023 payment determination and requiring hospitals to submit the core clinical data elements and linking variables used in the measure as early as CY 2020 to support a dry run of the measure during which hospitals would receive a confidential preview of their results in 2021. We would propose to require reporting on this measure in future rulemaking after we collect and analyze information from voluntary reporting.

We are inviting public comment on our proposal to adopt the Hybrid HWR measure (NQF #2879) for the Hospital IQR Program as a voluntary measure for the CY 2018 reporting period as described above.

8. Proposed Changes to Policies on Reporting of eCQMs

a. Background

For a discussion of our previously finalized eCQMs and policies, we refer readers to the FY 2014 IPPS/LTCH PPS final rule (78 FR 50807 through 50810; 50253 through 50256), the FY 2015 IPPS/LTCH PPS final rule (79 FR 50241 through 50253; 50256 through 50259; and 50273 through 50276), the FY 2016 IPPS/LTCH PPS final rule (80 FR 49692 through 49698; and 49704 through 49709), and the FY 2017 IPPS/LTCH PPS final rule (81 FR 57150 through 57161; and 57169 through 57172). In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57172), we finalized that hospitals must submit eCQM data by the end of two months following the close of the calendar year for the CY 2017 reporting period/FY 2019 payment determination and subsequent years.

In this proposed rule, we are proposing two modifications to our
finalized eCQM reporting policies for the CY 2017 reporting period/FY 2019 payment determination and the CY 2018 reporting period/FY 2020 payment determination. Specifically, we are proposing to: (1) Decrease the number of eCQMs for which hospitals must submit data; and (2) decrease the number of calendar quarters for which hospitals are required to submit data, as further detailed below. These proposals are made in conjunction with our proposals discussed in sections IX.E.2.b. of the preamble of this proposed rule to align requirements for the Hospital IQR Program and the Medicare and Medicaid EHR Incentive Programs for hospitals and CAHs.

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57150 through 57159), we finalized a policy to require hospitals to submit one full calendar year of data (consisting of four quarterly data reporting periods) for 8 self-selected eCQMs out of the available eCQMs for both the CY 2017 reporting period/FY 2019 payment determination and the CY 2018 reporting period/FY 2020 payment determination.

Since the conclusion of the public comment period for the FY 2017 IPPS/LTCH PPS final rule, we have continued to receive frequent feedback (via email, webinar questions, help desk questions, and conference call discussions) from hospitals and EHR vendors about ongoing challenges of implementing eCQM reporting. A summary of the main concerns identified by these data submitters were as follows:

- The timing of the transition to a new EHR system during 2017 (or system upgrades or new EHR vendor) affects hospitals’ ability to report on an increased number of measures in a timely manner;
- There is a need for at least one year between new EHR requirements due to the varying 6- to 24-month cycles needed for vendors to code new measures, test and institute measure updates, train hospital staff, and rollout other upgraded features;
- Hospitals have difficulty identifying applicable measures that reflect their patient population, given the reduction in the number of available eCQMs (from 28 to 15) for CY 2017 reporting and
- Hospitals have had challenges with data mapping (aligning the information available in an electronic health record (EHR), particularly if the information is not located in a structured field (for example, PDF attachment, free text section) to the required fields in a QRDA I (QRDA I file), and workflow (the process of extrapolating the pertinent patient data from an EHR, transferring that data to a QRDA I file, and submission of the QRDA I file to CMS) because hospitals still need to collect CY 2017 data while still reporting CY 2016 data.

In response to these issues, we are proposing to modify the eCQM reporting requirements for both the CY 2017 reporting period/FY 2019 payment determination and the CY 2018 reporting period/FY 2020 payment determination as discussed in more detail below:

b. Proposed Modifications to the eCQM Reporting Requirements for the Hospital IQR Program for the CY 2017 Reporting Period/FY 2019 Payment Determination

For the CY 2017 reporting period/FY 2019 payment determination, we are proposing to modify our policies to require hospitals to: (1) report on at least six of the available eCQMs, instead of eight as previously finalized, and (2) submit two self-selected quarters of data, instead of four quarters. The modified reporting cycle for the CY 2017 reporting period will be two calendar year of data as previously finalized. We believe that reducing the number of eCQMs required to be reported from eight to six and reducing the quarters of data to be reported from four quarters to any two quarters will ease the burden on data submitters, allowing them to shift resources to support system upgrades, data mapping, and staff training related to eCQMs. We also believe that the reduction in the number of required eCQMs will lessen the burden of identifying measures to report on and vendor coding of new measures; under the modified policy, hospitals will only be required to identify two additional measures between CY 2016 and CY 2017, as opposed to four additional measures. Further, successful reporting in CY 2016 should streamline CY 2017 reporting because hospitals can re-use the same measures submitted to satisfy the CY 2016 reporting requirements.

Although the publication of the FY 2018 IPPS/LTCH PPS final rule will not occur until on or about August 1, 2017, the data submission deadline for the CY 2017 reporting period/FY 2019 payment determination is not until February 28, 2018, giving hospitals ample time to adjust to these proposed modified policies. Any hospital that was prepared to submit one full calendar year of data for eight eCQMs in accordance with the previously finalized CY 2017 reporting requirements should be able to submit two self-selected quarters of data for six eCQMs in accordance with the proposed modifications to the CY 2017 reporting requirements. Reducing the number of data reporting periods to two quarters, rather than four, and allowing hospitals to select which two quarters of CY 2017 to report also will offer greater reporting flexibility and allow hospitals and their vendors more time to plan for reporting and to account for and schedule hospital-specific scenarios, such as EHR upgrades or system transitions. We believe these modified reporting requirements directly address stakeholder concerns while remaining consistent with our goal to incrementally transition to electronic reporting (80 FR 49694).

We note that we are making similar proposals in the EHR Incentive Program for eligible hospitals and CAHs to reduce confusion and reporting burden. In addition, we are not proposing any changes to the February 28, 2018 submission deadline for CY 2017 reporting (81 FR 57712) to ensure that APU determinations for FY 2019 are not affected and to maintain the established alignment with the Medicare EHR Incentive Program’s submission deadline (81 FR 57255).

We are inviting public comment on our proposals to modify the eCQM reporting requirements for the CY 2017 reporting period/FY 2019 payment determination for the Hospital IQR Program as described above.

c. Proposed Modifications to the eCQM Reporting Requirements for the Hospital IQR Program for the CY 2018 Reporting Period/FY 2020 Payment Determination

As stated above, in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57150 through 57159), we finalized a policy requiring submission of 8 self-selected eCQMs out of the available eCQMs in the Hospital IQR Program for both the CY 2017 reporting period/FY 2019 payment determination and CY 2018 reporting period/FY 2020 payment determination. In addition for the CY 2018 reporting period/FY 2020 payment determination, hospitals are required to submit the data by February 28, 2019 (the end of two months following the close of the calendar year, as set out in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57172)). For the same reasons as discussed above, we are proposing similar modifications for the CY 2018 reporting period/FY 2020 payment determination. Specifically, we are proposing to require hospitals to report on at least six of the available eCQMs for the CY 2018 reporting period/FY 2020 payment determination, instead of
eight as previously finalized. These six eCQMs may be the same or a different set of six eCQMs a hospital reports for the CY 2017 reporting period. In addition, we are proposing to decrease the number of required reporting periods, from four quarters as previously finalized, to the first three quarters of the CY 2018 reporting period (that is, Q1, Q2, and Q3 of CY 2018). We note that this differs from our proposal for the CY 2017 reporting period as discussed above, which would only require two self-selected quarters of data.

In crafting this proposal, we considered several alternatives. Specifically, we considered aligning the CY 2018 reporting period requirements with the proposed CY 2017 reporting period requirements, such that hospitals would report on at least six of the available eCQMs and submit two self-selected quarters of data for both years. We also considered retaining the reporting requirements finalized in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57153 through 57159), such that hospitals would submit one full calendar year of data for 8 self-selected eCQMs for the CY 2018 reporting period/FY 2020 payment determination. Ultimately, we believe that our proposals as stated above balance our goal to progressively shift towards electronic reporting of quality measure data with hospitals’ concerns of the burden this increase may cause. In addition, hospitals will have had several years to report data electronically for the Hospital IQR and Medicare and Medicaid EHR Incentive Programs. Therefore, we believe that hospitals will be better prepared to submit an additional quarter of data for the CY 2018 reporting period compared to the number of quarterly reporting periods we are proposing for the CY 2017 reporting period. We also believe that hospitals will be better prepared to submit additional eCQMs in the future, since hospitals will have had a sufficient number of cycles of eCQM reporting.

Our proposals for the CY 2018 reporting period/FY 2020 payment determination are being made in conjunction with proposals discussed in section IX.E.3. of the preamble of this proposed rule that fully align requirements for the Hospital IQR Program with the requirements for the CQM electronic reporting option in the Medicare EHR Incentive Program for eligible hospitals and CAHs. We note that the deadline for submission would be the same as previously finalized, two months following the end of the reporting period calendar year, specifically February 28, 2019 (81 FR 57172).

We are inviting public comment on our proposals to modify the CY 2018 reporting period/FY 2020 payment determination eCQM reporting requirements for the Hospital IQR Program as described above.

The proposed modifications to the CY 2017 reporting period/FY 2019 payment determination and CY 2018 reporting period/FY 2020 payment determination requirements, if finalized as proposed, would also have implications for eCQM validation in the Hospital IQR Program. Validation of eCQM data under the Hospital IQR Program is set to begin using FY 2017 reported data as finalized in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57153 through 57181). We refer readers to section IX.A.11. of the preamble of this proposed rule where we discuss our proposal to modify those requirements in order to align the eCQM validation process with these proposals.

9. Possible New Quality Measures and Measure Topics for Future Years

In the FY 2013 IPPS/LTCH PPS final rule (77 FR 53510 through 53512), we outlined considerations to guide us in selecting new quality measures to adopt into the Hospital IQR Program. Specifically, we seek to adopt measures for the Hospital IQR Program that would: (1) Promote better, safer, more efficient care; (2) expand the pool of measures to include measures that aim to improve patient safety; (3) support the NQS’ three-part aim of better health care for individuals, better health for populations, and lower costs for health care by creating transparency around the quality of care at inpatient hospitals to support patient decision-making and quality improvement; (4) collect data in a manner that balances the need for information related to the full spectrum of quality performance and the need to minimize the burden of data collection and reporting; (5) weigh the relevance and utility of the measures compared to the burden on hospitals in submitting data under the Hospital IQR Program; (6) to the extent practicable, consider measures that have been nationally endorsed by a multi-stakeholder organization, developed with the input of providers, purchasers/payers, and other stakeholders, and aligned with best practices among other payers and the needs of the end users of the measures; (7) in the case of a specified area or medical topic determined appropriate by the Secretary for which a feasible and practical measure has not been endorsed, give due consideration to measures that have been endorsed or adopted by a consensus organization identified by the Secretary; (8) give priority to measures that assess performance on conditions that result in the greatest mortality and morbidity in the Medicare population, are high volume and high cost for the Medicare program, and for which wide cost and treatment variations in the Medicare population have been reported across populations or geographic areas despite established clinical guidelines; (9) focus on selecting measures that will also meet the Hospital VBP Program measure inclusion criteria and advance the goals of the Hospital VBP Program by targeting hospitals’ ability to improve patient care and patient outcomes; and (10) align with the HHS Strategic Plan and Initiatives and the CMS Strategic Plan.

In keeping with these considerations, we are inviting public comment on the potential future inclusion of the following seven measures in the Hospital IQR Program (one measure related to the quality of informed consent documents, four measures that evaluate end-of-life processes and outcomes for cancer patients, and two measures that evaluate nursing skill mix):

- Quality of Informed Consent Documents for Hospital-Performed, Elective Procedures measure;
- Proportion of Patients Who Died from Cancer Receiving Chemotherapy in the Last 14 Days of Life measure (NQF #0210);
- Proportion of Patients Who Died from Cancer Not Admitted to Hospice measure (NQF #0215);
- Proportion of Patients Who Died from Cancer Admitted to the ICU in the Last 30 Days of Life measure (NQF #0213);
- Proportion of Patients Who Died from Cancer Admitted to Hospice for Less Than Three Days measure (NQF #0216);
- Skill Mix (Registered Nurse [RN], Licensed Vocational/Practical Nurse [LVN/LPN], Unlicensed Assistive Personnel [UAP], and contract) (Nursing Skill Mix) Measure (NQF #0204); and
- Nursing Hours per Patient Day Measure (NQF #0205).

We also are considering newly specified eCQMs for possible inclusion in future years of the Hospital IQR and Medicare and Medicaid EHR Incentive Programs. These measures are listed and these topics are further discussed below.

145 HHS Strategic Plan, available at: https://www.hhs.gov/about/strategic-plan/.
Safe Use of Opioids—Concurrent Prescribing; Completion of a Malnutrition Screening within 24 Hours of Admission; Completion of a Nutrition Assessment for Patients Identified as At-Risk for Malnutrition within 24 Hours of a Malnutrition Screening; Nutrition Care Plan for Patients Identified as Malnourished after a Completed Nutrition Assessment; Appropriate Documentation of a Malnutrition Diagnosis; Tobacco Use Screening (TOB–1); Tobacco Use Treatment Provided or Offered (TOB–2)/Tobacco Use Treatment (TOB–2a); Tobacco Use Treatment Provided or Offered at Discharge (TOB–3)/Tobacco Use Treatment at Discharge (TOB–3a); Alcohol Use Screening (SUB–1); Alcohol Use Brief Intervention Provided or Offered (SUB–2)/Alcohol Use Brief Intervention (SUB–2a); and Alcohol & Other Drug Use Disorder Treatment Provided or Offered at Discharge (SUB–3)/Alcohol & Other Drug Use Disorder Treatment at Discharge (SUB–3a).

a. Potential Inclusion of the Quality of Informed Consent Documents for Hospital-Performed, Elective Procedures Measure

(1) Background

The process and documentation of informed consent for surgical procedures is an ethical obligation and legal mandate intended to uphold patient autonomy. It is also a standard part of clinical practice performed prior to most procedures and therapies with material risks. This process provides information to patients about the associated risks and benefits, alternative treatment options, and what to expect during and after the procedure. As described in the literature and reported by patients, comprehensive informed consent documents can improve patient comprehension and satisfaction, and support patients in making decisions that are aligned with their expectations, preferences, and goals.147 148 149 150 151 152 153


Despite their importance, and our regulations in the Conditions for Participation Guidelines,154 informed consent documents are frequently generic, lack information that is relevant to the procedure, and include illegible, hand-written information. Moreover, patients are often given and asked to sign the informed consent document minutes before the start of a procedure when they are most vulnerable and least likely to ask questions.

Therefore, we developed the Measure of Quality of Informed Consent Documents for Hospital-Performed, Elective Procedures (hereinafter referred to as, Quality of Informed Consent Documents measure). This measure was developed in conjunction with feedback from patients and patient advocates convened by the measure developers, all of whom affirmed that the measure captured the most salient elements of informed consent documents, and represented a minimum, though significant, standard all hospitals should meet. We recognize the Quality of Informed Consent Documents measure does not capture all aspects of the informed consent process or all aspects of quality related to patient engagement in shared decision making. However, we view the Quality of Informed Consent Documents measure as a critical first step to incentivize hospitals to improve the informed consent process and to ensure patients receive basic information in a written format which is understandable, legible and presented with sufficient time allowed for questions and deliberation. The members of the patient workgroup involved in measure development also agreed with this determination and supported the measure.


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We are considering including the Quality of Informed Consent Documents measure in the Hospital IQR Program in future rulemaking.

(2) Overview of Measure

Improving the quality of informed consent documents is fundamental step for advancing patient-centered decision making.155 156 157 158 159 160 The written quality of informed consent documents is a critical component of the informed consent process, and hospitals have a role in ensuring their patients have the information they need in a readable form and with time to consider their options. We expect the Quality of Informed Consent Documents measure will help to pave the way for future measures which evaluate other components of the informed consent process, including shared decision-making.

The measure focuses on the quality of informed consent documents for elective procedures. Further, with a focus on ensuring that each person and family is engaged as partners in their care, this measure addresses the NQS priority of promoting effective communication and coordination of care. Elective procedures were chosen as the focus of the measure because all elective procedures have informed consent documents as standard practice. In addition, we believe patients undergoing elective, rather than emergent surgery, will benefit from a measure aimed at optimizing communications about the risk, benefits, and purpose of the procedure because there are typically reasonable alternatives to elective procedures and different patients may choose different options depending on their preferences.
values, and goals. Further, elective procedures usually allow ample decision time and do not require expedited explanations and decisions due to life threatening situations.

The measure would require hospitals to evaluate a sample of their informed consent documents from elective procedures performed among Medicare FFS patients aged 18 years and older hospitalized at acute care hospitals. The measure uses administrative claims to select a stratified random sample of elective procedures across specialties that are performed in hospitals. The informed consent documents associated with these procedures are reviewed and abstracted by trained personnel using a validated Abstraction Tool. Abstractors are trained using standard instructions, videos, and test documents with audit review we have developed. For additional information about the training materials and procedures, see the measure methodology report on our Web site at: http://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInitiatives/Measure-Methodology.html.

(3) Data Sources

The measure uses two sources of data to calculate the Quality of Informed Consent Documents measure: Medicare Part A administrative claims, specified below, to generate a random sample of qualified elective procedures performed at each hospital; and a sample of each hospital’s informed consent documents and the first page of the procedure/operative report for those elective procedures. Basing the sample selection on administrative data to identify medical records of elective procedures ensures a diversity of informed consent documents on a range of procedures will be reviewed, and minimizes selection bias.

(4) Outcome

The outcome for the Quality of Informed Consent Documents measure is a quality score which is calculated by aggregating the scores for individual informed consent documents from each hospital assessed with the Abstraction Tool. The items selected for inclusion in the Abstraction Tool were important to patients, supported by evidence in the literature and published standards and guidelines, applicable to the cohort of elective procedures, easily abstracted from medical records without undue burden on patients and hospitals, and feasible and reliably measured. These elements are also meaningful components of informed consent document quality from the patient perspective. Further, we received consistent feedback from all participating hospitals during testing of this measure that this information was useful for hospitals’ efforts to improve their informed consent documents and processes by identifying important gaps in existing documentation. Quality scores on each informed consent document will be aggregated to derive a hospital-level performance score.

The outcome for this measure does not overlap with our current regulations holding hospitals accountable for informed consent pursuant to our Conditions of Participation or The Joint Commission 2009 Requirements Related to the Provision of Culturally Competent Patient-Centered Care Hospital Accreditation Program (HAP), and fully aligns with State laws within the few States which have more specified informed consent rules. Current Conditions of Participation regulations focus on whether informed consent occurred and emphasize informed consent documents should include the name of the hospital, procedure, and practitioner performing the procedure along with a statement certifying the procedure, anticipated benefits, material risks, and alternative treatment options were explained to the patient or the patient’s legal representative. The Joint Commission offers additional guidance for best practices. However, there are no regulations to ensure hospitals provide patients with adequate written information about the procedure. We believe the use of this measure would supplement and augment existing standards by incentivizing hospitals to provide a minimum set of critical information about an elective procedure to the patient within a reasonable time before the patient undergoes the procedure and to enable the patient to receive and process the information prior to signing and providing informed consent.

(5) Cohort

The cohort for the Quality of Informed Consent Documents measure includes informed consent documents for a randomly selected sample of qualifying elective surgical procedures performed within non-federal acute care hospitals performed on Medicare FFS beneficiaries, aged 18 years and over who are enrolled in Part A at the time of the procedure. The list of qualifying elective procedures includes procedures for which informed consent is standard practice. The list of qualifying procedures is broad, capturing 10 specialties and various levels of invasiveness. For example, electively-performed knee replacements and coronary artery bypass surgeries are both included. For more information about the list of qualifying procedures, we refer readers to the measure methodology report on our Web site available at: http://www.cms.gov/...
(6) Inclusion and Exclusion Criteria
Qualifying electively-performed procedures were identified using the AHRQ Clinical Classification Software (CCS) codes from the list of potentially planned procedures and the list of acute discharge diagnosis AHRQ CCS codes in the CMS Planned Readmission Algorithm. The Planned Readmission Algorithm used for existing CMS readmission measures was refined in the FY 2015 IPPS/LTCFPS final rule (79 FR 50211 through 50216).

A complete description of the CMS Planned Readmission Algorithm, which includes lists of potentially planned procedures and acute discharge diagnoses, can be found on the CMS Web site at: http://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInitiatives/Measure-Methodology.html.

The CMS Planned Readmission Algorithm identifies a list of potentially planned procedures that have a potentially planned procedure without an acute discharge diagnosis code in the CMS Planned Readmission Algorithm. The Quality of Informed Consent Documentation measure does not use the Planned Readmission Algorithm to identify planned versus unplanned readmissions. The measure builds upon the established approach of the Planned Readmission Algorithm to identify only electively-performed procedures because planned procedures are also commonly electively-performed. We used clinical expert review to further narrow the list of potentially planned procedures from the Planned Readmission Algorithm to those which are consistently elective-performed and likely to have informed consent obtained prior to every procedure.

The measure excludes highly specialized procedures, such as organ transplantation because they typically use unique informed consent processes; non-invasive radiographic diagnostic tests because informed consent standards may be different than standards for invasive procedures and surgeries; and procedures that are conducted over several encounters since informed consent is likely only conducted prior to the first procedure. For more information about the list of qualifying procedures and excluded procedures, refer readers to the measure methodology report on our Web site available at: http://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInitiatives/Measure-Methodology.html.

(7) Abstraction Tool
The Abstraction Tool is an instrument used to evaluate the quality of a hospital’s informed consent documents based on a score of 0–20; a higher score indicates better quality. The Abstraction Tool is a checklist evaluating the presence of the following items in the consent document: A description of the procedure; how the procedure will be performed; the rationale for why the procedure will be performed; and the risks, benefits, and alternatives to the procedure. The Abstraction Tool also includes an item to assess whether patients received the document at least one calendar day in advance of the procedure date. Inclusion of the timing item ensures informed consent documents are not shared for the first time with patients on the day of the procedure. The abstraction tool provides an option for hospitals to note if a patient chose to opt out of signing their informed consent document 24 or more hours before surgery, enabling full credit to be given to the hospital for this item in that scenario. In addition, the tool gives credit for sharing the document prior to the day of the procedure, even if the patient does not sign the document until the day of the procedure. These aspects were raised with the patient and patient advocate workgroup and deemed to be more flexible to a range of scenarios and contexts, and therefore more patient-sensitive. To assess the reliability of the Abstraction Tool, we examined the inter-rater reliability (the degree of agreement among abstractors) of each item on the Abstraction Tool as well as the document scores produced by the Abstraction Tool for 80 of the 800 documents tested from the pilot project hospitals. For additional information about testing refer to our Web site at: http://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInitiatives/Measure-Methodology.html.

Abstractors enter responses for each item evaluated in each informed consent document. We would provide comprehensive standardized training materials including an instruction manual with guidance and examples of what meets criteria for each item in the Abstraction Tool, a training video, and sample test documents. This process has previously been piloted and found to be effective and efficient. For more information about the Abstraction Tool and instructions manual, refer readers to our Web site at: http://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInitiatives/Measure-Methodology.html.

(8) Calculating the Measure Score
The measure will be calculated by aggregating the scores of the sample of hospitals’ informed consent documents, as assessed using the Abstraction Tool. Based on input from stakeholders during the measure development stage, including the Technical Expert Panel convened by the measure developer, and feedback from patients and patient advocates, we are considering reporting the proportion of a hospital’s sampled informed consent documents that achieve a pre-specified threshold score. For example, the proportion of a hospital’s sampled informed consent documents which meet a minimum, patient-centered standard. We are considering setting the threshold score at 10 (out of 20 total points), and increasing the threshold score over time. The stakeholders we sought input from during the measure development process agreed with incrementally increasing the threshold score over time. This would establish an initial target that hospitals could feasibly meet in a short period of time, and allow for further informed consent improvement. Ultimately, we envision this measure would either evolve to include additional components or could complement a measure of shared decision making when an appropriate measure becomes available for potential use in the Hospital IQR Program.

Using this scoring approach, performance scores among the 25 hospitals in the testing sample were poor. The median hospital level score, based on evaluation of 100 informed consent documents, ranged from 0 (95 percent CI: 0–5) to 12 (95 percent CI: 10–12) out of a total of 20 points. The proportion of documents achieving a threshold score of at least 10 (out of 20 points) per hospital, ranged from 0 percent to 70 percent, demonstrating that the quality of informed consent documents varies both within and between hospitals.

(9) Implementation
We are considering two implementation approaches. One approach implements the measure in a
centralized fashion where hospitals send their sample of informed consent documents directly to CMS or to an entity contracted by us for central abstraction and measure score calculation. Another approach is local; hospitals abstract their own informed consent documents and transmit the abstraction results to CMS for measure calculation. During measure development, we worked closely with hospitals to evaluate the burden associated with each approach. The greatest burden was associated with copying and electronically sending informed consent documents, making centralized abstraction a more burdensome option for hospitals. Using a brief formal training process and materials to prepare abstractors, we found hospital abstractors can reliably abstract documents at a rate of 15–20 documents per hour or 3–4 minutes per document. The final sample size required for measure reporting has not been determined but will not exceed 100 documents and may be substantially fewer than 100 documents per hospital.

Implementation would entail identifying a hospital’s elective procedures which meet eligibility for the Quality of Informed Consent Documents measure using administrative claims data. We would then provide hospitals with a list of procedures and encounter dates selected from a hospital’s eligible elective procedures, along with the HIC number and date of birth of the patient who had the procedure in order to identify the medical record, the qualifying procedure, and the corresponding informed consent document and operative report. Hospitals would then locally evaluate the informed consent documents for these procedures using the Abstraction Tool and transmit the results of the abstraction through a secure data file transfer or similar process, such as the QualityNet Secure Portal or the External File Online Tool. We would then calculate and report the proportion of a hospital’s sample informed consent documents to us or an entity contracted by us via a secure mechanism for review and validation. The Quality of Informed Consent Documents for Hospital-Performed, Elective Procedures (MUC16–262) measure is included in a publicly available document entitled “2016–2017 Spreadsheet of Final Recommendations to HHS and CMS,” which is available on the NQF Web site. The MAP did not support this measure, indicating concern about the lack of evidence that implementation will affect hospital practices and the complexity of existing guidelines, regulations and State laws related to informed consent. Further, the MAP noted that this measure captures the quality of informed consent documents rather than the quality of communication between patients and their providers. However, the MAP noted that this measure concept is critical for shared decision making, and recommended that future measures on informed consent be patient-centered. In addition, the MAP noted that this measure should demonstrate reliability and validity, at the facility level, in the hospital setting, prior to being suitable for inclusion in the Hospital IQR Program measure set. Lastly, the MAP recommended that the measure be submitted to NQF for review and endorsement.

We are inviting public comment on multiple aspects of the measure. Specifically, we are seeking public comment on the potential scoring approach described above, reporting the proportion of a hospital’s sample informed consent documents, and setting a threshold score of 10 out of 20. In addition, we are seeking input on how the measure should be implemented, either through local abstraction where hospitals provide us with the results of their own abstraction work or by transmitting informed consent documents to us for centralized abstraction. We also are seeking public comment on the frequency of measure reporting for this measure, whether annually, quarterly or at some other interval. More frequent reporting updates would require hospitals to abstract documents and submit the results more often than less frequent reporting. Finally, we are seeking input on a potential validation process for the Quality of Informed Consent Documents measure.

b. Potential Inclusion of Four End-of-Life (EOL) Measures for Cancer Patients

(1) Background

The quality of palliative and end-of-life care has been identified as a measurement gap in the Hospital IQR Program. End-of-life care may be defined as “comprehensive care that addresses medical, emotional, spiritual, and social needs during the last stages of a person’s terminal illness.” While end-of-life care may include palliative care, palliative care is generally defined as multi-faceted, holistic care that anticipates, prevents, and alleviates suffering. Both palliative and end-of-life care can be provided when a patient is receiving hospice services, but it is not necessary for a patient to be admitted to hospice to receive such care. Hospitals are encouraged to counsel patients about palliative and end-of-life care; however, the National Academy of Medicine (NAM) of the National Academies has noted that “too few patients and families receive this help [palliative and end-of-life care] in a timely manner.” Despite evidence that this care improves patient quality of life. In the same report, the NAM proposed a number of core components of quality palliative and end-of-life care. These proposals included offering a referral to hospice if a patient “has a prognosis of 6 months or less” and regular revision of a patient’s care plan to address the patient’s changing needs, as well as the changing needs of the patient’s caregivers. The four palliative and end-of-life measures described below seek to improve the quality of care for cancer patients.

(2) Overview of Measures

All four of these end-of-life measures seek to assess the quality of end-of-life care for patients who died of cancer in order to improve the quality of end-of-life care for future cancer patients. As such, the four palliative and end-of-life measures all address the NQS priority of communication and care coordination. The Proportion of Patients Who Died from Cancer Receiving Chemotherapy in the Last 14 Days of Life (EOL-Chemo) (NQF #0210) measure evaluates the...
proportion of patients who died from cancer who received chemotherapy in the last 14 days of life. This measure was finalized for CY 2017 for the Merit Based Incentive Payment Program (MIPS) (81 FR 77672). The Proportion of Patients Who Died from Cancer Not Admitted to Hospice (EOL-Hospice) (NQF #0215) measure assesses the proportion of patients who died from cancer who were not admitted to hospice and evaluates whether or not patients were admitted to hospice. The Proportion of Patients Who Died from Cancer Admitted to Hospice for Less Than Three Days (EOL–3DH) (NQF #0216) measure evaluates whether patients who were admitted to hospice were admitted to hospice late in the course of their illness, defined as within three days of their death. The Proportion of Patients Who Died from Cancer Admitted to the ICU in the Last 30 Days of Life (EOL–ICU) (NQF #0213) measure assesses whether cancer patients were admitted to the ICU in the last 30 days of their lives. 

These measures were reviewed by the MAP in December of 2016 for the PPS-Exempt Cancer Hospital Quality Reporting (PCHQR) Program (MUC16–271, MUC16–273, MUC16–274, and MUC16–275). 173  The MAP Hospital Workgroup supported the inclusion of these measures in the PCHQR Program. Specifically, the MAP stressed the importance of end-of-life care as an area of cancer care that needs improvement and noted that these measures could help improve the patient and caregiver experience. The MAP also noted these measures could help encourage the use of hospice care and help avoid aggressive treatment in the last days of life, as unnecessary treatment at the end of life has been found to negatively impact a person’s quality of life. 174  We note that prior to implementation in the Hospital IQR Program, these measures would require a subsequent review from the MAP to assess appropriateness for programmatic inclusion.

We believe that these measures would be suitable for the Hospital IQR Program because they provide insight on the quality of end-of-life care for cancer patients provided in inpatient settings other than at PPS-exempt cancer hospitals. Currently, the Hospital IQR Program measure set does not contain any measure that assesses end-of-life care. As such, the future inclusion of these measures could promote the expansion of the Hospital IQR Program measure set to include a more robust set of measures that evaluate end-of-life care and address the NQS priority of improving person and family engagement. In addition, because these measures are specific to cancer patients, future inclusion would promote programmatic alignment between the Hospital IQR and PCHQR Programs should these measures be finalized as proposed in section IX.B.4.b. of the preamble of this proposed rule for inclusion in the PCHQR Program. Additional information on these measures is available at: http://www.qualityforum.org/Publications/2016/12/Palliative_and_End_of-Life_Care_2015-2016.aspx.

We are inviting public comment on the possible future inclusion of one or more of these end-of-life measures in the Hospital IQR Program.

c. Potential Inclusion of Two Nurse Staffing Measures

(1) Background

Nursing care is a core service of hospitals, and accordingly, hospital nurse staffing practices are increasingly recognized as a tool to improve the quality and value of care. 175  Studies have shown there is a link between nurse staffing and care quality and patient outcomes. For example, the AHRQ conducted a systematic review and meta-analysis examining the relationship between nurse staffing and patient outcomes. The review of 96 studies, published between 1990 and 2006, found that increased nurse staffing is associated with a reduction in hospital-related mortality and adverse patient events, such as respiratory failure, cardiac arrest, and hospital-acquired conditions. 176  A review of studies examining the impact of nurse staffing on hospital costs and patient length of stay found that an increased level of registered nurse (RN) staffing may result in reduced patient length of stay and hospital costs. 177  Further, more recent literature has demonstrated that nursing skill mix (licensure level) and increased RN nursing hours are associated with decreased rates of


their data directly shared with CMS while those that do not currently participate in that program have the opportunity to send their data directly to CMS. In addition, the MAP noted that, at the time, there was no gold standard for these measures, and thus it is difficult to access relative performance on these measures. The final recommendation from that review was to conditionally support the inclusion of these measures, contingent upon review and endorsement from the NQF. We note that these measures, initially obtained NQF endorsement on August 5, 2009, and after subsequent review by NQF for aggregation at the hospital level, the measures retained their endorsement as of December 10, 2015. Further, we note that approximately half of hospitals are already reporting this information to the NDNQI, founded by the American Nurses Association (ANA). NDNQI data are not publicly reported.

(2) Skill Mix (Registered Nurse [RN], Licensed Vocational/Practical Nurse [LVN/LPN], Unlicensed Assistive Personnel [UAP], and Contract) (Nursing Skill Mix) Measure (NQF #0204)

(a) Overview of Measure

The NQF-endorsed Nursing Skill Mix measure assesses the percentage of productive nursing care hours worked by nursing staff with direct patient care responsibilities for each nursing licensure category (RN, LPN/LVN, and UAP) and staff employment status (contract/agency versus employee), by eligible hospital unit. The intent of this measure is to enable hospitals to track and assess their nursing skill mix, given that research demonstrates a relationship between skill mix and certain quality outcomes. The measure focuses on the structure of care quality and includes the skill mix for adult and pediatric medical-surgical hospital units. Medical-surgical hospital units include hospitals areas of care related, such as the following:

- Medication administration.
- Nursing treatments.
- Nursing rounds.
- Admission, transfer, and discharge activities.
- Patient education.
- Patient communication.
- Coordination of patient care.
- Documentation time.
- Treatment planning.
- Patient screening and assessment.

Unlicensed assistive personnel (UAP) are defined as individuals trained to function in an assistive role to nursing in the provision of patient care, as delegated by and under the supervision of a registered nurse. UAPs include nursing assistants, patient care technicians/assistants, and graduate nurses not yet licensed who have completed orientations.

The measure includes: All nursing staff employed by the hospital; temporary staff who are not employed by the hospital (contract or agency); and float staff who are hospital employees temporarily assigned to provide direct patient care on an eligible unit other than their usual unit of employment.

(b) Data Source

Data collection for this structural measure would occur quarterly for each eligible unit from January 1 through December 31 of each calendar year, with data submission occurring 4.5 months after the end of each reporting quarter.

For staff with direct patient care responsibilities, the measure assesses the percentage of total productive nursing hours worked by either employee or contract RNs, LPNs/LVNs, and UAPs, as well as at the percentage of total productive nursing hours worked for contract or agency staff. Accordingly, four rates (percentages) are determined for each eligible hospital unit, one for each type of nursing staff, and one for contract and agency nursing staff. The four separate rates are as follows:

1. RN hours—Productive nursing care hours worked by RNs (employee and contract) with direct patient care responsibilities for each eligible inpatient unit/the total number of productive hours worked by employee or contract nursing staff with direct patient care responsibilities (RN, LPN/LVN, and UAP) for each eligible inpatient unit;
2. LPN/LVN hours—Productive nursing care hours worked by LPNs/LVNs (employee and contract) with direct patient care responsibilities for each eligible inpatient unit/the total number of productive hours worked by employee or contract nursing staff with direct patient care responsibilities (RN, LPN/LVN, and UAP) for each eligible inpatient unit;
3. UAP hours—Productive nursing care hours worked by UAPs (employee and contract) with direct patient care responsibilities for each eligible inpatient unit/the total number of productive hours worked by employee or contract nursing staff with direct patient care responsibilities (RN, LPN/LVN, and UAP) for each eligible inpatient unit; and
4. Contract or agency hours—Productive nursing care hours worked by contract or agency staff with direct patient care responsibilities (RN, LPN/LVN, and UAP) with direct patient care responsibilities for each eligible inpatient unit/the total number of productive hours worked by employee or contract nursing staff with direct patient care responsibilities (RN, LPN/LVN, and UAP) for each eligible inpatient unit.

The data collected and the rates calculated are aggregate nursing care hours worked by each licensure category, by unit type. Hospital rates are weighted for patient volume (patient days) to account for differences in unit sizes.
(d) Cohort

Employee, contract, or agency RNs, LPN/LVNs, and UAPs with direct patient care responsibilities are included in the numerator and denominator statements. The measure numerator and denominator include nursing staff assigned to the eligible unit who have direct patient care responsibilities for greater than 50 percent of their shift who are counted in an eligible unit’s staffing matrix, are replaced if they call in sick, and whose work hours are charged to the unit’s cost center. The measure numerator and denominator exclude the following: Nursing staff with no direct patient care responsibilities whose primary responsibility is administrative in nature; specialty teams (for example, wound care), patient educators, or case managers who are not assigned to a specific unit; unit clerks, monitor technicians, and secretaries with no direct patient care responsibilities; sitter not providing routine UAP activities; therapy assistants; student nurses fulfilling educational requirements; and nursing staff undergoing orientation who are not included in the eligible units staffing matrix. For more information regarding the Nursing Skill Mix measure, we refer readers to the NQF measure information page available at: http://www.qualityforum.org/QPS/0204.

We are inviting public comment on the future inclusion of the Skill Mix (Registered Nurse [RN], Licensed Vocational/Practical Nurse [LVN/LPN], Unlicensed Assistive Personnel [UAP], and Contract) (Nursing Skill Mix) measure for the Hospital IQR Program. Specifically, we are seeking public comments on narrowing the number of hospital units included in the measures’ calculation, which units we should consider for inclusion, and the burden of data collection on hospitals.

(3) Nursing Hours per Patient Day Measure (NQF #0203)

(a) Overview of Measure

The NQF-endorsed Nursing Hours per Patient Day for eligible adult and pediatric medical-surgical inpatient hospital units. Medical-surgical hospital units include hospitals areas for the evaluation of patients with medical and/or surgical conditions. Eligible adult and pediatric medical-surgical units can be mapped to the CDC’s National Healthcare Safety Network (NHSN) Healthcare Service locations codes as defined in the NHSN Patient Safety Component Manual. Similar to the Nursing Skill Mix Measure, additional unit types, such as adult and pediatric critical-care, step-down, medical, and surgical units could be included, but at this time, we believe that limiting the measure to adult and pediatric medical-surgical units would allow hospitals to become accustomed to collecting and reporting staffing data while also providing important staffing information to consumers. However, we are seeking comment on how many inpatient units to include and which units should be prioritized.

Productive hours are defined as the hours worked by nursing staff (RN, LPN/LVN, and UAP) with direct patient care responsibilities, including overtime, not budgeted, or scheduled hours. Direct patient care responsibilities are nursing activities performed by unit-based staff in the presence of the patients and activities that occur away from the patient that are patient related, such as the following:

- Medication administration
- Nursing treatments
- Nursing rounds
- Admission, transfer, and discharge activities
- Patient education
- Patient communication
- Coordination of patient care
- Documentation time
- Treatment planning
- Patient screening and assessment

UAPs are individuals trained to function in an assistive role to nursing staff in the provision of patient care, as delegated by and under the supervision of a registered nurse. UAPs include nursing assistants, patient care technicians/assistants, and graduate nurses not yet licensed who have completed orientations.

The measure includes all nursing staff employed by the hospital; temporary staff who are not employed by the hospital (contract or agency); and float staff who are hospital employees temporarily assigned to provide direct patient care on an eligible unit other than their usual unit of employment.

(b) Data Source

Data collection for this structural measure for hospitals occur quarterly, for each eligible unit, from January 1 through December 31 of each calendar year, with data submission occurring 4.5 months after the end of each reporting quarter. These data would be collected via a web-based tool available on the QualityNet Web site.

(c) Measure Calculation

For staff with direct patient care responsibilities, the measure assesses the number of productive hours per patient day worked by both RNs and by total nursing staff (RNs, LPN/LVNs, and UAPs). Accordingly, two rates are determined for each eligible hospital unit. The two separate rates are as follows: (1) RN hours per patient day—Total number of productive hours worked by RN nursing staff (contract and employee) with direct patient care responsibilities for each eligible inpatient unit/total number of patient days for each eligible inpatient unit; and (2) Total nursing care hours per patient day—Total number of productive hours worked by RN, LPN/LVN, and UAP nursing staff (contract and employee) with direct patient care responsibilities for each eligible inpatient unit/total number of patient days for each eligible inpatient unit. Patient days must be from the same unit in which nursing care hours are reported. The data collected and the rates calculated are aggregate nursing hours per patient day, by unit type. Hospital rates are weighted for patient volume (patient days) to account for differences in unit sizes.

(d) Cohort

RNs, LPN/LVNs, and UAPs with direct patient care responsibilities are included in the numerator and denominator statement. The measure numerator includes nursing staff assigned to the eligible inpatient unit who have direct patient care responsibilities for greater than 50 percent of their shift, who are counted in an eligible unit’s staffing matrix, are replaced if they call in sick, and whose work hours are charged to the unit’s cost center. The numerator excludes the following: Nursing staff with no direct patient care responsibilities whose primary responsibility is administrative in nature; specialty teams (for example, wound care), patient educators, or case managers who are not assigned to a specific unit; unit clerks, monitor technicians, and secretaries with no direct patient care responsibilities; sitters not providing routine UAP activities; therapy assistants; student
nurses fulfilling educational requirements; and nursing staff undergoing orientation who are not included in the eligible units staffing matrix. The measure denominator excludes patient days from ineligible units. For more information regarding the Nursing Hours Per Day measure, we refer readers to the National Quality Forum measure information page available at: http://www.qualityforum.org/QPS/0205.

We are inviting public comment on the possible future inclusion of the Nursing Hours Per Day measure for the Hospital IQR Program. Specifically, we are seeking comments on narrowing the number of hospital units included in the measures’ calculation, which units we should consider for inclusion, and the burden of data collection on hospitals.

d. Potential Inclusion of Additional Electronic Clinical Quality Measures (eCQMs) in the Hospital IQR and Medicare and Medicaid EHR Incentive Programs

As we previously indicated in the FY 2013 IPPS/LTCH PPS final rule, EHR technology continues to evolve and additional infrastructure is being put in place to afford us the capacity to accept enhanced electronic reporting of many of the clinical chart-abstracted measures that are currently part of the Hospital IQR Program (77 FR 53534). We continue to believe that electronic reporting of quality measure data derived from the EHR will, in the long run, reduce the burden on hospitals to collect and submit data for the Hospital IQR Program.

In keeping with this goal, we are soliciting feedback on the potential inclusion of additional eCQMs in the Hospital IQR and Medicare and Medicaid EHR Incentive Programs. These measures assess opioid prescribing practices, malnutrition, tobacco use, and substance use among the adult, inpatient population. As we continue to make strides with electronic reporting, we want to ensure that we provide hospitals with a robust selection of eCQMs. As we state in section IX, help to fill this identified gap area. The table below lists the eCQMs being considered for future inclusion in the Hospital IQR and Medicare and Medicaid EHR Incentive Programs and for which we are seeking public feedback.

**Electronic Clinical Quality Measures (eCQMs) for Future Consideration in the Hospital IQR and Medicare and Medicaid EHR Incentive Programs**

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<td>Completion of a Malnutrition Screening within 24 Hours of Admission</td>
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<tr>
<td>Completion of a Nutrition Assessment for Patients Identified At-Risk for Malnutrition within 24 Hours of a Malnutrition Screening</td>
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(1) Safe Use of Opioids-Concurrent Prescribing Measure

(a) Background

Unintended opioid overdose fatalities have reached epidemic proportions in the last 20 years and are a major public health concern in the United States. 188 Reducing the number of unintended opioid overdoses has become a priority for numerous HHS agencies. Concurrent prescriptions of opioids or opioids and benzodiazepines put patients at greater risk of unintended opioid overdose due to increased risk of respiratory depression. 189 190 Despite this risk, studies of multiple claims and prescription databases have shown that between 5 to 15 percent of patients receive concurrent opioid prescriptions, and 5 to 20 percent of patients receive concurrent opioid and benzodiazepine prescriptions across various settings. 191 192 193 In addition, an analysis


of more than 1 million hospital admissions in the United States found that over 43 percent of all patients with nonsurgical admissions were exposed to multiple opioids during their hospitalization.

(b) Overview of Measure

The Safe Use of Opioids—Concurrent Prescribing (MUC16–167) measure assesses patients (excluding cancer patients or patients receiving palliative care), ages 18 years and older with active, concurrent prescriptions for opioids, or opioids and benzodiazepines, at discharge.

This measure addresses the following NQS priorities: (1) Making care safer by reducing harm caused in the delivery of care; (2) promoting effective communication and coordination of care; and (3) promoting the most effective prevention and treatment practices for the leading causes of mortality, starting with cardiovascular disease.

This measure was reviewed by the MAP in December 2016 and received the recommendation to refine and resubmit for consideration for programmatic inclusion. MAP stakeholders acknowledged the significant health risks associated with concurrent prescribing of opioids, and opioids and benzodiazepines, but expressed concern with the measure specifications, indicating the need for a stronger evidence base for clinical guidelines and refinement of the measure exclusions to reduce the risk of unintended consequences.

Additional information on this measure can be found in the 2016 Measures Under Consideration Spreadsheet, available at: http://www.qualityforum.org/ProjectMaterials.aspx?projectID=75367

We are inviting public comment on the possible future inclusion of this opioid prescribing measure in the Hospital IQR Program.

(2) Malnutrition Measures

(a) Background

Malnutrition is associated with many adverse outcomes including depression of the immune system, impaired wound healing, muscle wasting, and increased mortality. Patients who are malnourished during a hospital stay have an increased risk of complications, readmissions, and length of stay. In addition, evidence demonstrates an association between malnutrition risk and increased inpatient costs. One study found that patients identified with undernutrition risk and high undernutrition risk experience increased costs by 28.8 percent and 21.1 percent, respectively, when compared to non-malnourished patients. Malnutrition risk screening, using a validated screening tool, can be useful in predicting certain patient outcomes including length of stay, mortality, and post-operative complications.

Nutrition assessments for patients identified as at-risk for malnutrition have been associated with improved patient outcomes including less weight loss, reduced length of stay, improved muscle function, better nutritional intake, and fewer readmissions. Further, there is evidence of a performance gap with regard to nutrition screening and assessment. A national survey of hospital-based professionals in the United States focused on nutrition screening and assessment practices demonstrated that out of 1,777 unique respondents, only 36.7 percent reported completing nutrition screening at admission and 50.8 percent reported doing so within 24 hours. Thus, there is an opportunity for hospitals to improve nutrition screening and assessment.

(b) Overview of Measures

The malnutrition measure set consists of the following four measures:

- Completion of a Malnutrition Screening within 24 Hours of Admission (MUC16–294);
- Completion of a Nutrition Assessment for Patients Identified as At-Risk for Malnutrition within 24 Hours of a Malnutrition Screening (MUC16–296);
- Appropriate Documentation of a Nutrition Diagnosis (MUC16–344); and
- Nutrition Care Plan for Patients Identified as Malnourished after a Completed Nutrition Assessment (MUC16–372).

These malnutrition measures are new eCQMs that collectively evaluate the quality of care rendered to adult patients that are identified as malnourished. These measures address the NQS priorities of: (1) Making care safer by reducing harm caused in the delivery of care; and (2) promoting effective communication and coordination of care. The Completion of a Malnutrition Screening within 24 Hours of Admission measure (MUC16–294) assesses whether patients age 18 years or older are screened for malnutrition within 24 hours of admission to the hospital. The Completion of a Nutrition Assessment for Patients Identified as At-Risk for Malnutrition measure (MUC16–296) assesses whether patients age 65 years or older, who screen positive for being at-risk for malnutrition, have a nutrition assessment documented in the medical record within 24 hours of the most recent malnutrition screening. The Appropriate Documentation of a Malnutrition Diagnosis measure (MUC16–344) assesses whether patients age 65 years and older, who are found to be malnourished on the nutrition assessment, have adequate documentation of a malnutrition diagnosis in their medical record. This measure is important because there is often a disconnect between screening for malnutrition and documentation of a diagnosis of malnutrition, which is necessary for appropriate follow-up after hospital discharge. Data analyzed from the Healthcare Cost and Utilization Project (HCUP), a nationally-representative data set describing U.S. hospital discharges, indicated that approximately 3.2 percent of hospital discharges in 2010 included malnutrition as a diagnosis. However, this same research article notes that the prevalence of a malnutrition diagnosis may be significantly higher as past...
researchers, using validated screening tools, indicate a significantly higher prevalence of undiagnosed malnutrition in the hospital, ranging from 33 to 54 percent.204 Lastly, the Nutrition Care Plan for Patients Identified as Malnourished after a Completed Nutrition Assessment measure (MUC16–372) assesses whether patients age 65 years and older, who are found to be malnourished on a completed nutrition assessment, have a nutrition care plan documented in their medical record.

These measures were reviewed by the MAP in December 2016 and received mixed support. The Nutrition Care Plan for Patients Identified as Malnourished after a Completed Nutrition Assessment (MUC16–372), Completion of a Malnutrition Screening within 24 Hours of Admission (MUC16–294), and Completion of a Nutrition Assessment for Patients Identified as At-Risk for Malnutrition within 24 Hours of a Malnutrition Screening (MUC16–296) measures were recommended to be refined and resubmitted for consideration for programmatic inclusion. For these three measures, the MAP encouraged providing more evidence to prove clinical importance and recommended that the exclusions continue to be tested for validity.204 The Appropriate Documentation of a Malnutrition Diagnosis measure (MUC16–344) was not supported because there was concern that there was insufficient evidence to support the link between documenting a malnutrition diagnosis and improved patient outcomes.

The MAP concluded that completing a malnutrition assessment provided the most potential value to the measure set and quality of care. The MAP also encouraged the measure developer to test the individual malnutrition measures as a composite in an effort to balance the number of measures in the Hospital IQR Program with the need to fill the measure gap addressing malnutrition.205 We note that we received written support (formal letters addressed to CMS) of these measures from other stakeholders who noted that addressing malnutrition among beneficiaries is an important clinical issue.

Additional information on these measures is available at: http://www.qualityforum.org/ProjectMeasures.aspx?ProjectID=80741.

We are inviting public comment on the possible future inclusion of one or more of these malnutrition measures in the Hospital IQR Program. In addition, we are inviting public comment on the possible future inclusion of a composite measure comprised of all or a subset of these individual malnutrition measures in the Hospital IQR Program. (3) Tobacco Use Measures

(a) Background

Tobacco use is the single greatest cause of disease in the United States today and accounts for more than 480,000 deaths each year.206 Tobacco use creates a heavy cost to society as well as to individuals. Smoking is a known cause of multiple cancers, heart disease, stroke, complications of pregnancy, chronic obstructive pulmonary disease, other respiratory problems, poorer wound healing, and many other diseases.207 Smoking-attributable health care expenditures are estimated to cost at least $130 billion per year in direct medical expenses for adults and over $150 billion in lost productivity.208 There is strong and consistent evidence that tobacco dependence interventions, if delivered in a timely and effective manner, significantly reduce the user’s risk of suffering from tobacco-related disease and improve outcomes for those already suffering from a tobacco-related disease.209 210 211 212 Effective, evidence-based tobacco dependence interventions have been clearly identified and include brief clinician advice, individual, group, or telephone counseling, and use of FDA-approved medications. Tobacco cessation treatments are clinically effective and extremely cost-effective relative to other commonly used disease prevention interventions and medical treatments.213

Performance on the chart-abstracted versions of these measures, as reported by The Joint Commission, yields that the Tobacco Use Screening (TOB–1) measure had a screening rate of 98.15 percent, based on a reporting period of July 2015–June 2016.214 TOB–1 is necessary to operationalize Tobacco Use Treatment Provided or Offered (TOB–2)/Tobacco Use Treatment (TOB–2a) and Tobacco Use Treatment Provided or Offered at Discharge (TOB–3)/Tobacco Use Treatment at Discharge (TOB–3a) measures. The goal of TOB–1 is to achieve 100 percent screening so that all tobacco users are consistently identified and offered appropriate interventions, which are evaluated by TOB–2/a and TOB–3/a. As noted in the table 215 below, the performance rates for the chart-abstracted versions of TOB–2/a and TOB–3/a measures suggest that there is an opportunity for hospitals to improve tobacco use treatment during the hospital stay and at discharge.

### Tobacco Use Measures Screening Results July 2015–June 2016

<table>
<thead>
<tr>
<th>Measure name</th>
<th>Screening rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tobacco Use Treatment Provided or Offered (TOB–2)</td>
<td>66.41</td>
</tr>
<tr>
<td>Tobacco Use Treatment (TOB–2a)</td>
<td>32.97</td>
</tr>
<tr>
<td>Tobacco Use Treatment Provided or Offered at Discharge (TOB–3)</td>
<td>46.20</td>
</tr>
<tr>
<td>Tobacco Use Treatment at Discharge (TOB–3a)</td>
<td>10.71</td>
</tr>
</tbody>
</table>

(b) Overview of Measures

The tobacco use measure set consists of the following three measures:

- Tobacco Use Screening (TOB–1) (MUC16–50):


208 Ibid.


210 Baumsteiger SE., Schumann A, Meyer C, et al. Effects of smoking cessation on health care use: is smoking attributable health care expenditures are estimated to cost at least $130 billion per year in direct medical expenses for adults and over $150 billion in lost productivity.


213 Ibid.

214 Joint Commission Quality Check Data, available at: https://www.qualitycheck.org/. (Data download.)

215 Joint Commission Quality Check Data available at: https://www.qualitycheck.org/.
• Tobacco Use Treatment Provided or Offered (TOB–2)/Tobacco Use Treatment (TOB–2a) (MUC16–51); and
• Tobacco Use Treatment Provided or Offered at Discharge (TOB–3)/Tobacco Use Treatment at Discharge (TOB–3a) (MUC16–52).

The TOB measures are eCQMs that assess tobacco use screening and treatment for patients age 18 years or older during the hospital stay and at discharge. We note that these measures were derived from the chart-abstracted version by The Joint Commission. The Joint Commission has been using the chart-abstracted versions of these measures for voluntary reporting since January 1, 2012. In addition, the chart-abstracted versions of these measures (TOB–1, TOB–2/TOB–2a, and TOB–3/TOB–3a) are also part of the IPFQR Program measure set (81 FR 57246). These measures address the NQS priority of promoting the most effective prevention and treatment practices for the leading causes of mortality.

TOB–1 assesses the proportion of hospitalized patients who are screened, or refuse screening, within the three days prior to admission through 1 day after admission, for tobacco use during the 30 days prior to the screening. TOB–2 assesses the proportion of patients who are light tobacco users who received or refused practical counseling to quit within 3 days prior to or anytime during admission. TOB–2 also assesses the proportion of heavy tobacco users who received or refused practical counseling to quit and received, had a medical reason not to receive, or refused FDA-approved cessation medications within 3 days prior to or anytime during admission. The subset measure TOB–2a only assesses light tobacco users who received practical counseling to quit within 3 days prior to or anytime during admission, and heavy tobacco users who received practical counseling to quit and received, or had a medical reason not to receive, FDA-approved cessation medications within 3 days prior to or anytime during admission. TOB–3 assesses the proportion of patients who are light tobacco users who were referred to or refused counseling within 3 days prior to admission through 1 day after discharge. TOB–3 also assesses the proportions of heavy tobacco users who were referred to or refused evidence-based counseling and received, had a medical reason not to receive, or refused a prescription for FDA-approved cessation medication upon discharge. The subset measure TOB–3a assesses light tobacco users who were referred to counseling within 3 days prior to admission through one day after discharge, and heavy tobacco users who were referred to evidence-based counseling and received, or had a medical reason not to receive, a prescription for FDA-approved cessation medication upon discharge. We note that we previously solicited comments on the future inclusion of electronically-specified versions of the tobacco use measures TOB–1, TOB–2/2a and TOB–3/3a, previously referred to as TAM–1, TAM–2, and TAM–3, respectively, in the FY 2013 IPPS/LTC PPS final rule (77 FR 53535).

Commenters equally supported and opposed the future inclusion of the tobacco use measures in the Hospital IQR Program. Commenters highlighted the importance of high validation rates such as 95 percent, across the electronic data capture method and manual chart- abstraction (77 FR 53535). We note that at the time we sought public comments on these measure concepts related to tobacco use, electronically-specified measures were not yet developed.

In the most recent MAP deliberations in December 2016, only the Tobacco Use Screening (TOB–1) eCQM (MUC16–50) was reviewed. The TOB–2/TOB–2a (MUC16–51) and TOB–3/TOB–3a (MUC16–52) eCQMs were on the December 2016 MUC List, but were not submitted for MAP review because they were still undergoing field testing. We anticipate that these measures should be ready for review by the MAP in the winter of CY 2017.

The TOB–1 eCQM was recommended to be refined and resubmitted for consideration for programmatic inclusion. The MAP indicated that the measure should be tested to ensure that it returns accurate, reliable results. In addition, the MAP Hospital Workgroup noted that it will be important to carefully assess feasibility and burden of data collection. As previously stated, the chart-abstracted versions of the Tobacco Use Screening measures (TOB–1, TOB–2/TOB–2a, and TOB–3/TOB–3a) are part of the IPFQR Program measure set (81 FR 57246); thus, future inclusion of the eCQM versions of these measures in the Hospital IQR Program measure set would promote programmatic alignment across these quality reporting programs.

Additional information on the chart-abstracted version of these measures is available at: http://www.qualityforum.org/docs/ContentServer?c=Page&pageName=QnetPublic%2FPage%2FQnetTier3&cid=12287757490207.

We are inviting public comment on the possible future inclusion of one or more of the eCQM versions of these tobacco use measures (TOB–1, TOB–2/2a and TOB–3/3a) in the Hospital IQR Program. In addition, we are inviting public comment on the possible future inclusion of a composite measure comprised of all or a subset of these individual tobacco use measures in the Hospital IQR Program.

(4) Substance Use Measures
(a) Background

Excessive alcohol consumption and drug misuse or abuse have a significant impact on the health of the U.S. population. Excessive alcohol consumption is a leading cause of preventable death and disability, resulting in approximately 88,000 deaths per year with an estimated economic cost of $249 billion, including $28 billion (2010 dollars) in direct health care costs. In 2015, approximately 20.8 million individuals were classified as having a substance use disorder. Of those individuals with substance use disorders, 13.1 million had an alcohol use disorder, 5.1 million had an illicit drug use disorder, and 2.7 million had an alcohol and illicit drug use disorder. Excessive alcohol consumption and substance use disorders can increase the risk of preventable injury, vision loss, chronic diseases, such as mental illness, and lead to the development of diseases, such as heart disease, cancer, and liver
disease. Studies show the majority of individuals who consume alcohol excessively do not meet the clinical criteria for diagnosis of a substance use disorder; yet evidence demonstrates screening and brief interventions, especially prior to the onset of a substance use disorder, can improve health and reduce costs. Similar benefits have been observed for individuals with substance use disorders who are identified and referred to treatment. The table below provides performance rates based on the July 2015–June 2016 reporting period for the chart-abstracted versions of these measures, as reported by The Joint Commission. The results show that there is an opportunity for hospitals to improve substance use screening, brief intervention, and treatment.

**SUBSTANCE USE MEASURES SCREENING RESULTS JULY 2015–JUNE 2016**

<table>
<thead>
<tr>
<th>Measure name</th>
<th>Screening rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alcohol Use Screening (SUB–1)</td>
<td>85.30</td>
</tr>
<tr>
<td>Alcohol Use Brief Intervention Provided or Offered (SUB–2)</td>
<td>62.68</td>
</tr>
<tr>
<td>Alcohol Use Brief Intervention (SUB–2a)</td>
<td>57.43</td>
</tr>
<tr>
<td>Alcohol &amp; Other Drug Use Disorder Treatment Provided or Offered at Discharge (SUB–3)</td>
<td>65.46</td>
</tr>
<tr>
<td>Alcohol &amp; Other Drug Use Disorder Treatment at Discharge (SUB–3a)</td>
<td>54.27</td>
</tr>
</tbody>
</table>

(b) Overview of Measures

The substance use measure set consists of the following three measures:

- Alcohol Use Screening (SUB–1) (MUC16–179);
- Alcohol Use Brief Intervention Provided or Offered (SUB–2)/Alcohol Use Brief Intervention (SUB–2a) (MUC16–178); and
- Alcohol & Other Drug Use Disorder Treatment Provided or Offered at Discharge (SUB–3)/Alcohol & Other Drug Use Disorder Treatment at Discharge (SUB–3a) (MUC16–180).

The SUB–1, SUB–2/2a and SUB–3/3a measures assess whether hospital patients age 18 years of age and older are screened for alcohol use using a validated screening questionnaire for excessive drinking during their inpatient stay. A validated screening questionnaire is defined as an instrument that has been psychometrically tested for reliability (the ability of the instrument to produce consistent results), validity (the ability of the instrument to produce true results), and sensitivity (the probability of correctly identifying a patient with the condition).

As previously noted, these measures are intended to be implemented as a set. At such, it would be necessary to adopt the SUB–1 measure in order to implement the other two measures. The SUB–2/2a measure assesses whether hospital patients age 18 years of age or older who screened positive for excessive alcohol use or an alcohol use disorder receive or refuse a brief intervention during the hospital stay (SUB–2). Subset measure SUB–2a includes only those patients who receive a brief intervention. A brief intervention is defined as a single session or multiple sessions conducted by a qualified healthcare professional or trained peer support person, which includes motivational discussion focused on increasing patient insight and awareness regarding alcohol use and motivating behavioral change. The SUB–3/3a measures assess whether hospitals patients 18 years of age or older with a substance use disorder (alcohol or drug) receive or refuse at discharge a medication prescription for treatment or receive or refuse a referral for substance use disorder treatment (SUB–3). Subset measure SUB–3a includes only those patients who receive a medication prescription or treatment referral at discharge.

The chart-abstracted versions of these three measures, not the eCQM versions, were added to the MUC List in the summer of 2016 and reviewed by the MAP in December 2016 as discussed in the MAP Pre-Rulemaking Report and Spreadsheet entitled “2016–2017 Spreadsheet of Final Recommendations to HHS and CMS.” The MAP noted that the measure encourages hospitals to screen patients for excessive alcohol use and can prevent life-threatening alcohol withdrawal syndrome, but recommended that the measure be paired with an appropriate intervention and follow-up measure. The MAP did not support the SUB–2/2a measure (MUC16–179) for adoption into the Hospital IQR Program. Proponents of the SUB–2/2a measure supported the incorporation of behavioral health measures into the Hospital IQR Program and noted that hospitalization is a prime opportunity to discuss harmul substance use because patients may be more amenable to a brief intervention during a hospital stay. Other stakeholders acknowledged the significant health impact of screening and brief intervention for substance use, but cited the burden of chart-abstracted data collection and encouraged the continued development of an electronic measure. MAP stakeholders also expressed concern the use of the measure in the hospital inpatient setting, rather than a primary care setting, was not strongly linked to improved patient outcomes. The MAP also did not support SUB–3/3a (MUC16–180) due to similar concerns as identified with the SUB–2/2a measure regarding the measure’s link to improved outcomes.

With respect to MAP stakeholder concerns regarding the evidence supporting the use of the measures in the inpatient setting, we note such supporting evidence, including the evidence of the generalizability of studies to the acute inpatient setting, was included as part of the endorsement.
process and these measures received NQF endorsement. Sufficient evidence exists linking the measures to improved patient outcomes.\textsuperscript{320} In the inpatient setting,\textsuperscript{232} In addition, in light of the significant health impact of harmful substance use, and its associated healthcare costs, we believe the benefits of collecting these measure data from hospitals and publicly reporting the information outweigh the burden, and address a critical topic impacting a patient’s quality of care and health outcomes.

We note that The Joint Commission has been using these chart-abstracted measures for optional reporting since January 1, 2012.\textsuperscript{233} The chart-abstracted versions of the Substance Use measures (SUB–1, SUB–2/2a and SUB–3/3a) are also part of the IPFQR Program measure set (81 FR 57246); thus, future inclusion of the eCQM versions of these measures in the Hospital IQR Program measure set would promote programmatic alignment across these quality reporting programs. Lastly, we note that electronic versions of these measures are in development by SAMHSA; we anticipate that the eCQM versions will be ready for review within the next 18–24 months.

Additional information on the chart-abstracted versions of these measures is available in TJC’s Specification Manual for National Hospital Inpatient Quality Measures at: https://www.jointcommission.org/specifications_manual_for_national_hospital_inpatient_quality_measures.aspx.

We are inviting public comment on the possible future inclusion of one or more of the eCQM versions of the Substance Use measures (SUB–1, SUB–2/2a and SUB–3/3a) in the Hospital IQR Program. In addition, we are inviting public comment on the possible future inclusion of a composite measure comprised of all of these individual substance use measures in the Hospital IQR Program.


\textsuperscript{233} The Joint Commission, Substance Use Measures overview, available at: https://www.jointcommission.org/core_measure_sets.aspx.
quarters of eCQM data. We refer readers to section IX.E.3. of the preamble of this proposed rule, where we are proposing aligned policies for the CQM electronic reporting option in the Medicare EHR Incentive Program for eligible hospitals and CAHs.

(2) Proposed Changes to the Reporting and Submission Requirements for eCQMs for the FY 2019 Payment Determination and Subsequent Years

In this proposed rule, we are not proposing any changes to our file format requirements or reporting deadlines. However, we are proposing changes to our requirements related to eCQM electronic specification and certification. These are discussed in more detail below.

(a) File Format

In the FY 2016 IPPS/LTCH PPS final rule (80 FR 49705 through 49708), we finalized that hospitals must submit eCQM data via the Quality Reporting Document Architecture Category I (QRDA I) file format for the CY 2016 reporting period/FY 2018 payment determination. In addition, we finalized that for the CY 2016 reporting period/FY 2018 payment determination, hospitals may use third parties to submit QRDA I files on their behalf and can either use abstraction or pull the data from non-certified sources in order to then input these data into CEHRT for capture and reporting QRDA I (80 FR 49706). In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57170), we finalized our proposal to continue these eCQM reporting policies for the CY 2017 reporting period/FY 2019 payment determination and subsequent years. These finalized requirements align with those of the Medicare EHR Incentive Program for eligible hospitals and CAHs (81 FR 57255 through 57257). We are not proposing any changes to these requirements in this proposed rule.

(b) Proposed Changes to the Certification Requirements for eCQM Reporting

(i) Background

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57170 through 57171), we finalized policies that hospitals must: (1) Report eCQM data using EHR technology certified to either the 2014 or 2015 Edition for the CY 2017 reporting period/FY 2019 payment determination; and (2) report eCQM data using EHR technology certified to the 2015 Edition beginning with the CY 2018 reporting period/FY 2020 payment determination and subsequent years. As we discuss in further detail in section IX.G.4. of the preamble of this proposed rule where the same considerations are discussed in detail for the Medicare and Medicaid EHR Incentive Programs, based on our past experience with the transition from the 2011 Edition to the 2014 Edition and concerns expressed by stakeholders, we understand that transitioning to technology certified to a new Edition can be complex and can require more resources and time than anticipated, including the time necessary to effectively deploy the upgraded system and make the necessary patient safety, staff training, and workflow investments. We understand and appreciate these concerns, and are working in cooperation with our federal partners at ONC to monitor progress on the 2015 Edition upgrade. Furthermore, we believe that there are many benefits for switching to EHR technology certified to the 2015 Edition. We will work with ONC to monitor the status of EHR technology certified to the 2015 Edition and the deployment and implementation of such technology. If we identify a change in the current trends and significant issues with the certification and deployment of the 2015 Edition, we will consider additional methods to offer flexibility in CY 2018 for those hospitals that are not able to implement 2015 Edition of CEHRT. One possibility is the flexibility to use technology certified to the 2014 Edition or the 2015 Edition in CY 2018. Another option is allowing a combination of EHR technologies certified to the 2014 Edition and 2015 Edition to be used in CY 2018, for those hospitals that are not able to fully implement EHR technology certified to the 2015 Edition. We invite public comment on these options for offering flexibility in CY 2018 with regard to EHR certification requirements.

In this proposed rule, we are proposing two changes related to certification requirements with regard to eCQM reporting: (1) To require EHR technology certified to all eCQMs that are available to report; and (2) to note that certified EHR technology does not need to be recertified each time it is updated to a more recent version of the eCQM specifications, to align with the Medicare EHR Incentive Program requirements for eligible hospitals and CAHs. These proposals are discussed in more detail below.

(ii) Proposal To Require EHR Technology To Be Certified to All eCQMs That Are Available To Report for the CY 2017 Reporting Period/FY 2019 Payment Determination and the CY 2018 Reporting Period/FY 2020 Payment Determination

We refer readers to the FY 2016 IPPS/LTCH PPS final rule (80 FR 49705) where we noted that although we require CEHRT, eligible hospitals were not required to ensure that their CEHRT products were recertified to the most recent version of the electronic specifications for the clinical quality measures. In this proposed rule, we are proposing new policies regarding the Hospital IQR Program eCQM specification requirements to align with the Medicare EHR Incentive Program requirements.

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57256) for the Medicare EHR Incentive Program, we finalized the continuation of a policy that electronic submission of CQMs will require the use of the most recent version of the electronic specification for each eCQM to which the EHR is certified. For the Medicare EHR Incentive Program, we finalized that in the event an eligible hospital or CAH has EHR technology that is certified to the 2014 Edition and not certified to all of the eCQMs that are available to electronically report for the CY 2017 reporting period/FY 2019 payment determination, we require that a hospital needs to have its EHR technology certified to all such eCQMs in order to meet the reporting requirements for the CY 2017 reporting period/FY 2019 payment determination (81 FR 57256).

Further, for the Medicare EHR Incentive Program, we stated that for the CY 2017 reporting period/FY 2019 payment determination, eligible hospitals and CAHs be required to use the Spring 2016 version of the eCQM specifications available on the eCQI Resource Center Web site at: https://ecqi.healthit.gov/

In order to align with the Medicare EHR Incentive Program requirements for eligible hospitals and CAHs, in this proposed rule, we are proposing that for the CY 2017 reporting period/FY 2019 payment determination, a hospital using EHR technology certified to the 2014 or 2015 Edition, but for which such EHR technology is not certified to all 15 available eCQMs, would be required to have its EHR technology certified to all 15 eCQMs that are available to report under the Hospital IQR Program for the CY 2017 reporting period/FY 2019 payment determination in order to meet the eCQM reporting requirements for the CY 2017 reporting period/FY 2019
payment determination. We further propose that for the CY 2017 reporting period/FY 2019 payment determination, hospitals would be required to use the most recent version of the eCQM electronic specifications (in other words, the Spring 2016 version of the eCQM specifications and any applicable addenda) available on the eCQI Resource Center Web site at: https://ecqi.healthit.gov/.

For the CY 2018 reporting period/FY 2020 payment determination, we are proposing to continue our policy regarding the reporting of eCQMs, which would require the use of the most recent version of the eCQM specifications for each eCQM to which the EHR is certified. For the CY 2018 eCQM reporting period, this means hospitals would be required to use the most recent version of the eCQM electronic specifications (in other words, the Spring 2017 version of the eCQM electronic specifications and any applicable addenda) available on the eCQI Resource Center Web page https://ecqi.healthit.gov/. In addition, we are proposing to require that a hospital would need to have its EHR technology certified to all 15 available eCQMs in order to meet the reporting requirements for the CY 2018 reporting period/FY 2020 payment determination. As described in the 2015 EHR Incentive Programs final rule (80 FR 62767) and as previously finalized for the Hospital IQR Program’s eCQM reporting requirements, starting with the CY 2018 reporting period, hospitals are required to use EHR technology certified to the 2015 Edition. Furthermore, we are proposing that an EHR certified for eCQMs under the 2015 Edition certification criteria would not need to be recertified each time it is updated to a more recent version of the eCQMs. We believe it is not necessary for EHRs certified for eCQMs under the 2015 Edition certification criteria to be recertified each time it is updated to the most recent version of the eCQMs. This is because the EHR technology continues to meet the 2015 Edition certification criteria and any updates to the eCQM specifications would not impact any elements regarding certification. Therefore, we are proposing that recertification would not be necessary and would reduce the burden associated with recertification. For further discussion regarding EHR certification requirements, we refer readers to section IX.G.4. of the preamble of this proposed rule.

We are inviting public comment on these proposals.

(c) Electronic Submission Deadlines for the FY 2020 Payment Determination and Subsequent Years

We refer readers to the FY 2015 IPPS/LTCH PPS final rule (79 FR 50256 through 50259) and the FY 2016 IPPS/LTCH PPS final rule (80 FR 49705 through 49708) for our previously adopted policies to align eCQM data reporting periods and submission deadlines for both the Hospital IQR Program and the Medicare EHR Incentive Program for eligible hospitals and CAHs.

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57172), we established eCQM submission deadlines for the Hospital IQR Program. We are not proposing any changes to the eCQM submission deadlines for the FY 2020 payment determination or subsequent years.

(d) Summary

As noted in the FY 2016 IPPS/LTCH PPS final rule (80 FR 49759) and the FY 2017 IPPS/LTCH PPS final rule (81 FR 57257), we continue to encourage health IT developers to test any updates on an annual basis, including any updates to the eCQMs and eCQM reporting requirements for the Hospital IQR and Medicare EHR Incentive Programs based on the CMS Implementation Guide for Quality Reporting Document Architecture (QRDA) Category I and Category III Eligible Professional Programs and Hospital Quality Reporting (HQR) (CMS Implementation Guide for QRDA). The CMS Implementation Guide for QRDA, program specific performance calculation guidance, and eCQM electronic specifications and guidance documents are available on the eCQI Resource Center Web site at: https://ecqi.healthit.gov/.

As noted in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57172), we also continue to encourage all hospitals and vendors to submit QRDA files early, and to use one of the pre-submission testing tools for electronic reporting, such as the CMS Pre-Submission Validation Application (PSVA), to allow additional time for testing and to make sure all required data files are successfully submitted by the deadline. The PSVA can be downloaded from the Secure File Transfer (SFT) section of the QualityNet Secure Portal at: https://cportal.qualitynet.org/QNet/pgn-select.jsp.

In summary, in this FY 2018 IPPS/LTCH PPS proposed rule, for the CY 2017 reporting period/FY 2019 payment determination, we are proposing for the Hospital IQR Program that: (1) A hospital using EHR technology certified to the 2014 or 2015 Edition of CEHRT, but for which such EHR technology is not certified to all available eCQMs, would be required to have its EHR technology certified to all eCQMs that are available to report; and (2) EHR technology that is certified to all available eCQMs would not need to be recertified each time the eCQMs are updated to a more recent version of the eCQM specifications.

For the CY 2018 reporting period/FY 2020 payment determination, we are proposing for the Hospital IQR Program that: (1) A hospital using EHR technology certified to the 2015 Edition of CEHRT, but for which such EHR technology is not certified to all available eCQMs, would be required to have its EHR technology certified to all eCQMs that are available to report; and (2) EHR technology that is certified to all available eCQMs would not need to be recertified each time the eCQMs are updated to a more recent version of the eCQM specifications. Further, we are proposing that: (1) For the CY 2017 reporting period, hospitals would be required to use the most recent version of the eCQM electronic specifications (in other words, the Spring 2016 version of the eCQM specifications, and any applicable addenda); and (2) for the CY 2018 reporting period, hospitals would be required to use the most recent version of the eCQM electronic specifications (in other words, the Spring 2017 version of the eCQM specifications, and any applicable addenda). eCQM specifications are available on the eCQI Resource Center Web site at: https://ecqi.healthit.gov/.

We are inviting public comment on our proposals related to the reporting and submission requirements of eCQM data for the Hospital IQR Program. We refer readers to section IX.E.3.c. of the preamble of this proposed rule, where similar policies are described for the Medicare EHR Incentive Program for eligible hospitals and CAHs.

e. Proposed Submission Form and Method for the Proposed Voluntary Hybrid Hospital-Wide Readmission Measure With Claims and Electronic Health Record Data (NQF #2879)

(1) Background

In section IX.A.7. of the preamble of this proposed rule, we are proposing voluntary reporting of the Hybrid Hospital-Wide Measure with Claims and Electronic Health Record Data. In the FY 2016 IPPS/LTCH PPS final rule (80 FR 49701 through 49704), we signaled our intent to use core clinical data elements in the Hospital IQR Program and
requested comment on the use of the QRDA Category I (QRDA I) file format for this purpose. In that rule, we noted that many commenters supported submitting the core clinical data elements using an EHR technology certified by the ONC. In addition, some commenters were supportive of our suggested use of QRDA I specifically for reporting core clinical data elements and recommended aligning the standards for data transmission requirements with those used in other reporting programs.  

(2) Proposed Certification and File Format Requirements for Core Clinical Data Element Submissions

We are proposing that hospitals that voluntarily report data for the Hybrid Hospital-Wide Readmission measure use EHR technology certified to the 2015 Edition. We also refer readers to our discussion of EHR certification requirements for eCQM reporting above and in section IX.G.4. of the preamble of this proposed rule where the same proposed requirements are discussed in detail for the Medicare EHR Incentive Program for eligible hospitals and CAHs. In addition, we are proposing that the 13 core clinical data elements and six linking variables for the Hybrid Hospital-Wide Readmission measure be submitted using the QRDA I file format. In order to ensure that the data have been appropriately connected to the encounter, the core clinical data elements specified for risk adjustment need to be captured in relation to the start of an inpatient encounter. The QRDA I standard enables the creation of an individual patient-level quality report that contains quality data for one patient for one or more quality measures. We note that as described in section IX.A.7. of the preamble of this proposed rule, participating hospitals are expected to successfully submit data values for vital signs and six linking variables required to merge with the CMS claims data on more than 95 percent of all Medicare FFS patients who are 65 years and older discharged from the hospital during the voluntary data collection period. In addition, participating hospitals are expected to successfully submit values for laboratory test results on more than 50 percent of these patients discharged over the same time period. For further detail on QRDA I, the most recently available QRDA I specifications can be found at: http://www.hl7.org/implement/standards/product_brief.cfm?product_id=35.

We are inviting public comment on our proposals related to the reporting and submission requirements of core clinical data elements and linking variables for the proposed, voluntary Hybrid Hospital-Wide Readmission measure as discussed above.

f. Sampling and Case Thresholds for the FY 2020 Payment Determination and Subsequent Years

We refer readers to the FY 2011 IPPS/LTCH PPS final rule (75 FR 50221), the FY 2012 IPPS/LTCH PPS final rule (76 FR 51641), the FY 2013 IPPS/LTCH PPS final rule (77 FR 53537), the FY 2014 IPPS/LTCH PPS final rule (78 FR 50819), and the FY 2016 IPPS/LTCH PPS final rule (80 FR 49709) for details on our sampling and case thresholds for the FY 2016 payment determination and subsequent years. We are not proposing any changes to our sampling and case threshold policies.

g. HCAHPS Administration and Submission Requirements for the FY 2020 Payment Determination and Subsequent Years

We refer readers to the FY 2011 IPPS/LTCH PPS final rule (75 FR 50220), the FY 2012 IPPS/LTCH PPS final rule (76 FR 51641 through 51643), the FY 2013 IPPS/LTCH PPS final rule (77 FR 53537 through 53538), and the FY 2014 IPPS/LTCH PPS final rule (78 FR 50819 through 50820) for details on previously-adopted HCAHPS requirements. We also refer hospitals and HCAHPS Survey vendors to the official HCAHPS Web site at: http://www.hcahpsonline.org for new information and program updates regarding the HCAHPS Survey, its administration, oversight, and data adjustments. We refer readers to section IX.A.6.a. of the preamble of this proposed rule for details on our proposal to refine the three questions of the Pain Management measure in the HCAHPS Survey. While we are proposing to refine the survey with respect to the questions about pain management in section IX.A.6.a. of the preamble of this proposed rule, we are not proposing any changes to the HCAHPS administration nor the HCAHPS submission requirements.

h. Data Submission Requirements for Structural Measures for the FY 2020 Payment Determination and Subsequent Years

We refer readers to the FY 2012 IPPS/LTCH PPS final rule (76 FR 51643 through 51644) and the FY 2013 IPPS/LTCH PPS final rule (77 FR 53538 through 53539) for details on the data submission requirements for structural measures. We are not proposing any changes to data submission requirements for structural measures.

i. Data Submission and Reporting Requirements for HAI Measures Reported via NHSN

For details on the data submission and reporting requirements for HAI measures reported via the CDC’s NHSN Web site, we refer readers to the FY 2012 IPPS/LTCH PPS final rule (76 FR 51629 through 51633; 51644 through 51645), the FY 2013 IPPS/LTCH PPS final rule (77 FR 53539), the FY 2014 IPPS/LTCH PPS final rule (78 FR 50821 through 50822), and the FY 2015 IPPS/LTCH PPS final rule (79 FR 50259 through 50262). The data submission deadlines are posted on the QualityNet Web site at: http://www.QualityNet.org/. We are not proposing any changes to data submission and reporting requirements for HAI measures reported via the NHSN.

11. Proposed Modifications to the Validation of Hospital IQR Program Data

a. Background

In the FY 2013 IPPS/LTCH PPS final rule (77 FR 53539 through 53553), we finalized the processes and procedures for validation of chart-abstracted measures in the Hospital IQR Program for the FY 2015 payment determination and subsequent years; the FY 2013 IPPS/LTCH PPS final rule also contains a comprehensive summary of all procedures finalized in previous years that are still in effect. We refer readers to the FY 2014 IPPS/LTCH PPS final rule (78 FR 50822 through 50835), the FY 2015 IPPS/LTCH PPS final rule (79 FR 50262 through 50273), and the FY 2016 IPPS/LTCH PPS final rule (80 FR 49710 through 49712) for detailed information on the modifications to these processes finalized for the FY 2016, FY 2017, and FY 2018 payment determinations and subsequent years. In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57173 through 57181), we finalized our proposal to update the validation procedures in order to incorporate a process for validating eCQM data for the FY 2020 payment determination and subsequent years. Specifically, we finalized a policy to: (1) Validate eCQM data submitted by up to 200 hospitals selected via random sample; (2) exclude any hospital selected for chart-abstracted measure validation as well as any hospital that has been granted a Hospital IQR Program Extraordinary Circumstances Exemption for the applicable eCQM reporting period; and (3) randomly select 32 cases from the QRDA I files submitted by each hospital selected for eCQM validation for the FY 2020 payment determination and subsequent years. As described in the FY 2017...
In this proposed rule, we are proposing to modify policies for eCQM validation for the FY 2020 payment determination and subsequent years. First, for hospitals selected to participate in validation of cEIQMs, we are proposing that we will select eight cases per quarter for the CY 2017 reporting period/FY 2020 payment determination and subsequent years. We note that this proposal is contingent upon whether or not our proposed modifications to eCQM reporting requirements for the FY 2017 IPPS/LTCH PPS final rule (81 FR 57178) will be applied before the random selection of 200 hospitals for eCQM validation, so that hospitals meeting any of these exclusions would not be eligible for selection. We believe that these proposals improve the likelihood that there would be sufficient data for validation obtained from the hospitals selected for eCQM data validation.

We are inviting public comment on our proposals to: (1) Exclude any hospital that does not have at least five discharges for at least one reported eCQM included among their QRDA I file submissions in eCQM validation; and/or (2) to exclude from selection hospitals meeting either of the two exclusion criteria finalized in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57178) as discussed above.

(b) Selection of Cases
We have not previously specified processes for the selection of cases for eCQM validation. For the FY 2020 payment determination and subsequent years, we are proposing to exclude the following cases from validation for those hospitals selected to participate in eCQM validation:
- Episodes of care that are longer than 120 days; and
- Cases with a zero denominator for each measure.

We believe that excluding episodes of care that are longer than 120 days will reduce the reporting burden on hospitals selected for eCQM validation, as the volume of data reported for longer cases is greater. Further, we believe that excluding cases with zero denominators for each measure would ensure that we perform validation on cases with applicable measure data. We note that this proposed exclusion applies to cases, rather than measures. However, a measure would not be validated if a hospital did not have any applicable cases for the measure.

We are inviting public comments on our proposal to exclude: (1) Episodes of care that are longer than 120 days; and (2) cases with a zero denominator for each measure from eCQM validation for the FY 2020 payment determination and subsequent years as discussed above.

(3) Medical Record Submission Requirements and Scoring
(a) Medical Record Submission Requirements
In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57179), we finalized that hospitals participating in eCQM validation would be required to submit: (1) 16 cases over two calendar quarters (eight cases × two quarters) for the CY 2017 reporting period/FY 2020 payment determination; and (2) 24 cases over three quarters (eight cases × three quarters) for the CY 2018 reporting period/FY 2021 payment determination.

We are inviting public comment on these proposals as discussed above.

(2) Selection of Hospitals and Cases
In this proposed rule, for the CY 2017 reporting period/FY 2020 payment determination and subsequent years, we are proposing changes to our policies related to the selection of hospitals and cases for eCQM validation to: (1) Expand the types of hospitals that could be excluded; and (2) expand the type of cases excluded from selection. These proposals are discussed in more detail below.

(a) Selection of Hospitals
As previously finalized in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57174–57178), we will validate eCQM data submitted by up to 200 hospitals selected via random sample. Further, we finalized that the following hospitals may be excluded from this random sample of 200 hospitals selected for eCQM validation (81 FR 57178):
- Any hospital selected for chart-abstracted measure validation; and
- Any hospital that has been granted a Hospital IQR Program Extraordinary Circumstances Exemption for the applicable eCQM reporting period.

In this proposed rule, we are proposing to expand the types of hospitals that could be excluded. For the FY 2020 payment determination and subsequent years, we are proposing to also exclude any hospital that does not have at least five discharges for at least one reported eCQM included among their QRDA I file submissions. In addition, we are proposing that the three exclusions described above would be applied before the random selection of 200 hospitals for eCQM validation, so that hospitals meeting any of these exclusions would not be eligible for selection.
validation for the FY 2020 payment determination and subsequent years are required to: (1) Submit data by 30 calendar days following the medical records request date listed on the CDAC request form; (2) provide sufficient patient level information necessary to match the requested medical record to the original Hospital IQR Program submitted eCQM measure data record; and (3) submit records in PDF file format through QualityNet using the Secure File Transfer (SFT). We also finalized, for the FY 2020 payment determination only, that for hospitals selected for eCQM validation, that: (1) We require submission of at least 75 percent of sampled eCQM measure medical records in a timely and complete manner; and (2) the accuracy of eCQM data submitted for validation would not affect a hospital’s validation score (81 FR 57180). In this proposed rule, we are not making any changes related to these operational procedures. However, we are proposing to continue these policies for the FY 2021 payment determination and subsequent years.

In this proposed rule, we are proposing to extend to the FY 2021 payment determination and subsequent years our previously finalized medical record submission policy for eCQM validation, as finalized in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57181), requiring submission of at least 75 percent of sampled eCQM measure medical records in a timely and complete manner. We are proposing to extend to the FY 2021 payment determination our previously finalized medical record submission policy for eCQM validation, as finalized in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57181), that the accuracy of eCQM data submitted for validation would not affect a hospital’s validation score. We note that if our proposals in section IX.A.8. of the preamble of this proposed rule to require two quarters of data for CY 2017 eCQM data submission and eight cases per quarter for hospitals selected for validation (16 total cases for the entire data collection period), are finalized as proposed, and hospitals selected for eCQM validation are required to submit complete information for 75 percent of requested cases as previously finalized, then those hospitals would be required to submit information for at least 12 records, or 75 percent of the requested 16 records for the FY 2020 payment determination. Similarly, if our proposals: (1) To continue our medical record submission policies for the FY 2021 payment determination and subsequent years; (2) to require three quarters of data for CY 2018 eCQM data submission and eight cases per quarter for hospitals selected for validation (24 total cases for the entire data collection period) as detailed in section IX.A.8. of the preamble of this proposed rule; and (3) that hospitals selected for eCQM validation are required to submit complete information for 75 percent of requested cases are all finalized as proposed, then those hospitals would be required to submit complete information for at least 18 records, or 75 percent of the requested 24 records for the FY 2021 payment determination.

Furthermore, as finalized in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57180) for the FY 2020 payment determination, we are proposing for the FY 2021 payment determination and subsequent years that any hospital that fails any validation requirement, such as submission of records in PDF file format within 30 days of the date listed on the CDAC medical records request, and/or submission of complete information for at least 75 percent of the requested records, would be considered not to have met the eCQM validation requirements and would be subject to a one-quarter reduction of the applicable percentage increase for not meeting all Hospital IQR Program requirements.

We are inviting public comment on our proposal as discussed above.

(b) Scoring

As finalized in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57178) for the FY 2020 payment determination only, the accuracy of eCQM data (the extent to which eCQM data reported for validation matches the data previously reported in the QRDA I files for eCQM reporting) submitted for validation will not affect a hospital’s validation score. In this proposed rule, we are proposing the continuation of this policy for the FY 2021 payment determination, such that the accuracy of eCQM data submitted for validation would not affect a hospital’s validation score. We intend for the accuracy of eCQM validation to affect validation scores in the future and would propose any changes related to this in future rulemaking. The data submission deadlines and additional details about the eCQM validation procedures will be posted on the QualityNet Web site at: http://www.QualityNet.org/.

We are inviting public comment on this proposal as discussed above.

c. Proposed Modifications to the Educational Review Process for Chart-Abstracted Measures Validation

(1) Background

In the FY 2015 IPPS/LTCH PPS final rule, we stated that we rely on hospitals to request an educational review or appeal cases to identify any potential CDAC or CMS errors (79 FR 50260). We also noted that a hospital may request from CMS at any time an educational review to better understand whether or not we reached a correct conclusion during validation; hospitals that fail to meet Hospital IQR Program validation requirements have 30 days to appeal after this determination (79 FR 50260). We have described our processes for educational review on the QualityNet Web site.234 We note that historically this process functioned as an outreach opportunity we provided hospitals, but based on our experience, and more robust validation requirements, we believe that it would benefit hospitals to propose formalizing this process.

Under the current process, if the results of an educational review indicate that CDAC or CMS has incorrectly scored a hospital, those scores are not changed unless and until the hospital submits a reconsideration request. Therefore, in this proposed rule, we are proposing: (1) To formalize this process; and (2) to update the process to specify that if the results of an educational review indicate that we incorrectly scored a hospital, the corrected score would be used to compute the hospital’s final validation score whether or not the hospital submits a reconsideration request. These proposals are discussed in more detail below.

Stakeholder feedback, provided via email, has indicated that while the educational review process is helpful to participating hospitals, it is limited in its impact, given that a hospital’s score is not corrected even after an educational review determines that CMS reached an incorrect conclusion regarding a hospital’s validation score for a given quarter. Based on this feedback, we are proposing to change the Hospital IQR Program’s chart-abstracted measure validation educational review process. Our goal is to reduce the number of reconsideration requests by identifying and correcting errors before the final yearly validation score is derived. By identifying and correcting any mistakes early on, this process could help decrease the burden

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during the annual reconsideration process, both for hospitals and CMS.

2) Proposed Educational Review Process Modifications for the FY 2020 Payment Determination and Subsequent Years

(a) Request for Educational Review

Under this proposal, the educational review request process, as well as our procedures for responding to requests, remain the same. Specifically, under the current process, hospitals may request an educational review if they believe they have been scored incorrectly or if they have questions about their score. We would provide the results of the educational review, outlining the findings of whether the scores were correct or incorrect, to the requesting hospital through secure file transfer.

In this proposed rule, we are proposing to formalize this process. In formalizing our current procedures, the educational review request process, as well as our procedures for responding to requests, would remain the same. First, we are proposing that, for the FY 2020 payment determination and subsequent years, a hospital may request from CMS an educational review to better understand whether or not CDAC or CMS reached a correct conclusion during validation for the first three quarters of validation. Specifically, upon receipt of an unsatisfactory score, a hospital would have 30 calendar days to contact the Validation Support Contractor (VSC) to solicit a written explanation of the provided score. We note that currently hospitals receive validation results on a quarterly basis, and that would not change under this proposed process. Accordingly, under this proposal, an educational review could be requested on a quarterly basis for the first three quarters of validation. Results of the educational review would be provided to hospitals via secure file transfer.

Second, we are proposing that the process used to evaluate whether or not validation results are correct would be the same in both an educational review and a reconsideration request. Specifically, as finalized in the FY 2012 IPPS/LTCH PPS final rule for the Hospital IQR Program’s reconsideration request process, we are proposing that upon receipt of an educational review request, we would review the data elements that were labeled as mismatched, as well as the written justifications provided by the hospitals, and make a decision on the educational review request.

(b) Scoring Update

For the FY 2020 payment determination and subsequent years, we are proposing that if an educational review, that is requested for any of the first 3 quarters of validation, yields incorrect CMS validation results for chart-abstracted measures, we would use the corrected quarterly score, as recalculated during the educational review process, to compute the final confidence interval (CI). These corrected scores would be applicable to the corresponding quarter, within the first 3 quarters of validation, for which a request was submitted. We note that under this proposal, the quarterly validation reports issued to hospitals would not be changed to reflect the updated score due to the burden associated with reissuing corrected reports. Beginning with the FY 2020 payment determination, we are proposing to use the revised score identified through an educational review when determining whether or not a hospital failed validation. Further, under this proposal, as with the current educational review process, corrected scores identified through the educational review would only be used if they indicate that the hospital performed more favorably than previously determined.

Under this proposal, the educational review request process, as well as our procedures for responding to requests, remain the same. We also note that, in accordance with our previously established policies, a hospital may still request reconsideration even if an educational review determined that a hospital was scored correctly. Hospitals that fail Hospital IQR Program requirements, which include validation, can request reconsideration at the end of the year after the annual payment update has been made. We refer readers to section IX.A.14. of the preamble of this proposed rule for a discussion about our reconsideration and appeals process. We note that under this proposal, corrected scores identified through the educational review would only be used if they indicate that the hospital performed more favorably than previously determined.

In addition, we note that for the last quarter of validation, because of the need to calculate the confidence interval in a timely manner and the insufficient time available to conduct educational reviews, the existing reconsideration process would be used to dispute an unsatisfactory validation result. If a hospital does not fail validation they still would have the opportunity to request an educational review within 30 days of receiving the results.

We are inviting public comment on our proposals to formalize the educational review process and use this process to correct scores for the first three quarters of chart-abstracted measure validation as discussed above.

13. Public Display Requirements for the FY 2020 Payment Determination and Subsequent Years

We refer readers to the FY 2008 IPPS/LTCH PPS final rule (72 FR 47364), the FY 2011 IPPS/LTCH PPS final rule (75 FR 50230), the FY 2012 IPPS/LTCH PPS final rule (76 FR 51650), the FY 2013 IPPS/LTCH PPS final rule (77 FR 53554), the FY 2014 IPPS/LTCH PPS final rule (78 FR 50836), the FY 2015 IPPS/LTCH PPS final rule (79 FR 50277), and the FY 2016 final rule (80 FR 49712 through 49713) for details on public display requirements. The Hospital IQR Program quality measures are typically reported on the Hospital Compare Web site at: http://www.medicare.gov/hospitalcompare, but on occasion are reported on other CMS Web sites such as: https://data.medicare.gov.

In this proposed rule, we are not proposing any changes to public display requirements. However, we are soliciting public comment on potential options for confidential and public reporting data sets that may be reported on the public display requirements, e.g., identifying measures that incorporate health equity. In this proposed rule, we are seeking additional public comment on the potential confidential and public reporting of Hospital 30-day, All-Cause, Risk-Standardized Readmission Rate Following Pneumonia Hospitalization (NQF #0506), the Pneumonia Readmission measure, and the Hospital 30-Day, All-Cause, Risk-Standardized
Mortality Rate Following Pneumonia Hospitalization (NQF #0468), (the Pneumonia Mortality measure), data stratified specifically by patient dual eligibility status.

b. Potential Options for Confidential and Public Reporting of Hospital IQR Measures Stratified by Patient Dual Eligibility Status

(1) Background

In section IX.A.1.d. of the preamble of this proposed rule, we discuss the importance of improving beneficiary outcomes including reducing health disparities, and our commitment to ensuring that medically complex patients, as well as those with social risk factors, receive excellent care. As we note in section IX.A.1.d. of the preamble of this proposed rule, studies show that social risk factors, such as earning a low-income, belonging to a racial or ethnic minority group, or living with a disability, to be associated with poor health outcomes and some of this disparity is related to the quality of health care.235 One of our core objectives is to improve health outcomes for all beneficiaries, and ensure that complex patients as well as those with social risk factors receive excellent care. Within this context, recent reports by the Office of the Assistant Secretary for Planning and Evaluation (ASPE) and the National Academies of Sciences, Engineering, and Medicine have examined the influence of social risk factors in CMS value-based purchasing programs.236 In addition, as noted in the FY 2017 IPPS/LTC PPS final rule (81 FR 57185), the NQF has undertaken a 2-year trial period in which certain new measures and ongoing maintenance review have been assessed to determine if risk adjustment for social risk factors is appropriate for these measures.237

As part of this effort, we are soliciting feedback on which social risk factors provide information that is most valuable to stakeholders. We also are seeking public comment on confidential reporting and future public reporting of some of our measures, specifically the Pneumonia Readmission measure (NQF #0506) and the Pneumonia Mortality measure (NQF #0468), stratified by patient dual eligibility. There are two potential purposes for providing information on hospital results stratified by dual eligibility. The approach we are considering would illuminate differences in outcome rates among patient groups within a hospital and would also allow for a comparison of those differences, or disparities, across hospitals. We also considered an alternative approach that would measure outcome rates for subgroups of patients, such as the dual eligible patients, across hospitals. However, this alternative would not allow for an examination of the difference in rates between groups (for example dual eligible compared to non-dual eligible).

The goal of measuring and monitoring disparities in patient outcomes for specific sub-groups of patients within hospitals is to reduce health inequities, improve health care quality for vulnerable populations, and promote greater transparency for health care consumers. This is in alignment with the CMS Quality Strategy and the ASPE report to Congress, which stated performance rates including readmission rates stratified by social risk should be developed and considered for hospital specific confidential preview reports and public reporting in places such as Hospital Compare, so hospitals, health systems, policymakers, and consumers can see and address important disparities in care.

Many levers exist for addressing and improving disparities in care and outcomes. The 21st Century Cures Act (Pub. L. 114–255) addresses payment penalty scoring in the Hospital Readmissions Reduction Program by identifying hospitals based on their proportion of dual eligible patients and supporting improvement efforts for hospitals caring for patients with social risk factors by setting penalty thresholds among similar peer hospitals.240 As discussed in sections V.I.7. through V.I.10. of the preamble of this proposed rule, the Hospital Readmissions Reduction Program, as required by the statute, is proposing to use dual eligibility as a marker of poverty, one key patient social risk factor, and we would like to move in that direction for the Hospital IQR Program as well in the future. In the Hospital IQR Program, we are exploring methods to distinguish vulnerable patients with social risk factors, such as poverty. As such, we intend to use dual eligible status among the over 65 year old patients included in the measures as a marker of poverty.

Dual eligible status describes whether Medicare beneficiaries are also enrolled in Medicaid. We use dual enrollment in Medicare and Medicaid as a marker for a beneficiary having low income and/or few assets. The recent report to Congress by ASPE has shown that dual eligibility was the most powerful predictor of poor health care outcomes among the social risk factors they tested.241

The Hospital Compare Web site currently displays readmission rates for each hospital’s patients together, but does not specifically highlight hospitals’ quality of care for vulnerable populations. We believe stratifying data by social risk factors would supplement the current reporting of the Pneumonia Readmission and Pneumonia Mortality measures by highlighting disparities, that is, differences in outcomes, within hospitals that are not simply due to differences in illness level, to the extent that such disparities exist for any given hospital. To do so, we developed a method to quantify the disparities of readmission and mortality, between these groups within each hospital after accounting for patient case mix. The disparities indicator used in the hospital specific confidential preview reports would provide information assessing the increased odds, or rates, of readmission for dual eligible patients admitted to the same hospital, after accounting for differences in age and comorbidities.

For the Hospital IQR Program, we are considering options to improve health disparities among patient groups within hospitals by increasing the transparency of disparities among patients within hospitals and the ability to compare these disparities across hospitals. This would be accomplished by the methods described below. Our alternative approach, also described below, to measure outcome rates for subgroups of patients, such as the dual eligible patients, across hospitals, would examine the performance of hospitals.

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239 Ibid.

240 Ibid.

241 Ibid.
on the subgroup of dual eligible patients.

We previously sought public comment on the potential public reporting of quality measures data stratified by race, ethnicity, sex, and disability and future hospital quality measures that incorporate health equity in the FY 2017 IPPS/LTC PPS final rule (81 FR 57167 through 57168). In general, commenters supported the development of health equity measures and their inclusion in the Hospital IQR Program (81 FR 57167). In particular, stakeholders noted that stratified measures could serve as tools for hospitals to identify gaps in outcomes for different groups of patients, improve the quality of health care for all patients, empower consumers to make informed decisions about health care, and encouraged us to stratify measures by other social risk factors such as age, income, and educational attainment (81 FR 57167). However, commenters raised concerns about the small denominator sample size associated with measure stratification by social risk factors, which would skew the reliability of stratified quality measures. Commenters also were concerned that it may not be a simple task to stratify measures by race, ethnicity, sex, and disability because specific considerations are required for every measure and each reporting mechanism to implement such a requirement (81 FR 57168). For more details on the public comments, we refer the readers to the FY 2017 IPPS/ LTC PPS final rule (81 FR 57167 through 57168).

We acknowledge the complexity of interpreting stratified outcome measures. Due to this complexity, prior to publicly reporting stratified outcome measure data, as early as the summer of 2018 using data from the FY 2019 reporting period (July 1, 2014 through June 30, 2017), we are considering first providing hospitals with confidential preview reports containing stratified results for certain Hospital IQR Program measures, specifically the Pneumonia Readmission measure and the Pneumonia Mortality (MORT–30–PN) measure; (3) a potential methodology for illuminating differences in outcomes rates among patient groups within a hospital that would also allow for a comparison of those differences, or disparities, across hospitals; (4) an alternative methodology that compares performance for patient subgroups across hospitals but does not provide information on hospital disparities and any additional suggested methodologies for calculating stratified results by patient dual eligibility within the hospital, or more specifically, differences in outcome rates for the dual eligible and non-dual eligible patients in the measures. This would allow us to obtain feedback on reporting options and to ensure the information is reliable, valid, and understandable prior to any future public display on Hospital Compare.

Our goal in producing stratified results is to provide information about disparities in patient outcomes within hospitals to the extent that they exist for a given hospital. This information would supplement the assessment of overall hospital quality provided through the current measures of readmission and mortality rates; these measures would remain unchanged. We discuss below the methods and results of stratification for the current Hospital 30-day, All-Cause, Risk-Standardized Readmission Rate Following Pneumonia Hospitalization (NQF #0506) (the READ–30–PN or Pneumonia Readmission measure).

The stratified results would provide hospitals with confidential reporting with information that could illuminate any disparities in care and outcome that can be targeted through quality improvement efforts. Then for the future, we are considering publicly posting both of these results on Hospital Compare to allow consumers and other stakeholders to view critical information about the care and outcomes of subgroups of patients, particularly those with social risk factors. This information could drive consumer choice and spark improvement efforts targeting dual eligible patients. In the future, we would also consider expanding this approach to other social risk factors and other measures.

We are inviting public comment on: (1) Which social risk factors provide information that is most valuable to stakeholders; (2) providing hospitals with confidential preview reports containing stratified results for certain Hospital IQR Program measures, specifically the Pneumonia Readmission measure and the Pneumonia Mortality (MORT–30–PN) measure; (3) a potential methodology for illuminating differences in outcomes rates among patient groups within a hospital that would also allow for a comparison of those differences, or disparities, across hospitals; (4) an alternative methodology that compares performance for patient subgroups across hospitals but does not provide information on hospital disparities and any additional suggested methodologies for calculating stratified results by patient dual eligibility within the hospital, or more specifically, differences in outcome rates for the dual eligible and non-dual eligible patients in the measures. These are discussed in more detail below.

(2) Hospital Specific Confidential Preview Reports Prior to Publicly Reporting Stratified Data

We are seeking public comment on the possibility of providing hospitals specific confidential preview reports containing the results of the Pneumonia Readmission (NQF #0506) and Pneumonia Mortality (NQF #0468) measures stratified by patient dual eligibility, as early as the summer of 2018 using data from the FY 2019 reporting period (July 1, 2014 through June 30, 2017), prior to any future potential public reporting of this data. The current publicly reported measures used in the Hospital IQR Program and reported on the Hospital Compare Web site would remain unchanged.

Following the time period during which hospitals received confidential preview reports, we may display stratified results on Hospital Compare solely for the purpose of “stratification,” that is, producing results to describe differences between subgroups within the hospital.

(3) Potential Methodology for Calculating Stratified Results by Patient Dual Eligibility Status

(a) Background

Under any future option to stratify measure results by patient dual eligibility status, we intend to focus on disparities between dual eligible and non-dual eligible patients because dual eligibility is an important social risk factor among the Medicare Fee-for-Service population and is feasible to measure.242 In order to provide information about differences in readmission outcomes for dual eligible and non-dual eligible patients within a hospital that may be due to quality differences, we need a methodology that accounts for any differences in comorbidities, age, and other risk factors between these groups of patients. Such a methodology ensures that differences in outcomes are not simply due to differences in clinical severity and comorbid conditions among the patient groups. Therefore, any approach to identifying within-hospital disparities for readmission measures by patients’ dual eligibility would build on the methodology used to calculate the currently implemented RSRRs.243 As the Pneumonia Readmission measure (NQF #0506) is currently specified, risk-adjusted rates are estimated using a hierarchical logistic regression to account for the clustering of observations within hospitals and differences in the number of admissions across hospitals.


(b) Option To Measure Difference in Outcomes by Adding Three Additional Factors to Current Statistical Models

There is both a hospital and patient-level effect of dual eligibility on readmission risk. We have considered the hospital fixed effect in our approaches to stratafication (described in the preceding section IX.A.13.b.(3)(a) of the preamble of this proposed rule) because without it, we will introduce bias in the patient-level dual eligibility, which would produce misleading results. The statistical approach we may employ in the future would use current statistical models and add three additional factors to the statistical model for the purposes of measuring differences in outcomes: (1) An indicator for patient-level dual eligibility; (2) a hospital-level dual eligible factor (for example, percentage of dual eligible patients in each hospital); and (3) a hospital-specific indicator (random coefficient) for dual eligibility. This third factor, the hospital-specific random coefficient for dual eligibility, assesses the disparity or difference in readmissions for dual eligible patients within a specific hospital after accounting for other factors, such as differences in clinical disease or comorbidity conditions. The first two factors, (the patient-level dual eligibility coefficient, which represents the overall difference between dual and non-dual groups in the entire country, and the hospital-level dual eligible factor, which reflects the difference in readmission rate between hospitals with different proportions of dual eligible patients) are only included in order to be able to interpret the third factor random coefficient and ensure it is specific to a particular hospital. It is the third factor hospital-specific indicator, which would be used to calculate the differences in readmission rates between the dual and non-dual eligible patients within the hospital.

Using this method, within-hospital disparities in readmissions between dual eligible and non-dual eligible patients would be included in confidential hospital specific preview reports in addition to the currently calculated and displayed Pneumonia Readmission (NQF #0506) and Pneumonia Mortality (NQF #0468) measures. For ease of interpretation, we would provide information in the form of odds ratios (that is, the increased odds of readmission for dual eligible patients at a given hospital) or, alternatively, the average difference in readmission rates between dual and non-dual patients after accounting for differences in other risk-factors.

To calculate odds ratios, we would convert hospital-specific coefficients for dual eligibility into odds ratios. Odds ratios compare dual eligible patients relative to non-dual eligible patients in terms of their risk of readmission, assuming that the two groups have the same case mix (that is, comorbidities). If the readmission rate is the same in both groups, the odds ratio is 1. If the odds ratio is greater than 1, it would mean that dual eligible patients have worse readmission rates, and vice versa. To estimate the average difference of readmission rates between dual and non-dual beneficiaries for each hospital, we would first calculate the predicted probabilities of being readmitted by assuming all patients are dual eligible or all patients are non-dual eligible in a hospital. The difference between the two predicted probabilities is the average difference in the readmission rate between the two groups of patients at each hospital.

Rather than assuming a uniform impact of dual eligible and non-dual eligible status across hospitals, this approach would assess the impact of dual eligibility across all hospitals separately, recognizing that socioeconomic disparities of patients may be greater or lesser at some hospitals as compared with others. Therefore, this approach would allow quantification of the difference in readmissions between dual and non-dual eligible patients within each hospital, as long as a hospital has a sufficient number of cases to produce a reliable estimate for both groups.

In summary, this statistical model would uniquely identify disparities in readmission rates for dual eligible beneficiaries compared to non-dual eligible beneficiaries, after controlling for patients’ prior medical history and age for each hospital. This random coefficient for dual eligibility within the statistical model would indicate how much readmission rates at the same hospital would differ between two patients at that hospital with exactly the same age and underlying risk factors (those comorbid clinical conditions included in the statistical model), but differ with respect to dual eligibility.

(c) Option To Measure Difference in Outcomes Using Current Statistical Models

Depending on the information that is most useful to stakeholders, an alternative approach to examining readmission rates among dual eligible patients could be considered. To examine the relative performance of hospitals on readmission rates for their dual eligible patients, rather than to compare hospitals on within-hospital disparities in rates, we could calculate the current measures’ statistical model (without the additional factors mentioned above) and include only dual eligible patients. Similarly, this could be done for non-dual eligible patients. This approach of using two separate models for the separate patient subgroups would produce information on readmission rates for dual eligible patients at one hospital compared to another (or non-dual eligible patients across hospitals). There is a trade-off; because of the use of two separate statistical models, this approach would not ensure consistent treatment of risk factors across patient groups and could not be used to compare readmission rates for two groups within a hospital.

(d) Summary of Statistical Method Options

We intend to provide information on the difference in readmission rates of dual or non-dual beneficiaries within hospitals and also provide information for hospitals and consumers on the relative disparities across hospitals. We are soliciting public comment on the information that stakeholders would find most useful and any additional suggested methodologies for calculating stratified results by patient dual eligible status.

The hospital specific confidential preview reports containing data stratified by patient dual eligibility would be modelled after current hospital specific confidential preview reports and include patient-level data for hospitalizations included in the measure. The current hospital specific confidential preview reports would be supplemented by information for each patient on their dual eligible status and a summary of the difference in readmission rates for dual eligible patients in the hospital as compared to other hospitals in the state and nation. We are inviting public comment on both methodologies, as described above, to produce stratified results by determining the differences in readmission and mortality by dual eligible status within a hospital, and a comparison of those disparities across
hospitals, accounting for differences in comorbidities, age, and other risk factors between dual eligible and non-dual eligible patients.

Data Sources

To provide an example of the statistical approach we could apply, below we describe stratified results by patient dual eligibility for the Pneumonia Readmission measure (NQF #0506), using the first calculation methodology described in section IX.A.13.b.(3)(b) of the preamble of this proposed rule, above. To calculate the example rate, we used the CMS administrative claims data from each index pneumonia hospitalization, as well as from inpatient and outpatient Medicare claims from the 12 months prior to the hospitalization from July 2012 to June 2015 to calculate the publicly reported RSRRs following pneumonia hospitalization (NQF #0506) in the July 2016 Hospital Compare update. Both the cohort and the risk-adjustment model remained unchanged. For more details on the publicly reported RSRRs following pneumonia as currently implemented, we refer readers to its measure methodology report and measure update zip file on our Web site at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInits/Measure-Methodology.html. The data was then linked to CMS denominator files (2012 to 2015) to derive the indicator of dual eligibility for each patient admission (1.3 percent index admissions were excluded because there is no information available in the denominator files).

We conducted preliminary analyses on the Pneumonia Readmission measure (NQF #0506) and determined that there is a total of 3,851 hospitals that have a least 25 included index hospitalizations overall, and at least 10 dual eligible and 10 non-dual eligible index hospitalizations for which we could report outcome disparity (82 percent of hospitals). The minimum sample size for 25 hospitalizations is consistent with the current publicly reported measures. We imposed an additional requirement of at least 10 dual eligible and 10 non-dual eligible index hospitalizations for this example to ensure we had adequate numbers to observe any meaningful differences in outcome. We used this requirement because if a hospital has fewer than 10 patients in one subgroup of patients, it is not clear that readmission rates for that group as compared to others would be reliable, nor that it is meaningful or has face validity to measure stratified rates for hospitals very few of one of the categories of patients. We welcome public comment on this sample size determination.

The observed readmission rate within 30-days of index discharge for all patients was 17.1 percent when we did not adjust for patients’ prior medical history, and dual eligible beneficiaries had an approximately 3 percent higher readmission rate. Results from the hierarchical model indicate there is a statistically significant association between dual eligibility and pneumonia readmission (adjusted odds ratio, 1.07; 95 percent CI, 1.06–1.08). In addition, there is substantial variation in the relationship between dual eligibility and readmission across hospitals (Median odd ratio, 1.06; Min., 0.95; Max., 1.22). Findings also revealed that dual eligible patients are more likely to get readmitted in 95 percent of hospitals.

Future Potential Public Display

We are inviting public comment on the potential future public reporting of certain outcomes measures, such as the Pneumonia Readmission (NQF #0506) and Pneumonia Mortality measures (NQF #0468), stratified by social risk factors, specifically dual eligibility status, to illuminate within-hospital disparities. If we decide to display measure data stratified by dual eligible status on the Hospital Compare Web site, we would clearly differentiate between the measure information we currently display and the measure information that is stratified by patients’ dual eligibility status. In addition, as discussed above, if we decide to display measure data stratified by dual eligible status on the Hospital Compare Web site, hospitals would receive information about their stratified readmission rates for a certain period of time through hospital specific confidential preview reports prior to the public reporting of any information. We are inviting public comment on this future consideration to display the stratified measure results, in addition to the current measure results, for certain Hospital IQR Program measures, in future reporting years. We note that public display of measure data stratified by social risk factors such as dual eligible status would not occur until after a period of confidential reporting.

Summary

To summarize, we are inviting public comment on: (1) Which social risk factors provide the most valuable information to stakeholders; (2) providing hospitals with confidential preview reports containing stratified results for certain Hospital IQR Program measures, specifically the Pneumonia Readmission measure and the Pneumonia Mortality measure; (3) a potential methodology for illuminating differences in outcomes rates among patient groups within a hospital and would also allow for a comparison of those differences, or disparities, across hospitals; (4) an alternative methodology that compares performance for patient subgroups across hospitals but does not provide information on within hospital disparities and any additional suggested methodologies for calculating stratified results by patient dual eligibility status; and (5) future public reporting of these same measures stratified by patient dual eligibility status on Hospital Compare as discussed above.

14. Reconsideration and Appeal Procedures for the FY 2020 Payment Determination and Subsequent Years

We refer readers to the FY 2012 IPPS/LTCH PPS final rule (76 FR 51650 through 51651), the FY 2014 IPPS/LTCH PPS final rule (78 FR 50836), and 42 CFR 412.140(e) for details on reconsideration and appeal procedures for the FY 2017 payment determination and subsequent years. We are not proposing any changes to the reconsideration and appeals procedures in this proposed rule.

15. Proposed Change to the Hospital IQR Program Extraordinary Circumstances Exceptions (ECE) Policy

a. Background

We refer readers to the FY 2012 IPPS/LTCH PPS final rule (76 FR 51651 through 51652), the FY 2014 IPPS/LTCH PPS final rule (78 FR 50836 through 50837), the FY 2015 IPPS/LTCH PPS final rule (79 FR 50277), the FY 2016 IPPS/LTCH PPS final rule (80 FR 49713), the FY 2017 IPPS/LTCH PPS final rule (81 FR 57181 through 57182), and 42 CFR 412.140(c)(2) for details on the current Hospital IQR Program ECE policy. We also refer readers to the
propose to make conforming changes to the regulations at 42 CFR 412.140(c)(2) to reflect our previously finalized policy that the ECE request form be submitted within 90 days following the date the extraordinary circumstance occurred (81 FR 57181 through 57182). In addition, we are proposing to make conforming changes to the regulations to codify our existing policies in the Hospital IQR Program: (1) At 42 CFR 412.140(c)(2)(ii), that a separate submission deadline of April 1 following the end of the reporting calendar year in which the extraordinary circumstance occurred and applies to a hospital that wishes to request an extraordinary circumstances exception with respect to the reporting of electronic clinical quality measure data (81 FR 57182); (2) at 42 CFR 412.140(c)(2)(ii), that at the discretion of CMS, an exception may be granted to a hospital if a systemic problem arises with CMS data collection systems which directly affected the ability of a hospital to submit data (78 FR 50837), and that CMS may also grant exceptions to hospitals that have not requested them if an extraordinary circumstance affects an entire region or locale (76 FR 51651).

We are inviting public comments on these proposals as discussed above.

B. PPS-Exempt Cancer Hospital Quality Reporting (PCHQR) Program

1. Background

Section 3005 of the Affordable Care Act added new sections 1866(a)(1)(W) and (k) to the Act. Section 1866(k) of the Act establishes a quality reporting program for hospitals described in section 1886(d)(1)(A)(v) of the Act (referred to as “PPS-Exempt Hospitals” or “PCHs”) that specifically
applies to PCHs that meet the requirements under 42 CFR 412.23(f). Section 1866(k)(1) of the Act states that, for FY 2014 and each subsequent fiscal year, a PCH must submit data to the Secretary in accordance with section 1866(k)(2) of the Act with respect to such fiscal year. For additional background information, including previously finalized measures and other policies for the PCHQR Program, we refer readers to the following final rules: FY 2013 IPPS/LTCH PPS final rule (77 FR 53556 through 53561); the FY 2014 IPPS/LTCH PPS final rule (76 FR 50838 through 50846); the FY 2015 IPPS/LTCH PPS final rule (79 FR 50277 through 50288); the FY 2016 IPPS/LTCH PPS final rule (80 FR 49713 through 49723); and the FY 2017 IPPS/LTCH PPS final rule (81 FR 57182 through 57193).

2. Criteria for Removal and Retention of PCHQR Program Measures

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57182 through 57183), we adopted policies for measure retention and removal. We generally retain measures from the previous year’s PCHQR Program measure set for subsequent years’ measure sets, except when we specifically propose to remove or replace a measure. We adopted the following measure removal criteria for the PCHQR Program, which are based on criteria established in the Hospital IQR Program (80 FR 49641 through 49642):

- Measure performance among PCHs is so high and unvarying that meaningful distinctions and improvements in performance can no longer be made (“topped-out” measures);
- A measure does not align with current clinical guidelines or practice;
- The availability of a more broadly applicable measure (across settings or populations) or the availability of a measure that is more proximal in time to desired patient outcomes for the particular topic;
- Performance or improvement on a measure does not result in better patient outcomes;
- The availability of a measure that is more strongly associated with desired patient outcomes for the particular topic;
- Collection or public reporting of a measure leads to negative unintended consequences other than patient harm; and
- It is not feasible to implement the measure specifications.

For the purposes of considering measures for removal from the program, we consider a measure to be “topped-out” if there is statistically indistinguishable performance at the 75th and 90th percentiles and the truncated coefficient of variation is less than or equal to 0.10.

However, we recognized that there are times when measures may meet some of the outlined criteria for removal from the program, but continue to bring value to the program. Therefore, we adopted the following criteria for consideration in determining whether to retain a measure in the PCHQR Program, which also are based on criteria established in the Hospital IQR Program (80 FR 49641 through 49642):

- Measure aligns with other CMS and HHS policy goals;
- Measure aligns with other CMS programs, including other quality reporting programs; and
- Measure supports efforts to move PCHs towards reporting electronic measures.

We are not proposing any changes to these policies in this proposed rule.

3. Retention and Proposed Removal of Previously Finalized Quality Measures for PCHs Beginning With the FY 2020 Program Year

a. Background

In the FY 2013 IPPS/LTCH PPS final rule (77 FR 53556 through 53561), we finalized five quality measures for the FY 2014 program year and subsequent years. In the FY 2014 IPPS/LTCH PPS final rule (76 FR 50837 through 50847), we finalized one new quality measure for the FY 2015 program year and subsequent years and 12 new quality measures for the FY 2016 program year and subsequent years. In the FY 2015 IPPS/LTCH PPS final rule (79 FR 50278 through 50280), we finalized one new quality measure for the FY 2017 program year and subsequent years. In the FY 2016 IPPS/LTCH PPS final rule (80 FR 49713 through 49719), we finalized three new CDC NHSN measures for the FY 2018 program year and subsequent years, and finalized the removal of six previously finalized measures for fourth quarter (Q4) 2015 discharges and subsequent years. In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57183 through 57184), FY 2019 program year, we finalized one additional quality measure and updated the Oncology: Radiation Dose Limits to Normal Tissues (NQF #0382) measure.

We refer readers to the final rules referenced in section IX.B.1. of the preamble of this proposed rule for more information regarding these previously finalized measures.

b. Proposed Removal of Measures From the PCHQR Program Beginning With the FY 2020 Program Year

Based on a review of the above criteria, we are proposing to remove the following clinical process/cancer specific treatment measures from the PCHQR Program beginning with the FY 2020 program year because they are topped-out:

- Adjuvant Chemotherapy is Considered or Administered Within 4 Months (120 Days) of Diagnosis to Patients Under the Age of 80 with AJCC II (Lymph Node Positive) Colon Cancer (PCH–01/NQF #0223);
- Combination Chemotherapy is Considered or Administered Within 4 Months (120 Days) of Diagnosis for Women Under 70 with AJCC T1c, or Stage II or III Hormone Receptor Negative Breast Cancer (PCH–02/NQF #0559); and
- Adjuvant Hormonal Therapy (PCH–03/NQF #0220).

We first adopted these three Clinical Process/Cancer Specific Treatment Measures for the FY 2014 program year in the FY 2013 IPPS/LTCH PPS final rule (77 FR 53556 through 53561). We refer readers to that rule for a detailed discussion of the measures. However, based on an analysis of data from January 1, 2014 through September 30, 2015, we have determined that these three measures meet our topped-out criteria. This analysis, performed by the HCQIS Reports and Analytics Team, evaluated data sets provided from Program Data Management and calculated the 5th, 10th, 25th, 50th, 75th, 90th, and 95th percentiles of national facility performance for each measure. For measures where higher values indicate better performance, the percent relative difference (PRD) between the 75th and 90th percentiles were obtained by taking their absolute difference divided by the average of their values and result multiplied by 100. To calculate the truncated coefficient of variation (TCV), the lowest 5 percent and the highest 5 percent of hospital rates were discarded before calculating the mean and standard deviation for reach measure.

The following criteria were applied to the results:

- For measures ranging from 0–100 percent, with 100 percent being best, national measure data for the 75th and 90th percentiles have a relative difference of <=5 percent, or for measures ranging from 0–100 percent, with 100 percent being the best, performance achieved by the median hospital is >=95 percent, and national
measure data have a truncated coefficient of variation <=0.10.

• For measures ranging from 0–100 percent, with 0 percent being best, national measure data for the complement of the 10th and 25th percentiles have a relative difference of <=5 percent, or for measures ranging from 0–100 percent, with 0% being best, national measure data for the median hospital is <=5 percent, or for other measures with a low number indicating good performance, national measure data for the 10th and 25th percentiles have a relative difference of <=5 percent, and national measure data have a truncated coefficient of variation <=0.10.

The results for 2014 and 2015 are set out in the tables below.

### TOPPED-OUT ANALYSIS RESULTS FOR PCHQR MEASURES (2014)

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<th>Mean</th>
<th>Median</th>
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<th>Relative difference (%)</th>
<th>TCV</th>
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### TOPPED-OUT ANALYSIS RESULTS FOR PCHQR MEASURES (2015)

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Based on this analysis, we have concluded that these three measures are topped out and, as discussed below, we believe that collecting PCH data on these measures does not further program goals.

We believe that continuing to collect PCH data on these measures does not further program goals of improving quality, given that measure performance is so high and unvarying that meaningful distinctions and improvements in performance can no longer be made. We believe that these measures also do not meet the criteria for retention of an otherwise topped-out measure, as they do not align with other HHS and CMS policy goals, such as moving toward outcome measures; do not align with other CMS programs; and do not support the movement to electronic clinical quality measures due to the chart extraction required to collect the data for these measures. If we determine at a subsequent point in the future that hospital adherence to these practices has unacceptably declined, we may propose to readopt these measures in future rulemaking.

We are inviting public comment on our proposal to remove these three measures from the PCHQR Program beginning with the FY 2020 program year.

4. Proposed New Quality Measures Beginning With the FY 2020 Program Year

a. Considerations in the Selection of Quality Measures

In the FY 2013 IPPS/LTCH PPS final rule (77 FR 53556), the FY 2014 IPPS/LTCH PPS final rule (78 FR 50837 through 50838), and the FY 2015 IPPS/LTCH PPS final rule (79 FR 50278), we indicated that we take a number of principles into consideration when developing and selecting measures for the PCHQR Program, and that many of these principles are modeled on those we use for measure development and selection under the Hospital IQR Program. In this proposed rule, we are not proposing any changes to the principles we consider when developing and selecting measures for the PCHQR Program.

Section 1866(k)(3)(A) of the Act requires that any measure specified by the Secretary must have been endorsed by the entity with a contract under section 1890(a) of the Act (the NQF is the entity that currently holds this contract). Section 1866(k)(3)(B) of the Act provides an exception under which, in the case of a specified area or medical topic determined appropriate by the Secretary for which a feasible and practical measure has not been endorsed by the entity with a contract under section 1890(a) of the Act, the Secretary may specify a measure that is not so endorsed as long as due consideration is given to measures that have been endorsed or adopted by a consensus organization.

Using the principles for measure selection in the PCHQR Program, we are proposing four new measures, described below.

b. Proposed New Quality Measures Beginning With the FY 2020 Program Year

For the FY 2020 PCHQR program year, we are proposing to adopt two intermediate clinical outcome quality measures. These measures meet the requirement under section 1866(k)(3)(A) of the Act that measures specified for the PCHQR Program generally be endorsed by the entity with a contract under section 1890(a) of the Act (currently the NQF). Although there is no financial incentive or penalty associated with the PCHQR Program, we encourage participation to further the goal of improving the quality of care for the PCH patient population. The proposed measures are:

• Proportion of Patients Who Died from Cancer Receiving Chemotherapy in the Last 14 Days of Life (NQF #0210);
• Proportion of Patients Who Died from Cancer Admitted to the ICU in the Last 30 Days of Life (NQF #0213);
• Proportion of Patients Who Died from Cancer Not Admitted to Hospice (NQF #0215); and
• Proportion of Patients Who Died from Cancer Admitted to Hospice for Less Than Three Days (NQF #0216).

In compliance with section 1890A(a)[2] of the Act, the proposed
measures were included on a publicly available document entitled “List of Measures under Consideration for December 1, 2016.” 249 A list of quality and efficiency measures under consideration for use in various Medicare programs, and were reviewed by the MAP Hospital Workgroup. The MAP Hospital Workgroup supported the inclusion of these measures in the PCHQR Program in final recommendations it made in its February 2017 report to HHS and CMS for 2016 to 2017. 250 Additional details on MAP discussions of these measures may be found in the “MAP Pre-Rulemaking Report: 2016 Recommendations on Measures Under Consideration by HHS,” with additional discussion in the “MAP 2017 Considerations for Implementing Measures in Federal Programs: Hospitals (Draft Report).” 251 The sections below outline our rationale for proposing these measures.

(1) Background

The quality of end-of-life care has been identified by the NQF as an area of care that continues to need improvement. 252 End-of-life care may be defined as “comprehensive care that anticipates, prevents, and alleviates suffering. 253 and may exclude palliative care. Palliative care is generally defined as multi-faceted, holistic care that anticipates, prevents, and alleviates suffering. 254 Both palliative and end-of-life care can be provided when a patient is receiving hospice services, but it is not necessary to be admitted to hospice to receive such care. The NQF notes that hospice is both a type of care team and a care philosophy, and is intended to enable patients to prepare for death while living as fully as possible. 255 The Institute of Medicine of the National Academies (IOM) has noted that while clinicians are encouraged to counsel patients about palliative care, which better chances of maintaining a high quality of life when dying, “too few patients and families receive this help in a timely manner.” 256 In the same report, the IOM proposed a number of core components of quality end-of-life care. These proposals included offering a referral to hospice if a patient “has a prognosis of 6 months or less” and regular revision of a patient’s care plan to address the patient’s changing needs, as well as the changing needs of the family. 257

In addition to all of the quality of care benefits of end-of-life care to patients and caregivers, there are financial cost benefits as well. In its Technical Report on palliative and end-of-life care, the NQF cited research indicating that the use of palliative care, including end-of-life care, results in various positive outcomes, including a reduction of costs. 258 For example, one study evaluated the impact of hospice enrollment at different time periods on Medicare expenditures, and found that regardless of when a patient was enrolled in hospice, such patients’ subsequent Medicare costs were significantly lowered. 259

Despite the benefits attributed to the use of palliative and end-of-life services and the increase in their availability, the NQF and others have noted that such services remain underutilized. By proposing to include two process measures and two intermediate clinical outcome measures related to end-of-life care in the PCHQR Program, our intent is to assess the quality of end-of-life care provided to patients in the PCH setting.

We recognize that these measures may also be used in the broader population of all hospitals providing cancer care; therefore, as discussed in section IX.A.9.b. of the preamble of this proposed rule, we are inviting public comment on the future inclusion of these measures in the Hospital IQR Program. These four measures are described in more detail below.

(2) Proportion of Patients Who Died From Cancer Receiving Chemotherapy in the Last 14 Days of Life (EOL-Chemo) Measure (NQF #0210)

Chemotherapy is typically used to treat cancer, but in patients with incurable cancer it may also be used with the goal of easing symptoms and improving survival. 260 One study estimated that 6.2 percent of cancer patients continue receiving chemotherapy close to the end of their lives (defined as within 2 weeks of death). 261 However, studies have shown that administering palliative chemotherapy to terminally ill cancer patients may not be beneficial, as it may be associated with higher rates of interventions such as cardiopulmonary resuscitation in the last week of life without any difference in survival. 262 Such patients may also be more likely to die in the intensive care unit (ICU), and less likely to die either at home or in the place where they had expressed preference to die. 263 In addition, research has shown that some patients may receive chemotherapy for treatment instead of palliative care at the end of life, even when treatment has been determined to be unnecessary. 264 While the impetus for continuing treatment may vary from case to case, the available evidence indicates continuing to receive chemotherapy—for palliation

259 Kelley AS et al., Hospice Enrollment Saves Money for Medicare and Improves Care Quality Across a Number of Different Lengths-of-Stay, Health Affairs (March 2013)12/3:552-561.

260 Wright A et al., Associations Between Palliative Chemotherapy and Adult Cancer Patients’ End of Life Care and Place of Death: Prospective Cohort Study, BMJ 2014;348:g1219.

261 Wright A et al., Associations Between Palliative Chemotherapy and Adult Cancer Patients’ End of Life Care and Place of Death: Prospective Cohort Study, BMJ 2014;348:g1219.

262 Wright A et al., Associations Between Palliative Chemotherapy and Adult Cancer Patients’ End of Life Care and Place of Death: Prospective Cohort Study, BMJ 2014;348:g1219.

263 Mack JW et al., Patient Beliefs that Chemotherapy May be Curotive and Care Received at the End of Life Among Patients with Metastatic Lung and Colorectal Cancer, Cancer (June 1, 2015)121:11:1891-1897.

264 Mack JW et al., Patient Beliefs that Chemotherapy May be Curotive and Care Received at the End of Life Among Patients with Metastatic Lung and Colorectal Cancer, Cancer (June 1, 2015)121:11:1891-1897.
or treatment—toward the end of a patient’s illness is associated with increased hospitalization and may be associated decreased experience of care.

Researchers have also observed that patients receiving chemotherapy late into the course of a terminal illness tended to be referred to hospice later, resulting in lower quality of life, distress for caregivers, and increased cost.266 They noted that their results could suggest that either less use chemotherapy at the end of life or more frequent end-of-life discussions could improve the quality of those patients’ end-of-life care.267 Another study of early engagement in palliative care in patients diagnosed with metastatic lung cancer found that patients who received palliative care and less chemotherapy survived longer, in addition to experiencing improvement in quality of life.268 In this study, palliative care was integrated into standard oncologic care, and included an assessment of physical and psychosocial symptoms as well as care decision assistance.269 Results from this study showed significantly higher quality of life in the patient cohort receiving palliative care compared to those receiving only the standard oncologic care.270 In particular, clinically meaningful improvements in quality of life and mood were noted.271

The proposed EOL-Chemo measure addresses the NQS Communication and Care Coordination and Affordable Care domains, and aligns with the CMS Quality Strategy goals of strengthening person and family engagement as partners in their care, and promoting effective communication and coordination of care. The proposed measure is a process measure that evaluates the proportion of patients who died from cancer who received chemotherapy in the last 14 days of life.

Similar to the other three end-of-life measures we are proposing, this proposed measure seeks to assess the use of chemotherapy at the end-of-life, a practice advanced with the intent to alleviate disease symptoms but which has been shown to also be associated with reduced quality of life and increased costs. This measure was revised for use in the Merit-based Incentive Payment System (MIPS) in the FY 2017 MIPS final rule with comment period (81 FR 77672). By introducing this measure here, we are seeking to evaluate how often chemotherapy is administered near the end of life in PCUs.

The proposed EOL-Chemo measure cohort includes all Medicare beneficiaries who died of cancer and who received chemotherapy at a PCH within the last 14 days of their lives. The proposed measure uses Medicare administrative claims data to derive the numerator and denominator. The numerator for this measure is defined as cancer patients who received chemotherapy (regardless of whether for treatment or palliative purposes) in the last fourteen days of life. The denominator is defined as patients who died from cancer. Patients for whom numerator or denominator data cannot be identified will not be included in the calculation. The measure specifications contain no exclusions, risk adjustments or risk stratifications because the measure is intended to evaluate the quality of care provided to all cancer patients at the end of life. The measure will be calculated as the numerator divided by the denominator. Measure specifications for the proposed EOL-Chemo measure can be accessed on the NQF’s Web site at: http://www.qualityforum.org/Publications/2016/12/Palliative_care_and_End-of-Life_Care_2015-2016.aspx.

We are inviting public comment on our proposal to adopt the Proportion of Patients Who Died from Cancer Receiving Chemotherapy in the Last 14 Days of Life (NQF #0210) measure for the FY 2020 program year and subsequent years.

(3) Proportion of Patients Who Died From Cancer Admitted to the ICU in the Last 30 Days of Life (EOL–ICU) Measure (NQF #0213)

A number of research studies have determined that cancer care can become more aggressive at the end of life, which can result in a lower quality of care and lower quality of life.272 Care defined as “aggressive” may include the “possible misuse of treatment resulting in high rates of emergency room visits, hospitalization, or ICU stays for terminal patients” in addition to overuse of chemotherapy close to death and the underuse of hospice.273 In a retrospective study of patients with advanced lung cancer, researchers found that between 1993 and 2002, the number of patients being admitted to the ICU near death increased, and while in the ICU, one in four of those patients received mechanical ventilation, despite the likelihood that neither intervention would necessarily have effect on the advanced cancer.274 In this study, two-thirds of the patients died within a month of their admission to the ICU, which the authors interpreted as demonstrating that ICU admission in the context of advanced lung cancer was potentially ineffective.275 The authors noted other studies that showed that in-hospital mortality during ICU admissions exact a toll on patients and families in terms of “financial cost, emotional burden, and failed expectations.”276 The impact of ICU admission at the end of life is also observed amongst caregivers, who report excellent end-of-life care less often for patients admitted to the ICU within 30 days of death compared to those who are not.277

Patients who are not admitted to the ICU or involved in other aggressive mechanisms of care in their final week of life have been shown to experience a higher quality of life via less physical and emotional distress.278 Researchers have theorized that while patients who die at home are able to have care that focuses on symptom management and...
comfort; hospitals and ICUs focus instead on keeping the patient alive.279 ICU admission at the end of life is also costly,280 with ICU admissions identified as one of the “key drivers of resource use and expenditures.”281 Studies of claims data indicate that aggressiveness of care given to Medicare beneficiaries with cancer at the end of life continues to increase, with nearly 25 percent of Medicare expenditures in the last month of such beneficiaries’ lives, despite limited evidence that such an intervention improves patient outcomes.282

The proposed EOL–ICU measure addresses the NQS Communication and Care Coordination and Affordable Care domains, and addresses several CMS Quality Strategy goals: Making care safer by reducing harm caused in the delivery of care; strengthening person and family engagement as partners in their care; and promoting effective communication and coordination of care. The proposed EOL–ICU measure is an intermediate clinical outcome measure that assesses whether cancer patients were admitted to the ICU in the last 30 days of their lives. As with the other three proposed end-of-life measures discussed in section IX.B.4.b. of the preamble of this proposed rule, this proposed measure seeks to evaluate the end-of-life care provided to patients at PCHs. In particular, we seek to assess the frequency of end-of-life admissions to the ICU in this setting, as the research has shown that interventions provided in the ICU to patients with irreversible disease can be futile and may negatively impact patients’ quality of life. We recognize, however, that in some cases ICU admissions may be appropriate, and note that this measure broadly assesses how many patients are admitted to the ICU close to death, without excluding admissions for specific reasons.

The proposed EOL–ICU measure cohort includes Medicare beneficiaries who are PCH patients who died of cancer and who were admitted to the ICU within the last thirty days of their lives. This proposed measure uses Medicare administrative claims data to derive the numerator and denominator. The numerator for this measure is defined as the number of patients who died from cancer and who were admitted to the ICU in the last 30 days of life. The denominator is defined as patients who died from cancer. The measure specifications do not contain exclusions from the denominator and do not provide for risk adjustment or risk stratification in order to assess the quality of care provided to all cancer patients at the end of life. The rate of ICU admissions in the last 30 days of life will be calculated from the numerator divided by the denominator. Measure specifications for the proposed EOL–ICU measure can be accessed on the NQF’s Web site at: http://www.qualityforum.org/Publications/2016/12/Palliative_and_End-of-Life_Care_2015-2016.aspx.

We are inviting public comment on our proposal to adopt the Proportion of Patients Who Died From Cancer Admitted to the ICU in the Last 30 Days of Life (NQF #0213) measure for the FY 2020 program year and subsequent years.

(4) Proportion of Patients Who Died From Cancer Not Admitted to Hospice (EOL-Hospice) Measure (NQF #0215)

A number of research studies have determined that cancer care can become more aggressive at the end of life, which can result in a lower quality of care and lower quality of life.283 Such aggressive care has been identified to include the underutilization of hospice,284 which is either lack of referral or late referral to hospice services.285 Patients with advanced cancer who die while admitted to the hospital have been shown to have lower quality of life than those who die at home with hospice services.286

By contrast, studies have shown that cancer patients enrolled in hospice were hospitalized less frequently and received fewer procedures than those who were not receiving hospice care.287 In addition, cancer patients who were enrolled in hospice 5 to 8 weeks prior to their deaths demonstrated significant cost savings, with savings decreasing as the time period enrolled shortens.288 Researchers theorize that one reason aggressive or “injudicious” treatment occurs at the end of life is that end-of-life discussions are not being held with patients, and note that it is “the physician’s responsibility to counsel patients and their families and . . . focus on the need for effective palliative care as patients approach the end of life.”289

The proposed EOL-Hospice measure addresses the NQS Communication and Care Coordination and Affordable Care domains, as well as the CMS Quality Strategy goals of strengthening person and family engagement as partners in their care and promoting effective communication and coordination of care. The proposed measure is a process measure that assesses the proportion of patients who died from cancer who were not admitted to hospice. This measure evaluates whether or not patients were admitted to hospice, and then ties in to the following measure (EOL–3DH), which evaluates whether patients who were admitted to hospice were admitted to hospice late in the course of their illness, defined as within 3 days of their death. We discuss this proposed follow-on measure, EOL–3DH, in more detail in section IX.B.4.b.(5) of the preamble of this proposed rule. In summary, EOL-Hospice seeks to evaluate, simply, whether patients were admitted to hospice or not; the proposed follow-on measure EOL–3DH will then assess whether those patients admitted to hospice were admitted in a timely fashion to derive maximum benefit from hospice services. We do not expect PCHs to achieve perfect rates on the EOL-Hospice measure because we understand that some patients may refuse hospice, or that there may be additional intervening events or circumstances that impact whether or not a patient is admitted to hospice.

The proposed EOL-Hospice measure cohort includes Medicare beneficiaries who are PCH patients who died of cancer. The proposed measure uses
Medicare administrative claims data to derive the numerator and denominator. The numerator in this proposed measure is defined as the proportion of PCH patients not enrolled in hospice. The denominator is defined as patients who died from cancer. The measure specifications contain no denominator exclusions nor any risk adjustment or risk stratification. The proposed measure is calculated by dividing the numerator by the denominator. Measure specifications for the proposed EOL–Hospice measure can be accessed on the NQF’s Web site at: [http://www.qualityforum.org/Publications/2016/12/Palliative_and_End-of-Life_Care_2015-2016.aspx](http://www.qualityforum.org/Publications/2016/12/Palliative_and_End-of-Life_Care_2015-2016.aspx).

We are inviting public comment on our proposal to adopt the Proportion of Patients Who Died from Cancer Not Admitted to Hospice (NQF #0215) measure for the FY 2020 program year and subsequent years.

(5) Proportion of Patients Who Died From Cancer Admitted to Hospice for Less Than 3 Days (EOL–3DH) Measure (NQF #0216)

Older studies of patient cohorts from the mid-1990s have shown that, though there was an increasing trend to admit cancer patients to hospice, the number of patients admitted close to death was also increasing, about which the authors surmised that hospice care was not being used to mitigate symptoms but only to manage death.

Patients with cancer have been identified as the largest users of hospice, but are also the cohort with the highest rates of hospice stays of less than 3 days.

In one study involving cancer patients’ family members, patients’ loved ones were more likely to report that the patients received excellent end-of-life care when hospice was initiated earlier than three days prior to death.

The researchers indicated that enhancing counseling of patients and families and early referral to palliative care services could result in more “preference-sensitive care for patients” and overall improvement in the quality of care cancer patients receive at the end of life.

Because this and other research indicates that earlier discussion with patients about palliative care can positively impact the care received at the end of life, including timely admission to hospice, we believe including the proposed EOL–3DH measure in the measure set will incentivize timely discussions and admissions to hospice within the PCH setting. We believe that the emphasis on timely admission to hospice may lead to improving the quality of care for cancer patients at PCHs.

The proposed EOL–3DH measure addresses the NQS Communication and Care Coordination domain. It also addresses two CMS Quality Strategy goals: Strengthening person and family engagement as partners in their care and promoting effective communication and coordination of care. The proposed EOL–3DH measure is an intermediate clinical outcome measure that assesses the proportion of patients who died from cancer who were admitted to hospice late in the course of their illness, within 3 days of their death. The measure ties in to the proposed process measure (EOL-Hospital) we discuss in section IX.B.4.b(4) of the preamble of this proposed rule, above, and assesses whether, if patients were admitted to hospice, they were admitted prior to or when death was immediately imminent. As discussed, research has shown that the longer patients receive hospice services before the end of life, the more improvements in their quality of life and mood are observed.

The proposed EOL–3DH measure cohort includes Medicare beneficiaries who are PCH patients who died of cancer and were admitted to hospice within the last 3 days of their lives. The proposed measure uses Medicare administrative claims data to derive the numerator and denominator. The numerator is defined as the number of patients who died from cancer and spent fewer than 3 days in hospice. The denominator is defined as the number of patients who died from cancer who were admitted to hospice. There are no exclusions from the denominator in the measure specifications, nor risk adjustment or risk stratification, because the goal of the measure is to assess the quality of care provided to all cancer patients at the end of life. Measure specifications for the proposed EOL–3DH measure can be accessed on the NQF’s Web site at: [http://www.qualityforum.org/Publications/2016/12/Palliative_and_End-of-Life_Care_2015-2016.aspx](http://www.qualityforum.org/Publications/2016/12/Palliative_and_End-of-Life_Care_2015-2016.aspx).

We are inviting public comment on our proposal to adopt the Proportion of Patients Who Died from Cancer Admitted to Hospice for Less Than 3 Days (NQF #0216) measure for the FY 2020 program year and subsequent years.

c. Summary of Previously Finalized and Newly Proposed PCHQR Program Measures for the FY 2020 Program Year and Subsequent Years

In summary, the previously finalized and newly proposed measures for the PCHQR Program for the FY 2020 program year and subsequent years are listed in the table below.

### PREVIOUSLY FINALIZED AND NEWLY PROPOSED PCHQR MEASURES FOR THE FY 2020 PROGRAM YEAR AND SUBSEQUENT YEARS

<table>
<thead>
<tr>
<th>Short name</th>
<th>NQF No.</th>
<th>Measure name</th>
</tr>
</thead>
<tbody>
<tr>
<td>SSI</td>
<td>0753</td>
<td>American College of Surgeons—Centers for Disease Control and Prevention (ACS–CDC) Harmonized Procedure Specific Surgical Site Infection (SSI) Outcome Measure [currently includes SSIs following Colon Surgery and Abdominal Hysterectomy Surgery].</td>
</tr>
<tr>
<td>CDI</td>
<td>1717</td>
<td>National Healthcare Safety Network (NHSN) Facility-wide Inpatient Hospital-onset <em>Clostridium difficile</em> Infection (CDI) Outcome Measure.</td>
</tr>
</tbody>
</table>

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291 Obermeyer Z et al., Association Between the Medicare Hospice Benefit and Health Care Utilization and Costs for Patients with Prostate Cancer, JAMA (2014)312:18;1888-1896.

292 Wright AA et al., Family Perspectives on Aggressive Cancer Care Near the End of Life, JAMA (2016)315:3;284–292.

293 Wright AA et al., Family Perspectives on Aggressive Cancer Care Near the End of Life, JAMA (2016)315:3;284–292.
5. Accounting for Social Risk Factors in the PCHQR Program

We understand that social risk factors such as income, education, race and ethnicity, employment, disability, community resources, and social support (certain factors of which are also sometimes referred to as socioeconomic status (SES) factors or socio-demographic status (SDS) factors) play a major role in health. One of our core objectives is to improve beneficiary outcomes including reducing health disparities, and we want to ensure that all beneficiaries, including those with social risk factors, receive high quality care. In addition, we seek to ensure that the quality of care furnished by providers and suppliers is assessed as fairly as possible under our programs while ensuring that beneficiaries have adequate access to excellent care.

We have been reviewing reports prepared by the Office of the Assistant Secretary for Planning and Evaluation (ASPE) and the National Academies of Sciences, Engineering, and Medicine on the issue of measuring and accounting for social risk factors in CMS’ value-based purchasing and quality reporting programs, and considering options on how to address the issue in these programs. On December 21, 2016, ASPE submitted a Report to Congress on a study it was required to conduct under section 2(d) of the Improving Medicare Post-Acute Care Transformation (IMPACT) Act of 2014. The study analyzed the effects of certain social risk factors of Medicare beneficiaries on quality measures and measures of resource use used in one or more of nine Medicare value-based purchasing programs. The report also included considerations for strategies to account for social risk factors in these programs. In a January 10, 2017 report released by the National Academies of Sciences, Engineering, and Medicine, that body provided various potential methods for measuring and accounting for social risk factors in our programs.
for social risk factors, including stratified public reporting.\textsuperscript{296} As noted in the FY 2017 IPPS/LTCH PPS final rule, the NQF has undertaken a 2-year trial period in which new measures, measures undergoing maintenance review, and measures endorsed with the condition that they enter the trial period can be assessed to determine whether risk adjustment for selected social risk factors is appropriate for these measures. This trial entails temporarily allowing inclusion of social risk factors in the risk-adjustment approach for these measures. At the conclusion of the trial, NQF will issue recommendations on the future inclusion of social risk factors in risk adjustment for these quality measures, and we will closely review their findings.

As we continue to consider the analyses and recommendations from these reports and await the results of the NQF pilot on risk adjustment for quality measures, we are continuing to work with stakeholders in this process. As we have previously communicated, we are concerned about holding providers to different standards for the outcomes of their patients with social risk factors because we do not want to mask potential disparities or minimize incentives to improve the outcomes for disadvantaged populations. Keeping this concern in mind, while we sought input on this topic previously, we continue to seek public comment on whether we should account for social risk factors in the PCHQR Program, and if so, what method or combination of methods would be most appropriate for accounting for social risk factors. Examples of methods include: Confidential reporting to providers of measure rates stratified by social risk factors; public reporting of stratified measure rates; and potential risk adjustment of a particular measure as appropriate based on data and evidence.

In addition, we are also seeking public comment on which social risk factors might be most appropriate for reporting stratified measure scores and/or potential risk adjustment of a particular measure. Examples of social risk factors include, but are not limited to, dual eligibility/low-income subsidy, race and ethnicity, and geographic area of residence. We are seeking comments on which of these factors, including current data sources where this information would be available, could be used alone or in combination, and whether other data should be collected to better capture the effects of social risk. We will take commenters’ input into consideration as we continue to assess the appropriateness and feasibility of accounting for social risk factors in the PCHQR Program. We note that any such changes would be proposed through future notice-and-comment rulemaking.

We look forward to working with stakeholders as we consider the issue of accounting for social risk factors and reducing health disparities in CMS programs. Of note, implementing any of the above methods would be taken into consideration in the context of how this and other CMS programs operate (for example, data submission methods, availability of data, statistical considerations relating to reliability of data calculations, among others), so we also welcome comment on operational considerations. CMS is committed to ensuring that its beneficiaries have access to and receive excellent care, and that the quality of care furnished by providers and suppliers is assessed fairly in CMS programs.

6. Possible New Quality Measure Topics for Future Years

a. Background

We discussed future quality measure topics and quality measure domain areas in the FY 2015 IPPS/LTCH PPS final rule (79 FR 50280), the FY 2016 IPPS/LTCH PPS final rule (80 FR4979), and the FY 2017 IPPS/LTCH PPS final rule (81 FR 25211). Specifically, we discussed public comment and suggestions for measure topics addressing the following CMS Quality Strategy domains: (1) Making care affordable; (2) communication and care coordination; and (3) working with communities to promote best practices of healthy living. We welcome public comment and specific suggestions for measure topics that we should consider for future rulemaking, including considerations related to risk adjustment and the inclusion of social risk factors in risk adjustment for any individual performance measures.

We are also seeking public comment on six measures for potential future inclusion in the PCHQR Program:

- Localized Prostate Cancer: Vitality;
- Localized Prostate Cancer: Urinary Incontinence;
- Localized Prostate Cancer: Urinary Frequency, Obstruction, and/or Irritation;
- Localized Prostate Cancer: Sexual Function;
- Localized Prostate Cancer: Bowel Function; and
- 30-Day Unplanned Readmissions for Cancer Patients.

These measures are discussed in more detail below.

b. Localized Prostate Cancer: Vitality; Localized Prostate Cancer: Urinary Incontinence; Localized Prostate Cancer: Urinary Frequency, Obstruction, and/or Irritation; Localized Prostate Cancer: Sexual Function; and Localized Prostate Cancer: Bowel Function

The Localized Prostate Cancer measures are five related, patient-reported outcome measures drawn from the Expanded Prostate Inventory Composite (EPIC), which is a survey intended to gather input from patients on their experience. The survey questions are intended to be administered to all non-metastatic prostate cancer patients undergoing radiation or surgical treatment for prostate cancer at the reporting facility (denominator); the numerator is patients with clinically significant changes in each of the listed areas from baseline to follow-up. The goal of the measurement is to identify issues of variation, suboptimal performance, and disparities in care. This measurement aligns with recent initiatives to include patient-reported outcomes and experience of care into quality reporting programs, as well as to incorporate more outcome measures generally. Patient-centered experience measures are also a component of the 2016 CMS Quality Strategy, which emphasizes patient-centered care by rating patient experience as a means for empowering patients and improving the quality of their care and care experience.\textsuperscript{297}

These measures were included on the publicly available document entitled “List of Measures under Consideration for December 1, 2016”\textsuperscript{298} but were not reviewed by the MAP. We anticipate that they will be included on a future list of measures under consideration for MAP review. For further information on these measures, we refer readers to the discussion from the Measures Application Partnership’s Hospital Workgroup Discussion at: http://public.qualityforum.org/MAP/MAP%20Hospital%20Workgroup/2016-2017%20Hospital%20MAP/MAP_Hospital_Workgroup_Discussion_...\textsuperscript{297}\textsuperscript{CMS Quality Strategy 2016. Available at: https://www.cms.gov/medicare/quality-summariespatient-assessment-instruments/qualitysummariespatient-assessment-instruments.pdf.}

The 30-Day Unplanned Readmissions for Cancer Patients measure would measure the number of hospital-specific 30-day unscheduled and potentially avoidable readmissions following hospitalization among diagnosed malignant cancer patients. The measure numerator is the total number of unscheduled readmissions within 30 days of index admission. The measure denominator is total PCH admissions within the reporting year for patients, aged 18 years or older, who were discharged alive from the facility with an active malignant cancer diagnosis.

For further information on this measure, we refer readers to the AHRQ National Quality Measure Clearinghouse at: https://www.qualitymeasures.ahrq.gov/summaries/summary/50490/cancer-30day-unplanned-readmission-rate-for-cancer-patients. We are requesting public comment on the possible inclusion of this measure in future years of the program.

### 7. Maintenance of Technical Specifications for Quality Measures

We maintain technical specifications for the PCHQR Program measures, and we periodically update those specifications. The specifications may be found on the QualityNet Web site at: https://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=1228774479863.

In the FY 2015 IPPS/LTCH PPS final rule (79 FR 50281), we adopted a policy under which we use a subregulatory process to make nonsubstantive updates to measures used for the PCHQR Program. We are not proposing any changes to this policy in this proposed rule.

### PREVIOUSLY FINALIZED PUBLIC DISPLAY REQUIREMENTS

<table>
<thead>
<tr>
<th>Measures</th>
<th>Public reporting</th>
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<tr>
<td>c. 30-Day Unplanned Readmissions for Cancer Patients</td>
<td></td>
</tr>
<tr>
<td>Adjuvant Chemotherapy Is Considered or Administered Within 4 Months (120 days) of Diagnosis to Patients Under the Age of 80 with AJCC III (lymph node positive) Colon Cancer (NQF #0223) *</td>
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</tr>
<tr>
<td>Combination Chemotherapy Is Considered or Administered Within 4 Months (120 days) of Diagnosis for Women Under 70 with AJCC T1cN0MO, or Stage IB—III Hormone Receptor Negative Breast Cancer (NQF #0559) *</td>
<td></td>
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<tr>
<td>Oncology: Radiation Dose Limits to Normal Tissues (NQF #0382) *</td>
<td></td>
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<tr>
<td>Oncology: Plan of Care for Pain—Medical Oncology and Radiation Oncology (NQF #0383)</td>
<td>2014 and subsequent years.</td>
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<tr>
<td>Oncology: Medical and Radiation—Pain Intensity Quantified (NQF #0384)</td>
<td>2015 and subsequent years.</td>
</tr>
<tr>
<td>Prostate Cancer: Adjuvant Hormonal Therapy for High Risk Prostate Cancer Patients (NQF #0390)</td>
<td></td>
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<tr>
<td>Prostate Cancer: Avoidance of Overuse of Bone Scan for Staging Low Risk Prostate Cancer Patients (NQF #0389)</td>
<td></td>
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<tr>
<td>HCAHPS (NQF #0166)</td>
<td>2016 and subsequent years.</td>
</tr>
<tr>
<td>CLABSI (NQF #0139) **</td>
<td>Deferred.</td>
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<tr>
<td>CAUTI (NQF #0138) **</td>
<td>Beginning at the first opportunity in 2017 and for subsequent years.</td>
</tr>
<tr>
<td>External Beam Radiotherapy for Bone Metastases (NQF #1822) ***</td>
<td></td>
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</tbody>
</table>

* Update newly finalized for display for the FY 2019 program year and subsequent years in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57192)—expanded cohort will be displayed as soon as feasible.

** Measure newly finalized for public display in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57192).

*** Measure newly finalized for public display in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57192).

a. Background

As we strive to publicly display data as soon as possible on a CMS Web site, in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57191 through 57192), we finalized an update to our public display policies. We believe it is best to not specify in rulemaking the exact timeframe during the year for publication as doing so may prevent earlier publication. Therefore, we finalized our policy to make these data available as soon as it is feasible during the year, starting with the first year for which we are publishing data for each measure. We will continue to propose in rulemaking the first year for which we intend to publish data for each measure. We intend to make the data available on at least a yearly basis.

As stated above, we are required to give PCHs an opportunity to review their data before the data are made public. Because we will make the data for this program available as soon as possible, and the timeframe for this publication may change year to year, we will not propose to specify in rulemaking the exact dates for review.

However, in that final rule, we stated that the time period for review would be approximately 30 days in length. We will announce the exact timeframes on a CMS Web site and/or on our applicable listservs. We are not proposing any changes to this policy in this proposed rule.

b. Deferment of Public Display of Two Measures

In the FY 2015 IPPS/LTCH PPS final rule (79 FR 50281 through 50282), we finalized public display of the CLABSI...
and CAUTI measures beginning no later than 2017 and subsequent years. However, in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57192), we finalized a proposal to continue to defer public reporting of the CLABSI and CAUTI measures pending ongoing collaboration with the CDC to identify an appropriate timeframe for public reporting and the analytic methods that will be used to summarize the CLABSI and CAUTI data for public reporting purposes. We continue to collaborate with the CDC on these issues and continue to defer the public reporting of these two measures accordingly.

9. Form, Manner, and Timing of Data Submission
   a. Background

   Section 1866(k)(2) of the Act requires that, beginning with the FY 2014 PCHQR program year, each PCH must submit to the Secretary data on quality measures specified under section 1866(k)(3) of the Act in a form and manner, and at a time, as specified by the Secretary. There are no financial incentives or penalties associated with the PCHQR Program.

   Data submission requirements and deadlines for the PCHQR Program are generally posted on the QualityNet Web site at: http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetaPublic%2FPage%2FQNetTier3&cid=1228772864228.

   In this proposed rule, we are not proposing any changes to previously finalized data submission requirements.

   b. Proposed Reporting Requirements for the Proposed New Measures

   As further described above, we are proposing four new measures beginning with the FY 2020 program year: Proportion of Patients Who Died from Cancer Receiving Chemotherapy in the Last 14 Days of Life (NQF #0210); Proportion of Patients Who Died from Cancer Admitted to the ICU in the Last 30 Days of Life (NQF #0213); Proportion of Patients Who Died from Cancer Admitted to Hospice for Less Than Three Days (NQF #0216). All four measures are claims-based measures. Therefore, there will be no data submission requirements for PCHs related to these measures. As these measures use Medicare administrative claims data, we are proposing to calculate these measures on a yearly basis. Specifically, we are proposing that the data collection period will be from July 1 from the year 3 years prior to the program year to June 30 from the year 2 years prior to the program year. Thus, for the FY 2020 program year, we would collect data from July 1, 2017 through June 30, 2018.

We are inviting public comment on this proposal.

10. Extraordinary Circumstances

   a. Background

In our experience with other quality reporting and performance programs, we have noted occasions when providers have been unable to submit required quality data due to extraordinary circumstances that are not within their control (for example, natural disasters). We do not wish to increase their burden unduly during these times. Therefore, in the FY 2014 IPPS/LTCH PPS final rule (78 FR 50848), we finalized our policy that, for the FY 2014 program year and subsequent years, PCHs may request and we may grant exceptions (formerly referred to as waivers) with respect to the reporting of required quality data when extraordinary circumstances beyond the control of the PCH warrant. The PCH may request a reporting extension or a complete exception from the requirement to submit quality data for one or more quarters. Under our current policy, PCHs can submit a request form to CMS with the following information:

- The PCH’s CCN;
- The PCH’s name;
- Contact information for the PCH’s CEO and any other designated personnel, including name, email address, telephone number, and mailing address (the address must be a physical address, not a post office box);
- The PCH’s reason for requesting an extension or exception;
- Evidence of the impact of extraordinary circumstances, including but not limited to photographs, newspaper and other media articles; and
- A date when the PCH will again be able to submit PCHQR Program data, and a justification for the proposed date.

In addition, we finalized that the form must be signed by the PCH’s CEO or designee and submitted within 30 days of the date that the extraordinary circumstances occurred. Lastly, we finalized that following the receipt of the request form, we would: (1) provide a written acknowledgement; and (2) provide a formal response notifying the PCH of our decision.

We also clarified that the above policy does not preclude us from granting exceptions (including extensions) to PCHs that have not requested them when we determine that an extraordinary circumstance has affected an entire region or locale. We stated that if we make the determination to grant such an exception, we would communicate this decision through routine communication channels.

   b. Proposed Modifications to the ECE Policy

We are proposing to modify the ECE policy for the PCHQR Program by: (1) Extending the deadline for a PCH to submit a request for an extension or exception from 30 days following the date that the extraordinary circumstance occurred to 90 days following the date that the extraordinary circumstance occurred; and (2) allowing CMS to grant an exception or extension due to CMS data system issues which affect data submission. These proposed modifications will better align our ECE policy with that adopted for the Hospital IQR Program (76 FR 51651 through 51652, 76 FR 50836 through 50837, and 81 FR 57181 through 57182), the Hospital OQR Program (77 FR 68489 and 81 FR 79795), as well as other quality reporting programs that already have such policies in place or are proposing to modify their policies to achieve alignment. We are proposing that these modifications would apply beginning in FY 2018 as related to extraordinary circumstances that occur on or after October 1, 2017.

We also believe that it is important for facilities to receive timely feedback regarding the status of ECE requests. We strive to complete our review of each ECE request as quickly as possible. However, we recognize that the number of requests we receive, and the complexity of the information provided impacts the actual timeframe to make ECE determinations. Therefore, to ensure transparency and understanding of our process, we are also taking this opportunity to clarify that we will strive to provide our response to an ECE request within 90 days of receipt.

(1) Proposal To Extend the ECE Request Deadline

In the past, we have allowed facilities to submit an ECE request form within 30 calendar days following the occurrence of an extraordinary circumstance that causes hardship and prevents them from providing data. In certain circumstances, however, it may be difficult for facilities to timely evaluate the impact of a certain extraordinary circumstance within 30 calendar days.

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299 ECEs were originally referred to as “waivers.” This term was changed to “exceptions” in the FY 2015 IPPS/LTCH PPS final rule (79 FR 50286).
We believe that extending the deadline to 90 calendar days would allow PCHs more time to determine whether it is necessary and appropriate to submit an ECE request and to provide a more comprehensive account of the extraordinary circumstance in their ECE request form to CMS. For example, if a PCH has suffered damage due to a hurricane on June 1, it would have until August 30 to submit an ECE form via the QualityNet Secure Portal, mail, email, or secure fax as instructed on the ECE form. We are inviting public comments on this proposal.

(2) Proposal To Grant Exceptions or Extensions Due to CMS Data System Issues

Although we do not anticipate this situation will happen often, there may be times where CMS experiences issues with its data systems that directly affects facilities’ abilities to submit data. In these circumstances, we are proposing to grant exceptions or extensions to one or more data reporting requirements. If we make the determination to grant exceptions or extensions to PCHs on this basis, we are proposing to communicate this decision through routine communication channels.

We are inviting public comment on this proposal.

G. Long-Term Care Hospital Quality Reporting Program (LTCH QRP)

1. Background and Statutory Authority

Section 3004(a) of the Affordable Care Act amended section 1886(m) of the Act by adding paragraph (5), requiring the Secretary to establish the Long-Term Care Hospital Quality Reporting Program (LTCH QRP). This program applies to all hospitals certified by Medicare as LTCHs. Beginning with the FY 2014 LTCH QRP, the Secretary is required to reduce any annual update to the LTCH PPS standard Federal rate for discharges occurring during such fiscal year by 2 percentage points for any LTCH that does not comply with the requirements established by the Secretary. Specifically, section 1886(m)(5) of the Act requires LTCHs to submit data on quality measures specified by the Secretary in a form and manner, and at a time, specified by the Secretary. For more information on the statutory history of the LTCH QRP, we refer readers to the FY 2015 IPPS/LTCH PPS final rule (79 FR 50286).

Please note that term “FY [year] LTCH QRP” refers to the fiscal year for which the LTCH QRP requirements applicable to that fiscal year must be met for an LTCH to receive the full annual update when calculating the payment rates applicable to it for that fiscal year.

The Improving Medicare Post-Acute Care Transformation Act of 2014 (IMPACT Act) (Pub. L. 113–185) amended Title XVIII of the Act, in part, by adding a new section 1899B of the Act, entitled “Standardized Post-Acute Care (PAC) Assessment Data for Quality, Payment, and Discharge Planning,” that enacts new data reporting requirements for certain post-acute care (PAC) providers, including LTCHs.

Specifically, new sections 1899B(a)(1)(A)(ii) and (iii) of the Act require LTCHs to report quality measures for inpatient rehabilitation facilities (IRFs), skilled nursing facilities (SNFs) and home health agencies (HHAs), under each of their respective quality reporting program (which, for LTCHs, is found at section 1886(m)(5) of the Act), to report data quality measures specified under section 1899B(c)(1), with respect to at least five domains, and data on resource use and other measures specified under section 1899B(d)(1) of the Act with respect to at least three domains. Section 1899B(a)(1)(A) of the Act further requires each of these PAC providers to report data on quality measures for use in the LTCH setting. We have previously adopted measures for the nursing home setting but not for the LTCH setting. For such measures, we intend to seek endorsement for the LTCH setting, and if the NQF endorses one or more of them, we will update the title of the measure to remove the reference to “application.”

b. Accounting for Social Risk Factors in the LTCH QRP

We consider related factors that may affect measures in the LTCH QRP. We understand that social risk factors such as income, education, race and ethnicity, employment, disability, community resources, and social support (certain factors of which are also sometimes referred to as socioeconomic status (SES) factors or socioeconomic status (SOS) factors) play a major role in health. One of our core objectives is to improve beneficiary outcomes including reducing health disparities, and we want to ensure that all beneficiaries, including those with social risk factors, receive high quality care. In addition, we seek to ensure that the quality of care furnished by providers and suppliers is assessed as fairly as possible under our programs while ensuring that beneficiaries have adequate access to excellent care.

We have been reviewing reports prepared by the Office of the Assistant Secretary for Planning and Evaluation
potential disparities or minimize incentives to improve the outcomes for disadvantaged populations. Keeping this concern in mind, while we sought input on this topic previously, we continue to seek public comment on whether we should account for social risk factors in measures in the LTCH QRP, and if so, what method or combination of methods would be most appropriate for accounting for social risk factors. Examples of methods include: confidential reporting to providers of measure rates stratified by social risk factors; public reporting of stratified measure rates; and potential risk adjustment of a particular measure as appropriate based on data and evidence.

In addition, we are also seeking public comment on which social risk factors might be most appropriate for reporting stratified measure scores and/or potential risk adjustment of a particular measure. Examples of social risk factors include, but are not limited to, dual eligibility/low-income subsidy, race and ethnicity, and geographic area of residence. We are seeking comments on which of these factors, including current data sources where this information would be available, could be used alone or in combination, and whether other data should be collected to better capture the effects of social risk. We will take commenters’ input into consideration as we continue to assess the appropriateness and feasibility of accounting for social risk factors in the LTCH QRP. We note that any such changes would be proposed through future notice and comment rulemaking.

We look forward to working with stakeholders as we consider the issue of accounting for social risk factors and reducing health disparities in CMS programs. Of note, implementing any of the above methods would be taken into consideration in the context of how this and other CMS programs operate (for example, data submission methods, availability of data, statistical considerations relating to reliability of data calculations, among others), so we also welcome comment on operational considerations. CMS is committed to ensuring that its beneficiaries have access to and receive excellent care, and that the quality of care furnished by providers and suppliers is assessed fairly in CMS programs.

3. Proposed Collection of Standardized Patient Assessment Data Under the LTCH QRP

a. Proposed Definition of Standardized Patient Assessment Data

Section 1886(m)(5)(F)(ii) of the Act requires that, for fiscal year 2019 and each subsequent year, LTCHs report standardized patient assessment data required under section 1899B(b)(1) of the Act. For purposes of meeting this requirement, section 1886(m)(5)(F)(iii) of the Act requires an LTCH to submit the standardized patient assessment data required under section 1899B(b)(1) of the Act using the standard instrument in a time, form, and manner specified by the Secretary.

Section 1899B(b)(1)(B) of the Act describes standardized patient assessment data as data required for at least the quality measures described in section 1899B(c)(1) of the Act and that is with respect to the following categories:

• Functional status, such as mobility and self-care at admission to a PAC provider and before discharge from a PAC provider;
• Cognitive function, such as ability to express ideas and to understand and mental status, such as depression and dementia;
• Special services, treatments and interventions such as the need for ventilator use, dialysis, chemotherapy, central line placement and total parenteral nutrition;
• Medical conditions and comorbidities such as diabetes, congestive heart failure and pressure ulcers;
• Impairments, such as incontinence and an impaired ability to hear, see or swallow; and
• Other categories deemed necessary and appropriate.

As required under section 1899B(b)(1)(A) of the Act, the standardized patient assessment data must be reported at least with respect to LTCH admissions and discharges, but the Secretary may require the data to be reported more frequently.

In this proposed rule, we are proposing to define the standardized patient assessment data that LTCHs must report to comply with section 1886(m)(5)(F)(ii) of the Act, as well as the requirements for the reporting of these data. The collection of standardized patient assessment data is critical to our efforts to drive improvement in health care quality across the four PAC settings to which the IMPACT Act applies. We intend to use these data for a number of purposes, including facilitating their exchange and

longitudinal use among health care providers to enable high quality care and outcomes through care coordination, as well as for quality measure calculation and identifying comorbidities that might increase the medical complexity of a particular admission.

LTCHs are currently required to report patient assessment data through the Long-Term Care Hospital Continuity Assessment Record and Evaluation Data Set (LTCH CARE Data Set or LCDS) by responding to an identical set of assessment questions using an identical set of response options (we refer to each solitary question/response option as a data element and we refer to a group of questions/responses as data elements), both of which incorporate an identical set of definitions and standards. The primary purpose of the identical questions and response options is to ensure that we collect a set of standardized data elements across LTCHs which can then be used for a number of purposes, including LTCH payment and measure calculation for the LTCH QRP.

SNFs, IRFs, and HHA s are also required to report patient assessment data through their applicable PAC assessment instruments, and they do so by responding to identical assessment questions developed for their respective settings using an identical set of response options (which incorporate an identical set of definitions and standards). Like the LCDS, the questions and response options for each of these other PAC assessment instruments are standardized across the PAC provider type to which the PAC assessment instrument applies. However, the assessment questions and response options in the four PAC assessment instruments are not currently standardized with each other. As a result, questions and response options that appear on the LCDS cannot be readily compared with questions and response options that appear, for example, on the Inpatient Rehabilitation Facility-Patient Assessment Instrument (IRF–PAI), the PAC assessment instrument used by IRFs. This is true even when the questions and response options are similar. This lack of standardization across the four PAC providers has limited our ability to compare one PAC provider type with another for purposes such as care coordination and quality improvement.

To achieve a level of standardization across SNFs, LTCHs, IRFs, and HHA s that enables us to make comparisons between SNFs, LTCHs, IRFs, and HHA s that enables us to make comparisons between them, we are proposing to define “standardized patient assessment data” as patient assessment questions and response options that are identical in all four PAC assessment instruments, and to which identical standards and definitions apply. Standardizing the questions and response options across the four PAC assessment instruments will also enable the data to be interoperable, allowing it to be shared electronically, or otherwise, between PAC provider types. It will enable the data to be comparable for various purposes, including the development of cross-setting quality measures and to inform payment models that take into account patient characteristics rather than setting, as described in the IMPACT Act.

We are inviting public comment on this proposed definition.

b. General Considerations Used for the Selection of Proposed Standardized Patient Assessment Data

As part of our effort to identify appropriate standardized patient assessment data for purposes of collecting under the LTCH QRP, we sought input from the general public, stakeholder community, and subject matter experts on items that would enable person-centered, high quality health care, as well as access to longitudinal information to facilitate coordinated care and improved beneficiary outcomes.

To identify optimal data elements for standardization, our data element contractor organized teams of researchers for each category, and each team worked with a group of advisors made up of clinicians and academic researchers with expertise in PAC. Information-gathering activities were used to identify data elements, as well as key themes related to the categories described in section 1899B(b)(1)(B) of the Act. In January and February 2016, our data element contractor also conducted provider focus groups for each of the four PAC provider types, and a focus group for consumers that included current or former PAC patients and residents, caregivers, ombudsmen, and patient advocacy group representatives. The Development and Maintenance of Post-Acute Care Cross-Setting Standardized Patient Assessment Data Focus Group Summary Report is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html. Our data element contractor also assembled a 16-member TEP that met on April 7 and 8, 2016, and January 5 and 6, 2017, in Baltimore, Maryland, to provide expert input on data elements that are currently in each PAC assessment instrument, as well as data elements that could be standardized. The Development and Maintenance of Post-Acute Care Cross-Setting Standardized Patient Assessment Data TEP Summary Reports are available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

As part of the environmental scan, data elements currently in the four existing PAC assessment instruments were examined to see if any could be considered for proposal as standardized patient assessment data. Specifically, this evaluation included consideration of data elements in OASIS–C2 (effective January 2017); IRF–PAI, v1.4 (effective October 2016); LCDS, v3.00 (effective April 2016); and MDS 3.0, v1.14 (effective October 2016). Data elements in the standardized assessment instrument that we tested in the Post-Acute Care Payment Reform Demonstration (PAC PRD)—the Continuity Assessment Record and Evaluation (CARE)—were also considered. A literature search was also conducted to determine whether additional data elements to propose as standardized patient assessment data could be identified.

We also held four Special Open Door Forums (SODFs) on October 27, 2015; May 12, 2016; September 15, 2016; and December 8, 2016, to present data elements we were considering and solicit input. At each SODF, some stakeholders provided immediate input, and all were invited to submit additional comments via the CMS IMPACT Mailbox at: PACQualityInitiative@cms.hhs.gov.

We also convened a meeting with federal agency subject matter experts (SMEs) on May 13, 2016. In addition, a public comment period was open from August 12 to September 12, 2016, to solicit comments on detailed candidate data element descriptions, data collection methods, and coding methods. The IMPACT Act Public Comment Summary Report containing the public comments (summarized and verbatim) and our responses, is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

We specifically sought to identify standardized patient assessment data that we could feasibly incorporate into the LTCH, IRF, SNF, and HHA.
assessments and that have the following attributes: (1) Being supported by current science; (2) testing well in terms of their reliability and validity, consistent with findings from the Post-Acute Care-Payment Reform Demonstration (PAC PRD); (3) the potential to be shared (for example, through interoperable means) among PAC and other provider types to facilitate efficient care coordination and improved beneficiary outcomes; (4) the potential to inform the development of quality, resource use and other measures, as well as future payment methodologies that could more directly take into account individual beneficiary health characteristics; and (5) the ability to be used by practitioners to inform their clinical decision and care planning activities. We also applied the same considerations that we apply with quality measures, including the CMS Quality Strategy which is framed using the three broad aims of the National Quality Strategy.

4. Policy for Retaining LTCH QRP Measures and Proposal To Apply That Policy to Standardized Patient Assessment Data

In the FY 2013 IPPS/LTCH PPS final rule (77 FR 53614 through 53615), we adopted a policy that would allow any quality measure adopted for use in the LTCH QRP to remain in effect until the measure is removed, suspended, or replaced. For further information on how measures are considered for removal, suspension, or replacement, we refer readers to the FY 2013 IPPS/LTCH PPS final rule (77 FR 53614 through 53615). We are proposing to apply this policy to the standardized patient assessment data that we adopt for the LTCH QRP.

We are inviting public comment on our proposal.

5. Policy for Adopting Changes to LTCH QRP Measures and Proposal To Apply That Policy to Standardized Patient Assessment Data

In the FY 2013 IPPS/LTCH PPS final rule (77 FR 53615 through 53616), we adopted a subregulatory process to incorporate updates to LTCH quality measure specifications that do not substantively change the nature of the measure. Substantive changes will be proposed and finalized through rulemaking. For further information on what constitutes a substantive versus a nonsubstantive change and the subregulatory process for nonsubstantive changes, we refer readers to the FY 2013 IPPS/LTCH PPS final rule (77 FR 53615 through 53616). We are proposing to apply this policy to the standardized patient assessment data that we adopt for the LTCH QRP.

We are inviting public comment on our proposal.

6. Quality Measures Previously Finalized for the LTCH QRP

The LTCH QRP currently has 17 finalized measures as outlined in the table below:

### QUALITY MEASURES CURRENTLY ADOPTED FOR THE LTCH QRP

<table>
<thead>
<tr>
<th>Short name</th>
<th>Measure name and data source</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>LTCH CARE Data Set</strong></td>
<td></td>
</tr>
<tr>
<td>Pressure Ulcers</td>
<td>Percent of Residents or Patients with Pressure Ulcers That Are New or Worsened (Short Stay) (NQF #0678).</td>
</tr>
<tr>
<td>Patient Influenza Vaccine</td>
<td>Percent of Residents or Patients Who Were Assessed and Appropriately Given the Seasonal Influenza Vaccine (Short Stay) (NQF #0680).</td>
</tr>
<tr>
<td>Application of Falls</td>
<td>Application of Percent of Residents Experiencing One or More Falls with Major Injury (Long Stay) (NQF #0674).*</td>
</tr>
<tr>
<td>Functional Assessment</td>
<td>Percent of Long-Term Care Hospital (LTCH) Patients with an Admission and Discharge Functional Assessment and a Care Plan That Addresses Function (NQF #2631).</td>
</tr>
<tr>
<td>Application of Functional Assessment</td>
<td>Application of Percent of Long-Term Care Hospital (LTCH) Patients with an Admission and Discharge Functional Assessment and a Care Plan That Addresses Function (NQF #2631).</td>
</tr>
<tr>
<td>Change in Mobility</td>
<td>Functional Outcome Measure: Change in Mobility Among Long-Term Care Hospital (LTCH) Patients Requiring Ventilator Support (NQF #2632).</td>
</tr>
<tr>
<td>DRR</td>
<td>Drug Regimen Review Conducted With Follow-Up for Identified Issues- Post Acute Care (PAC) Long-Term Care Hospital (LTCH) Quality Reporting Program (QRP).*</td>
</tr>
<tr>
<td><strong>NHSN</strong></td>
<td></td>
</tr>
<tr>
<td>MRSA</td>
<td>National Healthcare Safety Network (NHSN) Facility-wide Inpatient Hospital-onset Methicillin-resistant Staphylococcus aureus (MRSA) Bacteremia Outcome Measure (NQF #1716).</td>
</tr>
<tr>
<td>CDI</td>
<td>National Healthcare Safety Network (NHSN) Facility-wide Inpatient Hospital-onset Clostridium difficile Infection (CDI) Outcome Measure (NQF #1717).</td>
</tr>
<tr>
<td>HCP Influenza Vaccine</td>
<td>Influenza Vaccination Coverage among Healthcare Personnel (NQF #0431).</td>
</tr>
<tr>
<td>VAE</td>
<td>National Healthcare Safety Network (NHSN) Ventilator-Associated Event (VAE) Outcome Measure.*</td>
</tr>
<tr>
<td><strong>Claims-Based</strong></td>
<td></td>
</tr>
<tr>
<td>All-Cause Readmissions</td>
<td>All-Cause Unplanned Readmission Measure for 30-Days Post-Discharge from Long-Term Care Hospitals (LTCHs) (NQF #2512).</td>
</tr>
<tr>
<td>MSPB</td>
<td>Medicare Spending Per Beneficiary (MSPB)-Post Acute Care (PAC) Long-Term Care Hospital (LTCH) Quality Reporting Program (QRP).*</td>
</tr>
<tr>
<td>DTC</td>
<td>Discharge to Community-Post Acute Care (PAC) Long-Term Care Hospital (LTCH) Quality Reporting Program (QRP).*</td>
</tr>
</tbody>
</table>
7. LTCH QRP Quality Measures Proposed Beginning With the FY 2020 LTCH QRP

Beginning with the FY 2020 LTCH QRP, in addition to the quality measures we are retaining under our policy described in section IX.C.4. of the preamble of this proposed rule, we are proposing to remove the current pressure ulcer measure entitled Percent of Residents or Patients with Pressure Ulcers That Are New or Worsened (Short Stay) (NQF #0678) and replace it with a modified version of the measure entitled Changes in Skin Integrity Post-Acute Care: Pressure Ulcer/Injury and adopt two new measures (one process and one outcome) related to ventilator weaning. We are also proposing to characterize the data elements described below as standardized patient assessment data under section 1899(b)(1)(B) of the Act that must be reported by LTCHs under the LTCH QRP through the LTCH CARE Data Set.

The proposed measures are as follows:

- Changes in Skin Integrity Post-Acute Care: Pressure Ulcer/Injury
- Compliance with Spontaneous Breathing Trial (SBT) by Day 2 of the LTCH Stay
- Ventilator Liberation Rate

The measures are described in more detail below.

a. Proposal To Replace the Current Pressure Ulcer Quality Measure, Percent of Residents or Patients With Pressure Ulcers That Are New or Worsened (Short Stay) (NQF #0678), With a Modified Pressure Ulcer Measure, Changes in Skin Integrity Post-Acute Care: Pressure Ulcer/Injury

(1) Measure Background

In this proposed rule, we are proposing to remove the current pressure ulcer measure, Percent of Residents or Patients with Pressure Ulcers That Are New or Worsened (Short Stay) (NQF #0678), from the LTCH QRP measure set and to replace it with a modified version of that measure, Changes in Skin Integrity Post-Acute Care: Pressure Ulcer/Injury, beginning with the FY 2020 LTCH QRP.

The change in the measure name is to reduce confusion about the new modified measure. The modified version differs from the current version of the measure because it includes new or worsened unstageable pressure ulcers, including deep tissue injuries (DTIs), in the measure numerator. The proposed modified version of the measure also contains updated specifications intended to eliminate redundancies in the assessment items needed for its calculation and to reduce the potential for underestimating the frequency of pressure ulcers. The modified version of the measure would satisfy the IMPACT Act domain of skin integrity and changes in skin integrity.

(2) Measure Importance

As described in the FY 2012 IPPS/LTCNPS final rule (76 FR 51754 through 51756), pressure ulcers are high-cost adverse events and are an important measure of quality. For information on the history and rationale for the relevance, importance, and applicability of having a pressure ulcer measure in the LTCH QRP, we refer readers to the FY 2012 IPPS/LTCNPS final rule (76 FR 51748 through 51750) and the FY 2014 IPPS/LTCNPS final rule (78 FR 50861 through 50863).

We are proposing to adopt a modified version of the current pressure ulcer measure because unstable pressure ulcers, including DTIs, are similar to Stage 2, Stage 3, and Stage 4 pressure ulcers in that they represent poor outcomes, are a serious medical condition that can result in death and disability, are debilitating and painful, and are often an avoidable outcome of medical care. Studies show that most pressure ulcers can be avoided and can also be healed in acute, post-acute, and long-term care settings with appropriate medical care.

Furthermore, some studies indicate that DTIs, if managed using appropriate care, can be resolved without deteriorating into a worsened pressure ulcer. While there are few studies that provide information regarding the incidence of unstable pressure ulcers in PAC settings, an analysis conducted by a contractor suggests the incidence of unstable pressure ulcers varies according to the type of unstable pressure ulcer and setting. This analysis examined the national incidence of new unstable pressure ulcers in LTCHs at discharge compared with admission using LTCH discharges from January through December 2015. The contractor found a national incidence of 1.15 percent of new unstable pressure ulcers due to slough and/or eschar, 0.05 percent of new unstable pressure ulcers due to non-removable dressing/device, and 1.01 percent of new DTIs. In addition, an international study spanning the time period 2006 to 2017 provides some evidence to suggest that the proportion of pressure ulcers identified as DTI has increased over time. The study found DTIs increased by three fold, to nine percent of all observed ulcers in 2009, and that DTIs were more prevalent than either Stage 3 or 4 ulcers. During the same time period, the proportion of Stage 1 and 2 ulcers decreased, and the proportion of Stage 3 and 4 ulcers remained constant.


The inclusion of unstageable pressure ulcers, including DTIs, in the numerator of this measure is expected to increase measure scores and variability in measure scores, thereby improving the ability to discriminate among poor- and high-performing LTCHs. In the currently implemented pressure ulcer measure, Percent of Residents or Patients with Pressure Ulcers That Are New or Worse (Short Stay) (NQF #0678), analysis using data from Quarter 1 through Quarter 4 2015 data reveals that the LTCH mean score is 1.95 percent; the 25th and 75th percentiles are 0.53 percent and 2.49 percent, respectively; and 12.11 percent of facilities have perfect scores. In the proposed measure, Changes in Skin Integrity Post-Acute Care: Pressure Ulcer/Injury, during the same timeframe, the LTCH mean score is 3.73 percent; the 25th and 75th percentiles are 1.53 percent and 4.89 percent, respectively; and 5.46 percent of facilities have perfect scores.

(3) Stakeholder Feedback
Our measure development contractor sought input from subject matter experts, including Technical Expert Panels (TEPs), over the course of several years on various skin integrity topics and specifically those associated with the inclusion of unstageable pressure ulcers, including DTIs. Most recently, on July 18, 2016, a TEP convened by our measure development contractor provided input on the technical specifications of this proposed quality measure, including the feasibility of implementing the proposed measure’s updates across PAC settings. The TEP supported the updates to the measure across PAC settings, including the inclusion in the numerator of unstageable pressure ulcers due to slough and/or eschar that are new or worsened, new unstageable pressure ulcers due to a non-removable dressing or device, and new DTIs. The TEP also supported the use of different data elements for measure calculation. The TEP recommended supplying additional guidance to providers regarding each type of unstageable pressure ulcer. This support was in agreement with earlier TEP meetings, held on June 13 and November 15, 2013, which had recommended that CMS update the specifications for the pressure ulcer measure to include unstageable pressure ulcers in the numerator.315 316

Exploratory data analysis conducted by our measure development contractor suggests that the addition of unstageable pressure ulcers, including DTIs, will increase the observed incidence and variation in the rate of new or worsened pressure ulcers at the facility level, which may improve the ability of the proposed quality measure to discriminate between poor- and high-performing facilities.

We solicited stakeholder feedback on this proposed measure by means of a public comment period held from October 17 through November 17, 2016. In general, we received considerable support for the proposed measure. A few commenters supported all of the changes to the current pressure ulcer measure that resulted in the proposed measure, with one commenter noting the significance of the work to align the pressure ulcer quality measure specifications across the PAC settings.

Many commenters supported the inclusion of unstageable pressure ulcers due to slough and/or eschar that are new or worsened pressure ulcers. There are related pressure ulcer quality measures for PAC settings, but there is no universally accepted definition for this type of skin injury. Some commenters provided feedback on the data elements used to calculate the proposed quality measure. We believe that these data elements will promote facilitation of cross-setting quality comparison as mandated by the IMPACT Act, alignment between quality measures and payment, reduction in redundancies in assessment items, and prevention of inappropriate underestimation of pressure ulcers. The currently implemented pressure ulcer measure is calculated using retrospective data elements that assess the number of new or worsened pressure ulcers at each stage, while the proposed measure is calculated using the number of unhealed pressure ulcers at each stage after subtracting the number that were present upon admission. Some commenters did not support the data elements that would be used to calculate the proposed measure, and requested further testing of these data elements. Other commenters supported the use of these data elements stating that these data elements simplified the measure calculation process.


The NQF-convened Measures Application Partnership (MAP) Post-Acute Care/Long-Term Care (PAC/LTC) Workgroup met on December 14 and 15, 2016, and provided input about this measure. The Workgroup provided a recommendation of “support for rulemaking” for use of the proposed measure in the LTCH QRP. The MAP Coordinating Committee met on January 24 and 25, 2017, and provided a recommendation of “conditional support for rulemaking” for use of the proposed measure in the LTCH QRP. The MAP’s conditions of support include that, as a part of measure implementation, CMS provide guidance on the correct collection and calculation of the measure result, as well as guidance on public reporting Web sites explaining the impact of the specification changes on the measure result. The MAP’s conditions also specify that CMS continue analyzing the proposed measure in order to investigate unexpected results reported in public comment. We intend to fulfill these conditions by offering additional training opportunities and educational materials in advance of public reporting, and by continuing to monitor and analyze the proposed measure. More information about the MAP’s recommendations for this measure is available at: http://www.qualityforum.org/WorkArea/linkit.aspx?LinkIdentifier=id&ItemID=84452.

We reviewed the NQF’s consensus endorsed measures and were unable to identify any NQF-endorsed pressure ulcer quality measures for PAC settings that are inclusive of unstageable pressure ulcers. There are related measures, but after careful review, we determined these measures were not applicable for use in LTCHs based on the populations addressed or other...
aspects of the specifications. We are unaware of any other such quality measures that have been endorsed or adopted by another consensus organization for the LTCH setting. Therefore, based on the evidence discussed above, we are proposing to adopt the quality measure entitled, Changes in Skin Integrity Post-Acute Care: Pressure Ulcer/Injury, for the LTCH QRP beginning with the FY 2020 LTCH QRP. We plan to submit the proposed measure to the NQF for endorsement consideration as soon as feasible.

(4) Data Collection

The data for this quality measure would be collected using the LTCH CARE Data Set, which is currently submitted by LTCHs through the QIES ASAP System. The proposed standardized patient assessment data applicable to this measure that must be reported by LTCHs for admissions as well as discharges occurring on or after April 1, 2018 is described in section IX.C.11. of the preamble of this proposed rule. While the inclusion of unstageable wounds in the proposed measure results in a measure calculation methodology that is different from the methodology used to calculate the current pressure ulcer measure, the data elements needed to calculate the proposed measure are already included on the LTCH CARE Data Set. In addition, our proposal to eliminate duplicative data elements that were used in calculation of the current pressure ulcer measure will result in an overall reduced reporting burden for LTCHs with respect to the proposed measure. For more information on LTCH CARE Data Set submission using the QIES ASAP System, we refer readers to: http://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/ LTCHTechnicalInformation.html.

For technical information about this proposed measure, including information about the measure calculation and the standardized patient assessment data elements used to calculate this measure, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/ LTCH-Quality-Reporting-Measures-Information.html.

We are proposing that LTCHs would begin reporting the proposed pressure ulcer measure, Changes in Skin Integrity Post-Acute Care: Pressure Ulcer/Injury, which will replace the current pressure ulcer measure, with data collection beginning April 1, 2018.

We are inviting public comment on our proposal to replace the current pressure ulcer measure, Percent of Residents or Patients with Pressure Ulcers That Are New or Worsened (Short Stay) (NQF #0678), with a modified version of that measure, entitled Changes in Skin Integrity Post-Acute Care: Pressure Ulcer/Injury, for the LTCH QRP beginning with the FY 2020 LTCH QRP.

b. Proposed Mechanical Ventilation Process Quality Measure: Compliance With Spontaneous Breathing Trial (SBT) by Day 2 of the LTCH Stay

Invasive mechanical ventilation care was identified through technical expert panels convened by our measure development contractor and public comment periods as a gap in the LTCH QRP measure set and aligns with the National Quality Strategy priority and the CMS Quality Strategy goal of “promoting the most effective prevention and treatment practices” by reducing the risk of complications from unnecessarily prolonged mechanical ventilation. We are proposing to adopt the quality measure, Compliance with Spontaneous Breathing Trial (SBT) by Day 2 of the LTCH Stay, beginning with the FY 2020 LTCH QRP. The data applicable to this measure that must be reported by LTCHs for admissions as well as discharges occurring on or after April 1, 2018 is described in section IX.C.11. of the preamble of this proposed rule.

The Compliance with SBT by Day 2 of the LTCH Stay measure is a process quality measure. For patients on invasive mechanical ventilation support upon admission to the LTCH, except those who meet measure exclusion criteria, this measure assesses facility-level compliance with SBT, including TCT or CPAP breathing trial, by Day 2 of the LTCH stay, where Day 1 is the day of admission to the LTCH and Day 2 is the subsequent calendar day. This measure is calculated and reported for the following two components: (1) The percentage of patients admitted on invasive mechanical ventilation support upon admission to the LTCH; and (2) the percentage of patients deemed medically ready for SBT who received SBT by Day 2 of the LTCH stay. Higher percentages indicate better compliance. Patients are included in this quality measure if they are on invasive mechanical ventilation support upon admission to the LTCH, unless they meet measure exclusion criteria.

Patients on invasive mechanical ventilation support present a critical focus for assessment of high quality care because they comprise a substantial proportion of LTCH patient admissions. Mechanically ventilated patients are increasingly common in both acute care hospital intensive care units (ICUs), where up to 40 percent of patients require some duration of mechanical ventilation,\(^317\) and LTCHs, where patients are frequently transferred for weaning following treatment in ICUs.\(^318\)\(^319\)\(^320\) Patients who require invasive mechanical ventilation of longer than 14 or 21 days are undergoing a long-term mechanical ventilation (PMV). In 2012, about 22,000 or 15.8 percent of all LTCH discharges received PMV services during the LTCH stay.\(^321\)

This ventilator weaning-related process quality measure is important for encouraging implementation of evidence-based weaning guidelines as early during the LTCH patient stay as is beneficial to the patient. Although often necessary for life support, invasive mechanical ventilation is not without risk of harm to patients, and these risks increase as duration of ventilation continues.\(^322\)\(^323\)\(^324\)\(^325\) In both ICUs and LTCHs, unsuccessful weaning and delayed weaning increase patient exposure to a number of ventilator-associated negative health outcomes, including ventilator-associated pneumonia,\(^325\)\(^326\)\(^327\)\(^328\) ventilator-


associated lung injury,329 330 331 ventilator induced diaphragm dysfunction,332 psychological distress333 334 335 and post-traumatic stress disorder,336 disability337 and decreased functional status,338 339 and chronic critical illness syndrome.340 Furthermore, these ventilator-associated negative health outcomes particularly affect the LTCH population since a significant number of its patients are on PMV. The majority of mechanically ventilated patients who are transferred to an LTCH have received mechanical ventilation for at least 21 days.341 PMV increases the risk of patient morbidity and short-term and long-term mortality. According to a recent systematic review, the pooled mortality of patients with PMV (defined here as invasive mechanical ventilation for ≥14 days) undergoing weaning attempts in LTCHs was 31 percent (18 studies); however, the pooled mortality at one year significantly increased to 73 percent (8 studies).342 In addition to increased morbidity and mortality, mechanical ventilation is also associated with higher costs. While the literature on costs of mechanical ventilation is limited for the LTCH setting, studies in the acute care hospital ICU setting indicate that patients who require mechanical ventilation can have up to 50 percent higher costs than patients who do not receive mechanical ventilation.343 ICU patients who develop VAP incur at least $40,000 more in hospital costs than ventilated patients without VAP, and costs increase with increasing duration of mechanical ventilation.344 345 346 Although there is evidence regarding the benefit of daily assessments of patient readiness for weaning from invasive mechanical ventilation,347 as well as for the importance of adherence to weaning protocols,348 we are not aware of any studies in LTCHs that evaluate timing of assessment for readiness to wean with respect to the admission date. However, an international task force, convened in 2005, developed guideline recommendations to address the entire weaning process. Despite the limited evidence, this task force recommended that weaning be considered as soon as possible,349 because failure to assess the patient for readiness to wean may lead to undue prolonged mechanical ventilation,350 thus exposing patients unnecessarily to adverse ventilator-associated morbidity and mortality.351 Based on studies and observations of implementation of regular assessment for SBTs and weaning protocols in ICUs, adherence to the recommended weaning processes, including prompt assessment of weaning readiness and initiation of SBTs, appears quite variable, likely due to differences in clinicians’ intuitive thresholds for determination of patients’ readiness to wean.352 353 Clinician delays in recognizing that weaning may be possible and beginning assessment of weaning readiness are two common causes of weaning delays.354 In one study, 50 percent of the patients considered to be incapable of sustaining spontaneous ventilation by clinicians later were able to tolerate a weaning trial. The authors concluded that tests used to validate clinician intuition on a patient’s readiness for weaning are often inaccurate and that clinicians should follow explicit protocols to consistently test patients on their readiness to wean.355 Because prompt identification of patients’ readiness for SBTs has been shown to reduce weaning duration without harm to patients,356 such delays indicate less than optimal

mechanical ventilation. In addition, facilities can use results of this measure to improve timely compliance with evidence-based weaning guidelines and develop ventilator weaning quality improvement programs.

A TEP assembled by our measure development contractor convened nine meetings (two in-person meetings and seven webinars) between April 2014 and August 2016 in order to refine the quality measure’s technical specifications, including the measure target population, inclusion and exclusion criteria, and key definitions (for example, “non-weaning”). The TEP also offered feedback on the individual LTCH CARE Data Set ventilator weaning items and supported the feasibility of implementing this measure in the LTCH setting. The measure developer recruited two former patients successfully weaned from mechanical ventilation as well as the primary caregiver of one of the patients to solicit their views on the measures. The 2014–2016 Development of Long-Term Care Hospital (LTCH) Ventilator Weaning Quality Measures Technical Expert Panel Summary Report is available on the CMS Web site at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

We also solicited stakeholder feedback on the development of this measure through a public comment period held from May 19, 2016, through June 9, 2016. Several stakeholders and organizations supported this measure for implementation, including hospitals and professional organizations. The public comment summary report for the proposed measure is available on the CMS Web site at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

Our measure development contractor conducted a pilot test on the data elements used to calculate this quality measure. The pilot test was conducted in 10 LTCHs among approximately 150 LTCH patients and used a mixed methods research design to collect data. Quantitative data on the ventilator weaning items was collected from May 27, 2016 through September 10, 2016, and qualitative data on these items was collected from June 6, 2016 through October 4, 2016. The LTCHs who participated in the pilot test were selected to represent variation across several key facility-level characteristics: geographic location, size, and profit status.

The qualitative data from the pilot test of the ventilator weaning process measure supported the importance of the measure. Results from qualitative and quantitative analysis further support the feasibility of data collection for this quality measure. Data collection for this quality measure was not seen as burdensome by pilot sites. The pilot test summary report for this measure is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

The NQF-convened MAP PAC/LTC Workgroup met on December 12, 2014 and again on December 14 and 15, 2015. During these meetings, the MAP encouraged continued development of this proposed measure, acknowledging that there is evidence for interventions that improve ventilator care, that variation in quality of care exists among LTCHs, and that ventilator care is an important safety priority for LTCHs.

Since the MAP’s review and recommendation of continued development in 2015, we have continued to refine this proposed measure in compliance with the MAP’s recommendations. Results of continued development activities, including stakeholder feedback from the 2016 public comment period and 2016 pilot test findings, were presented to the MAP during the MAP feedback loop meeting in October 2016. The proposed measure is consistent with the information submitted to the MAP, and the original MAP submission and our continued refinements support its scientific acceptability for use in quality reporting programs. As discussed with the MAP, we fully anticipate that additional analyses will continue once data collection for the measure begins. More information about the MAP’s recommendations for this proposed measure is available at: http://www.qualityforum.org/Publications/2016/02/MAP_2016_Considerations_

We reviewed the NQF’s consensus endorsed measures and were unable to identify any NQF-endorsed ventilator weaning quality measures focused on assessment of readiness to wean for patients admitted on invasive mechanical ventilation in the LTCH setting. We are unaware of any other quality measures for weaning from invasive mechanical ventilation that have been endorsed or adopted by another consensus organization for the LTCH setting. Therefore, based on the evidence discussed above, we are proposing to adopt the quality measure entitled, Compliance with SBT by Day 2 of the LTCH Stay, for the LTCH QRP beginning with the FY 2020 LTCH QRP.

We plan to submit the quality measure to the NQF for consideration for endorsement.

We are proposing that data for this ventilator weaning quality measure be collected through the LTCH CARE Data Set, with submission through the QIES ASAP System. For more information on LTCH QRP reporting using the QIES ASAP System, we refer readers to our Web site at: http://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment- Instruments/LTCH-Quality-Reporting/LTCHTechnicalInformation.html. We intend to revise the LTCH CARE Data Set to include new items that assess processes for weaning from invasive mechanical ventilation, should this proposed measure be adopted.

This measure is calculated and reported for two components. The proposed measure numerator for Component 1 is the number of patients admitted on invasive mechanical ventilation during the reporting period who were ready for SBT and who received an SBT (including TCT or CPAP breathing trial) by Day 2 of the LTCH stay.

For technical information about this proposed measure, including information about the measure calculation and proposed measure denominator exclusions, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

We are inviting public comments on our proposal to adopt the quality measure, Compliance with SBT by Day 2 of the LTCH Stay, beginning with the FY 2020 LTCH QRP.

c. Proposed Mechanical Ventilation Outcome Quality Measure: Ventilator Liberation Rate

Invasive mechanical ventilation care was identified as an important gap in the LTCH QRP measure set, and aligns with the National Quality Strategy priority and the CMS Quality Strategy goal of “promoting the most effective prevention and treatment practices” by reducing the risk of complications from unnecessarily prolonged mechanical ventilation. We are proposing to adopt the quality measure, Ventilator Liberation Rate, for the LTCH QRP beginning with the FY 2020 LTCH QRP. The data applicable to this measure that must be reported by LTCHs for admissions as well as discharges occurring on or after April 1, 2018 is described in section IX.C.11. of the preamble of this proposed rule.

The Ventilator Liberation Rate measure is an outcome quality measure. This quality measure is a facility-level measure that reports the percentage of LTCH patients admitted on invasive mechanical ventilation during the reporting period who were assessed for readiness for SBT (including TCT or CPAP breathing trial) by Day 2 of the LTCH stay.

The proposed measure denominator for Component 2, Percentage of Patients Ready for SBT Who Received SBT by Day 2 of LTCH Stay, is the number of patients admitted on invasive mechanical ventilation during the reporting period who were ready for SBT and who received an SBT (including TCT or CPAP breathing trial) by Day 2 of the LTCH stay. The proposed measure numerator for Component 2, Percentage of Patients Ready for SBT Who Received SBT by Day 2 of LTCH Stay, is the number of patients admitted on invasive mechanical ventilation during the reporting period who were ready for SBT and who received an SBT (including TCT or CPAP breathing trial) by Day 2 of the LTCH stay.

For technical information about this proposed measure, including information about the measure calculation and proposed measure denominator exclusions, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

We are inviting public comments on our proposal to adopt the quality measure, Compliance with SBT by Day 2 of the LTCH Stay, beginning with the FY 2020 LTCH QRP.

c. Proposed Mechanical Ventilation Outcome Quality Measure: Ventilator Liberation Rate

Invasive mechanical ventilation care was identified as an important gap in the LTCH QRP measure set, and aligns with the National Quality Strategy priority and the CMS Quality Strategy goal of “promoting the most effective prevention and treatment practices” by reducing the risk of complications from unnecessarily prolonged mechanical ventilation. We are proposing to adopt the quality measure, Ventilator Liberation Rate, for the LTCH QRP beginning with the FY 2020 LTCH QRP. The data applicable to this measure that must be reported by LTCHs for admissions as well as discharges occurring on or after April 1, 2018 is described in section IX.C.11. of the preamble of this proposed rule.

The Ventilator Liberation Rate measure is an outcome quality measure. This quality measure is a facility-level measure that reports the percentage of LTCH patients admitted on invasive mechanical ventilation during the reporting period who were assessed for readiness for SBT (including TCT or CPAP breathing trial) by Day 2 of the LTCH stay.

The proposed measure denominator for Component 1, Percentage of Patients Assessed for Readiness by Day 2 of LTCH Stay, is the total number of patients admitted during the reporting period who were on invasive mechanical ventilation upon admission to an LTCH and expected or anticipated by the provider to undergo weaning attempts at admission. The proposed measure numerator for Component 1 is the number of patients admitted on invasive mechanical ventilation during the reporting period who were assessed for readiness for SBT (including TCT or CPAP breathing trial) by Day 2 of the LTCH stay.

The proposed measure denominator for Component 2, Percentage of Patients Ready for SBT Who Received SBT by Day 2 of LTCH Stay, is the subset of patients in the denominator of the Component 1, who were assessed and deemed ready for SBT by Day 2 of the LTCH stay. The proposed measure numerator for Component 2, Percentage of Patients Ready for SBT Who Received SBT by Day 2 of LTCH Stay, is the number of patients admitted on invasive mechanical ventilation during the reporting period who were ready for SBT and who received an SBT (including TCT or CPAP breathing trial) by Day 2 of the LTCH stay.

For technical information about this proposed measure, including information about the measure calculation and proposed measure denominator exclusions, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

We are inviting public comments on our proposal to adopt the quality measure, Compliance with SBT by Day 2 of the LTCH Stay, beginning with the FY 2020 LTCH QRP.

Numerous studies from 1991 through 2015 have reported a range of ventilator liberation rates among LTCHs. A review of nine single-center studies conducted between 1991 and 2001 reported that, among more than 3,000 patients with
prolonged mechanical ventilation increases the risk of negative patient outcomes and can be an indicator of poor quality care or of persistent illness. Based on the evidence, improving weaning processes and increasing weaning rates are expected to mitigate the risk of harm associated with invasive mechanical ventilation, thus contributing to more favorable clinical outcomes for patients and decreased costs. This quality measure, Ventilator Liberation Rate, will assess the proportion of patients discharged alive from an LTCH who are fully weaned, thereby promoting weaning efforts and encouraging quality management of LTCH patients on invasive mechanical ventilation. Kahn et al. (2013) noted that inclusion of a liberation outcome measure is key to providing a truly patient-centered measure related to invasive mechanical ventilation weaning among LTCH patients.

A TEP assembled by our measure development contractor convened nine meetings (two in-person meetings and seven webinars) between April 2014 and August 2016. TEP members provided input to guide the development of the quality measures, including feedback on the individual LTCH CARE Data Set ventilator weaning items, the target population, inclusion and exclusion criteria, and patient demographic and clinical factors that could affect ventilator weaning outcomes (risk adjstors). The TEP also supported the feasibility of implementing this measure in the LTCH setting. The measure developer recruited two former patients successfully weaned from mechanical ventilation as well as the primary caregiver of one of the patients to solicit their views on the measures. The 2014–2016 Development of Long-Term Care Hospital (LTCH) Ventilator Weaning Quality Measures Technical Expert Panel Summary Report is available on the CMS Web site at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

We also solicited stakeholder feedback on the development of this measure through a public comment period held from May 19, 2016, through June 9, 2016. Several stakeholders and organizations supported this measure for implementation, including hospitals and professional organizations. The public comment summary report for the proposed measure is available on the CMS Web site at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

Our measure development contractor conducted a pilot test on the proposed data elements used to calculate this quality measure. The pilot test was conducted in ten LTCHs among approximately 150 LTCH patients and used a mixed methods research design to collect data. Quantitative data on the ventilator weaning items was collected from May 27, 2016 through September 10, 2016, and qualitative data on these items was collected from June 6, 2016 through October 4, 2016. The LTCHs who participated in the pilot test were selected to represent variation across several key facility-level characteristics: geographic location, size, and profit status.

The qualitative data from the pilot test of the ventilator liberation quality measure supported the importance of the measure; results from qualitative and quantitative analysis also supported the feasibility of data collection. Data collection for this quality measure was not seen as burdensome by pilot sites. The pilot test summary report for this measure is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

The NQF-convened MAP PAC/LTC Workgroup met on December 12, 2014, and on December 14 and 15, 2015. During these meetings, the MAP provided input on the importance and

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specifications of this measure. The MAP encouraged continued development of the measure, stating that this measure has high value potential for the LTCH QRP because successful weaning is important for improving quality of life and decreasing morbidity, mortality, and resource use among patients. Since the MAP’s review and recommendation of continued development in 2015, we have continued to refine this proposed measure in compliance with the MAP’s recommendations. Results of continued development activities, including stakeholder feedback from the 2016 public comment period and 2016 pilot test findings, were presented to the MAP during the MAP feedback loop meeting in October 2016. The proposed measure is consistent with the information submitted to the MAP, and the original MAP submission and our continued refinements support its scientific acceptability for use in quality reporting programs. As discussed with the MAP, we fully anticipate that additional analyses will continue once data collection for the measure begins. More information about the MAP’s recommendations for this measure is available at: http://www.qualityforum.org/Publications/2016/02/MAP_2016_Considerations_for_Implementing_Measures_in_Federal_Programs_-PAC-LTC.aspx.

We reviewed the NQF’s consensus endorsed measures and were unable to identify any NQF-endorsed ventilator weaning quality measures focused on the liberation status at discharge for patients admitted on invasive mechanical ventilation in the LTCH setting. We are unaware of any other quality measures for liberation from invasive mechanical ventilation that have been endorsed or adopted by another consensus organization for the LTCH setting. Therefore, based on the evidence discussed above, we are proposing to adopt the quality measure entitled, Ventilator Liberation Rate, for the LTCH QRP beginning with the FY 2020 LTCH QRP. We plan to submit the quality measure to the NQF for consideration for endorsement.

We are proposing that data for this quality measure be collected through the LTCH CARE Data Set, with the submission through the QIES ASAP System. For more information on LTCH QRP reporting using the QIES ASAP system, we refer readers to our Web site at: http://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCQuality-Reporting/LTCHTechnicalInformation.html. We intend to revise the LTCH CARE Data Set to include new items that assess invasive mechanical ventilation liberation at discharge, should this proposed measure be adopted.

This measure reports facility-level Ventilator Liberation Rate for patients admitted to an LTCH on invasive mechanical ventilation, and for whom weaning attempts were expected or anticipated as reported on the Admission Assessment. The Ventilator Liberation Rate is defined as the percentage of patients on invasive mechanical ventilation upon admission who are alive and fully liberated at discharge. The proposed measure denominator is the number of patients requiring invasive mechanical ventilation support upon admission to an LTCH, except those who meet exclusion criteria. The proposed measure numerator is the number of patients who are discharged alive and fully liberated. This measure is risk-adjusted for variables such as age, neurological injury or disease, dialysis, and other comorbidities and treatments. If a patient has more than one LTCH stay during the reporting period, then each LTCH stay will be included in the measure calculation and reporting. For technical information about this measure, including information about the measure calculation, risk adjustment, and proposed measure denominator exclusions, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCQuality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

We are inviting public comments on our proposal to adopt the quality measure, Ventilation Liberation Rate, beginning with the FY 2020 LTCH QRP.

8. Proposed Removal of the All-Cause Unplanned Readmission Measure for 30 Days Post-Discharge From LTCHs From the LTCH QRP.

We are proposing to remove the All-Cause Unplanned Readmission Measure for 30 Days Post-Discharge from LTCHs (NQF #2512) from the LTCH QRP. In the FY 2016 IPPS/LTCPPS final rule (80 FR 49730 through 49731), we adopted the All-Cause Unplanned Readmission Measure for 30 Days Post-Discharge from LTCHs (NQF #2512) for the LTCH QRP. This measure assesses all-cause unplanned hospital readmissions from LTCHs. In the FY 2017 IPPS/LTCPPS final rule (81 FR 57215 through 57219), we adopted the Potentially Preventable 30-Day Post-Discharge Readmission Measure for LTCH QRP to fulfill IMPACT Act requirements. In response to the FY 2017 IPPS/LTCPPS proposed rule, we received public comments expressing concern over the multiplicity of readmission measures and the overlap between the All-Cause Readmission and Potentially Preventable Readmission (PPR) 30-Day Post-Discharge measures (see 81 FR 57217 through 57218). Commenters also stated that more than one readmission measure would create confusion and require additional effort by providers to track and improve performance.

We retained the All-Cause Readmission measure because it would allow us to monitor trends in both all-cause and PPR rates. In particular, we could compare facility performance on the All-Cause Readmission and PPR 30-Day Post-Discharge measures. However, upon further consideration of the public comments, we believe that removing the All-Cause Readmission measure and retaining the PPR 30-Day Post-Discharge measure in the LTCH QRP would prevent duplication, because potentially preventable readmissions are a subset of all-cause readmissions. Although there is no data collection burden associated with these claims-based measures, we recognize that having two hospital readmission measures in the LTCH QRP may create confusion. We agree with commenters that there is overlap between the All-Cause Readmission measure and the PPR 30-Day Post-Discharge measure, which identifies a subset of all-cause readmissions, and believe the PPR measure will be more actionable for quality improvement.

We are proposing to remove the All-Cause Unplanned Readmission measure beginning with the FY 2019 LTCH QRP. We are proposing that public reporting of this measure would end by October 2018 when public reporting of the PPR
In this proposed rule, we are also soliciting public comments on the use of survey-based experience of care measures for the LTCH QRP. We are currently developing an experience of care survey for LTCHs and survey-based measures will be developed from this survey. These survey-based measures may be considered for inclusion in the LTCH QRP through future notice-and-comment rulemaking. This survey was developed using a rigorous survey development methodology that included a public request for measures titled Request for Information To Aid in the Design and Development of a Survey Regarding Patient and Family Member Experiences With Care Received in Long-Term Care Hospitals (80 FR 72722 through 72725); focus groups and interviews with patients, family members, and caregivers; input from a TEP of LTCHs, researchers, and patient advocates; and cognitive interviewing. The survey has also been field tested. The survey explores experience of care across five main areas: (1) Beginning stay at the hospital; (2) interactions with staff; (3) experience during the hospital stay; (4) preparing for leaving hospital; and (5) overall hospital rating. We are specifically interested in comments regarding survey implementation and logistics, use of the survey-based measures in the LTCH QRP, and general feedback.

Also, we are considering a measure focused on pain that relies on the collection of patient-reported pain data, and another measure that documents whether a patient has an Advance Care Plan. Finally, we are considering a measure related to patient safety, specifically, Patients Who Received an Antipsychotic Medication. We are inviting public comment on the possible inclusion of such measures in future years of the LTCH QRP.

b. IMPACT Act Measure—Possible Future Update to Measure Specifications

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57207 through 57215), we finalized the Discharge to Community–PAC LTCH QRP measure, which assesses successful discharge to the community from an LTCH setting, with successful discharge to the community including no unplanned rehospitalizations and no death in the 31 days following discharge from the LTCH. We received public comments (see 81 FR 57211) recommending exclusion of baseline nursing facility residents from the measure, as these residents did not live in the community prior to their LTCH stay. At that time, we highlighted that, using Medicare FFS claims alone, we were unable to accurately identify baseline nursing facility residents. We stated that potential future modifications of the measure could include assessment of the feasibility and impact of excluding baseline nursing facility residents from the measure through the addition of patient assessment-based data. In response to these public comments, we are considering a future modification of the Discharge to Community–PAC LTCH QRP measure, which would exclude baseline nursing facility residents from the measure.

We are inviting public comment on the possibility of excluding baseline nursing facility residents from the Discharge to Community–PAC LTCH QRP measure in future years of the LTCH QRP.

c. IMPACT Act Implementation Update

As a result of the input and suggestions provided by technical experts at the TEPs held by our measure developer, and through public comment, we are engaging in additional development work, including performing additional testing, with respect to two measures that would satisfy the domain of accurately communicating the existence of and providing for the transfer of health information and care preferences when the individual transitions, in section 1899B(c)(1)(E) of the Act. The measures under development are: Transfer of Information at Post-Acute Care Admission, Start or Resumption of Care from other Providers/Settings; and Transfer of Information at Post-Acute Care Discharge, and End of Care to other Providers/Settings. We intend to specify these measures under section 1899B(c)(1)(E) of the Act no later than October 1, 2018, and we intend to propose to adopt them for the FY 2021 LTCH QRP, with data collection beginning on or about April 1, 2019.
10. Proposed Standardized Patient Assessment Data Reporting for the LTCH QRP

a. Proposed Standardized Patient Assessment Data Reporting for the FY 2019 LTCH QRP

Section 1886(m)(5)(F)(ii) of the Act requires that for fiscal year 2019 and each subsequent year, LTCHs report standardized patient assessment data required under section 1899B(b)(1) of the Act. As we describe in more detail above, we are proposing that the current pressure ulcer measure, Percent of Residents or Patients with Pressure Ulcers That Are New or Worsened (Short Stay) (NQF #0678), be replaced with the proposed pressure ulcer measure, Changes in Skin Integrity Post-Acute Care: Pressure Ulcer/Injury, beginning with the FY 2020 LTCH QRP. The current pressure ulcer measure will remain in the LTCH QRP until that time. Accordingly, with respect to the required standardized patient assessment data for the FY 2019 LTCH QRP, we are proposing that the data elements used to calculate that measure meet the definition of standardized patient assessment data with respect to medical conditions and co-morbidities under section 1899B(b)(1)(B)(iv) of the Act, and that the successful reporting of that data under section 1886(m)(5)(F)(i) of the Act with respect to admissions as well as discharges occurring during last three quarters of CY 2017 would also satisfy the requirement to report standardized patient assessment data for the FY 2019 LTCH QRP.

The collection of assessment data pertaining to skin integrity, specifically pressure related wounds, is important for multiple reasons. Clinical decision support, care planning, and quality improvement all depend on reliable assessment data collection. Pressure related wounds represent poor outcomes, are a serious medical condition that can result in death and disability, are debilitating, painful and are often an avoidable outcome of medical care.392 393 394 395 396 397

Pressure related wounds are considered healthcare acquired conditions. As we note above, the data elements needed to calculate the current pressure ulcer measure are already included on the LTCH CARE Data Set and reported by LTCHs, and exhibit validity and reliability for use across PAC providers. Item reliability for these data elements was also tested for the nursing home setting during implementation of MDS 3.0. Testing results are from the RAND Development and Validation of MDS 3.0 project.398 The RAND pilot test of the MDS 3.0 data elements showed good reliability and is also applicable to both the IRF–PAI and the LTCH CARE Data Set because the data elements tested are the same. Across the pressure ulcer data elements, the average gold-standard nurse to gold-standard nurse kappa statistic was 0.905. The average gold-standard nurse to facility-nurse kappa statistic was 0.937. Data elements used to risk adjust this quality measure were also tested under this same pilot test, and the gold-standard to gold-standard kappa statistic, or percent agreement (where kappa statistic not available), ranged from 0.91 to 0.99 for these data elements. These kappa scores indicate “almost perfect” agreement using the Landis and Koch standard for strength of agreement.399

The data elements used to calculate the current pressure ulcer measure received public comment on several occasions, including when that measure was proposed in the FY 2012 IRF PPS (76 FR 47876) and IPPS/LTC PPS proposed rules (76 FR 51574). Further, they were discussed in the past by TEPs held by our measure development contractor on June 13 and November 15, 2013, and recently by a TEP on July 18, 2016. TEP members supported the measure and its cross-setting use in PAC. The report, Technical Expert Panel Summary Report: Refinement of the Percent of Patients or Residents with Pressure Ulcers that are New or Worsened (Short Stay) (NQF #0678) Quality Measure for Skilled Nursing Facilities (SNFs), Inpatient Rehabilitation Facilities (IRFs), Long-Term Care Hospitals (LTCHs), and Home Health Agencies (HHAs), is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

We are inviting public comment on this proposal.

b. Proposed Standardized Patient Assessment Data Reporting Beginning With the FY 2020 LTCH QRP

We describe below our proposals for the reporting of standardized patient assessment data by LTCHs beginning with the FY 2020 LTCH QRP. LTCHs would be required to report these data with respect to LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018, with the exception of three data elements (Brief Interview of Mental Status (BIMS), Hearing, and Vision) that would be required with respect to LTCH admissions only that occur between April 1, 2018 and December 31, 2018. The BIMS, Hearing, and Vision data elements would be assessed at admission only due to the relatively stable nature of the types of cognitive function, hearing impairment, and vision impairment, making it unlikely that these assessments would change between the start and end of the PAC stay. Assessment of the BIMS, Hearing, and Vision data elements at discharge would introduce additional burden without improving the quality or usefulness of the data, and is unnecessary. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting.

In selecting the data elements proposed below, we carefully weighed the balance of burden in assessment-based data collection and aimed to minimize additional burden through the utilization of existing data in the assessment instruments. We also took into consideration the following factors with respect to each data element: overall clinical relevance; ability to support clinical decisions, care planning and interoperable exchange to facilitate care coordination during transitions in care; and the ability to capture medical complexity and risk factors that can inform both payment and quality. In addition, the data elements had to have strong scientific reliability and validity; be meaningful enough to inform longitudinal analysis by providers; had to have received general consensus agreement for its usability; and had to have the ability to collect such data once but support multiple uses. Further,
to inform the final set of data elements for proposal, we took into account technical and clinical subject matter expert review, public comment, and consensus input in which such principles were applied. We also took into account the consensus work and empirical findings from the PAC–PRD.

Below, we discuss the proposed standardized patient assessment data by category.

(1) Functional Status Data

We are proposing that the data elements currently reported by LTCHs to calculate the measure, Application of Percent of Long-Term Care Hospital Patients with an Admission and Discharge Functional Assessment and a Care Plan That Addresses Function (NQF #2631), would also meet the definition of standardized patient assessment data with respect to functional status under section 1899B(b)(1)(B)(ii) of the Act and that the successful reporting of that data under section 1886(m)(5F)(ii) of the Act would also satisfy the requirement to report standardized patient assessment data under section 1886(m)(5F)(ii) of the Act.

These patient assessment data for functional status are from the CARE Item Set. The development of the CARE Item Set and a description and rationale for each item is described in a report entitled “The Development and Testing of the Continuity Assessment Record and Evaluation (CARE) Item Set: Final Report on the Development of the CARE Item Set: Volume 1 of 3.”

(2) Cognitive Function and Mental Status Data

Cognitive function and mental status in PAC patient and resident populations can be affected by a number of underlying conditions, including dementia, stroke, traumatic brain injury, side effects of medication, metabolic and/or endocrine imbalances, delirium, and depression.

The assessment of cognitive function and mental status by PAC providers is important because of the high prevalence of patients and residents with these conditions, and the opportunity for improving the quality of care. Symptoms of dementia may improve with pharmacotherapy, occupational therapy, or physical activity.

We are proposing that the data elements that comprise the Brief Interview for Mental Status meet the definition of standardized patient assessment data with respect to cognitive function and mental status (for example, delirium), anticipating the patient or resident’s ability to understand and participate in treatments during a PAC stay, ensuring patient and resident safety (for example, risk of falls), and identifying appropriate support needs at the time of discharge or transfer. Standardized assessment data elements will enable or support clinical decision-making and early clinical intervention; person-centered, high quality care through: Facilitating better care continuity and coordination; better data exchange and interoperability between settings; and longitudinal outcome analysis. Hence, reliable data elements assessing cognitive impairment and mental status are needed in order to initiate a management program that can optimize a patient or resident’s prognosis and reduce the possibility of adverse events. 

• Brief Interview for Mental Status (BIMS)

We are proposing that the data elements that comprise the Brief Interview for Mental Status meet the definition of standardized patient assessment data with respect to cognitive function and mental status under section 1899B(b)(1)(B)(ii) of the Act. The proposed data elements consist of seven BIMS questions that result in a cognitive function score. For more information on the BIMS, we refer readers to the document titled, "Proposed Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/CARE-Item-Set-and-B-CARE.html."

For more information about this quality measure and the data elements used to calculate it, we refer readers to the FY 2016 IPPS/LTCH PPS final rule (80 FR 49739 through 49774).

We are inviting public comment on this proposal.


Dementia and cognitive impairment are associated with long-term functional dependence and, consequently, poor quality of life and increased health care costs and mortality.414 This makes assessment of mental status and early detection of cognitive decline or impairment critical in the PAC setting. The burden of cognitive impairment in PAC is high. The intensity of routine nursing care is higher for patients and residents with cognitive impairment than those without, and dementia is a significant variable in predicting readmission after discharge to the community from PAC providers.415 The BIMS is a performance-based cognitive assessment that assesses repetition, recall with and without prompting, and temporal orientation. It was developed to be a brief screener to assess cognition, with a focus on learning and memory. The BIMS data elements are currently in use in two of the PAC assessments: The MDS 3.0 in SNFs and the IRF-PAI in IRFs. The BIMS was tested in the PAC PRD where it was found to have substantial to almost perfect agreement for inter-rater reliability (kappa range of 0.71 to 0.91) when tested in all four PAC settings.416 Clinical and subject matter expert advisors working with our data element contractor agreed that the BIMS is a feasible data element for use by PAC providers. In addition, discussions during a TEP convened on April 6 and 7, 2016, demonstrated support for the BIMS. The Development and Maintenance of Post-Acute Care Cross-Setting Standardized Patient Assessment Data Technical Expert Panel Summary Report is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

To solicit additional feedback on the BIMS, we asked for public comment from August 12 to September 12, 2016. Many commenters expressed support for use of the BIMS, noting that it is reliable, feasible to use across settings, and will provide useful information about patients and residents. These comments noted that the data collected through the BIMS will provide a clearer picture of patient or resident complexity, help with the care planning process, and be useful during care transitions and when coordinating across providers. A full report of the comments is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

Therefore, we are proposing to adopt the BIMS for use in the LTCH QRP. We are proposing to add the data elements that comprise the BIMS to the LCDS, and that LTCHs would be required to report these data for the FY 2020 LTCH QRP with respect to LTCH admissions that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting. The BIMS data element would be assessed at admission only due to the relatively stable nature of the types of cognitive function assessed by the BIMS, making it unlikely that a patient’s score on this assessment would change between the start and end of the PAC stay. Assessment at discharge would introduce additional burden without improving the quality or usefulness of the data, and we believe that it is unnecessary.

We are inviting public comment on these proposals.

- Confusion Assessment Method (CAM)

We are proposing that the data elements that comprise the Confusion Assessment Method (CAM) meet the definition of standardized patient assessment data with respect to cognitive function and mental status under section 1899B(b)(1)[B][iii] of the Act. The CAM is a six-question instrument that screens for overall cognitive impairment, as well as distinguishes delirium or reversible confusion from other types of cognitive impairment. For more information on the CAM, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html. The CAM was developed to identify the signs and symptoms of delirium. It results in a score that suggests whether the patient or resident should be assigned a diagnosis of delirium. Because patients and residents with multiple comorbidities receive services from PAC providers, it is important to assess delirium, which is associated with a high mortality rate and prolonged duration of stay in hospitalized older adults.417 Assessing these signs and symptoms of delirium is clinically relevant for care planning by PAC providers.

The CAM is currently in use in two of the PAC assessments: the MDS 3.0 in SNFs and the LCDS in LTCHs. The CAM was tested in the PAC PRD where it was found to have substantial agreement for inter-rater reliability for the “Inattention and Disorganized Thinking” questions (kappa range of 0.70 to 0.73); and moderate agreement for the “Altered Level of Consciousness” question (kappa of 0.54).418 Clinical and subject matter expert advisors working with our data element contractor agreed that the CAM is feasible for use by PAC providers, that it assesses key aspects of cognition, and that this information about patient or resident cognition would be clinically useful both within and across PAC provider types. The CAM was also supported by a TEP that discussed and rated candidate data elements during a meeting on April 6 and 7, 2016. The Development and Maintenance of Post-Acute Care Cross-Setting Standardized Patient Assessment Data Technical Expert Panel Summary Report is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html. We asked for public comment on the CAM from August 12 to September 12, 2016. Many commenters expressed support for use of the CAM, noting that it would provide important information for care planning and care coordination, and therefore, contribute to quality improvement. The commenters noted it is particularly helpful in distinguishing


As noted above, the CAM is already included on the LCDS. For purposes of reporting for the FY 2020 LTCH QRP, LTCHs would be required to report these data with respect to LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting.

We are inviting public comment on these proposals.

- Behavioral Signs and Symptoms

We are proposing that the Behavioral Signs and Symptoms data elements meet the definition of standardized patient assessment data with respect to cognitive function and mental status under section 1899B(b)(1)(B)(ii) of the Act. The proposed data elements consist of three Behavioral Signs and Symptoms questions and result in three scores that categorize respondents as having or not having certain types of behavioral signs and symptoms. For more information on the Behavioral Signs and Symptoms data elements, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Insitutes/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

The questions included in the Behavioral Signs and Symptoms group assess whether the patient or resident has exhibited any behavioral symptoms that may indicate cognitive impairment or other mental health issues during the assessment period, including physical, verbal, and other disruptive or dangerous behavioral symptoms, but excluding patient wandering. Such behavioral disturbances can indicate unrecognized needs and care preferences and are associated most commonly with dementia and other cognitive impairment, and less commonly with adverse drug events, mood disorders, and other conditions. Assessing behavioral disturbances can lead to early intervention, patient- and resident-centered care planning, clinical decision support, and improved staff and patient or resident safety through early detection. Assessment and documentation of these disturbances can help inform care planning and patient transitions and provide important information about resource use.

Data elements that capture behavioral symptoms are currently included in two of the PAC assessments: the MDS 3.0 in SNFs and the OASIS–C2 in HHAs. In the MDS, each question includes four response options ranging from “behavior not exhibited” (0) to behavior “occurred daily” (3). The OASIS–C2 includes some similar data elements which record the frequency of disruptive behaviors on a 6-point scale ranging from “never” (0) to “at least daily” (5). Data elements that mirror those used in the MDS and serve the same assessment purpose were tested in post-acute providers in the PAC PRD and found to be clinically relevant, meaningful for care planning, and feasible for use in each of the four PAC settings.

The proposed data elements were supported by comments from the Standardized Patient Assessment Data TEP held by our data element contractor. The TEP identified patient and resident behaviors as an important consideration for resource intensity and care planning, and affirmed the importance of the standardized assessment of patient behaviors through data elements such as those in use in the MDS. The Development and Maintenance of Post-Acute Care Cross-Setting Standardized Patient Assessment Data Technical Expert Panel Summary Report is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Insitutes/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

Because the PAC PRD version of the Behavioral Signs and Symptoms data elements were previously tested across PAC providers, we solicited additional feedback on this version of the data elements by including these data elements in a call for public comment that was open from August 12 to September 12, 2016. Consistent with the TEP discussion on the importance of patient and resident behaviors, many commenters expressed support for use of the Behavioral Signs and Symptoms data elements, noting that they would provide useful information about patient and resident behavior at both admission and discharge and contribute to care planning related to what treatment is appropriate for the patient or resident and what resources are needed. Public comment also supported the use of highly similar MDS version of the data element in order to provide continuity with existing assessment processes in SNFs. A full report of the comments is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Insitutes/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

Therefore, we are proposing the MDS version of the Behavioral Signs and Symptoms data elements because they focus more closely on behavioral symptoms than the OASIS data elements, and include more detailed response categories than those used in the PAC PRD version, capturing more information about the frequency of behaviors. We are proposing to add the Behavioral Signs and Symptoms data elements to the LCDS, and that LTCHs would be required to report these data for the FY 2020 LTCH QRP with respect to LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting.

We are inviting public comment on these proposals.

- Patient Health Questionnaire-2 (PHQ–2)

We are proposing that the PHQ–2 data elements meet the definition of standardized patient assessment data with respect to cognitive function and mental status under section 1899B(b)(1)(B)(ii) of the Act. The proposed data elements consist of the PHQ–2 two-item questionnaire that assesses the cardinal criteria for depression: depressed mood and anhedonia (inability to feel pleasure). For more information on the PHQ–2, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Insitutes/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

Depression is a common mental health condition often missed and under-recognized. Assessments of depression help PAC providers better understand the needs of their patients

and residents by: Prompting further evaluation (that is, to establish a diagnosis of depression); elucidating the patient’s or resident’s ability to participate in therapies for conditions other than depression during their stay; and identifying appropriate ongoing treatment and support needs at the time of discharge. A PHQ–2 score beyond a predetermined threshold signals the need for additional clinical assessment in order to determine a depression diagnosis.

The proposed data elements that comprise the PHQ–2 are currently used in the OASIS–C2 for HHAs and the MDS 3.0 for SNFs (as part of the PHQ–9). The PHQ–2 data elements were tested in the PAC PRD, where they were found to have almost perfect agreement for inter-rater reliability (kappa range of 0.84 to 0.91) when tested by all four PAC providers.421

Clinical and subject matter expert advisors working with our data element contractor agreed that the PHQ–2 is feasible for use in PAC, that it assesses key aspects of mental status, and that this information about patient or resident mood would be clinically useful both within and across PAC provider types. We note that both the PHQ–9 and the PHQ–2 were supported by TEP members who discussed and rated candidate data elements during a meeting on April 6 and 7, 2016. They particularly noted that the brevity of the PHQ–2 made it feasible with low burden for both assessors and PAC patients or residents. The Development and Maintenance of Post-Acute Care Cross-Setting Standardized Patient Assessment Data Technical Expert Panel Summary Report is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

To solicit additional feedback on the PHQ–2, we asked for public comment from August 12 to September 12, 2016. Many commenters provided feedback on using the PHQ–2 for the assessment of mood. Overall, commenters believed that collecting these data elements across PAC provider types was appropriate, given the role that depression plays in well-being. Several commenters expressed support for an approach that would use PHQ–2 as a gateway to the longer PHQ–9 and would maintain the reduced burden on most patients and residents, as well as test administrators, which is a benefit of the PHQ–2, while ensuring that the PHQ–9, which exhibits higher specificity,422 would be administered for patients and residents who showed signs and symptoms of depression on the PHQ–2.


Therefore, we are proposing to add the PHQ–2 data elements to the LCDS, and that LTCHs would be required to report these data for the FY 2020 LTCH QRP with respect to LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting.

We are inviting public comment on these proposals.

(3) Special Services, Treatments, and Interventions Data

Special services, treatments, and interventions performed in PAC can have a major effect on an individual’s health status, self-image, and quality of life. The assessment of these special services, treatments, and interventions in PAC is important to ensure the continuing appropriateness of care for the patients and residents receiving them, and to support care transitions from one PAC provider to another, an acute care hospital, or discharge. Accurate assessment of special services, treatments, and interventions of patients and residents served by PAC providers are expected to have a positive impact on the National Quality Strategy’s domains of patient and family engagement, patient safety, care coordination, clinical process/ effectiveness, and efficient use of health care resources.

For example, standardized assessment of special services, treatments, and interventions used in PAC can promote patient and resident safety through appropriate care planning (for example, mitigating risks such as infection or pulmonary embolism associated with central intravenous access), and identifying life-sustaining treatments that must be continued, such as mechanical ventilation, dialysis, suctioning, and chemotherapy, at the time of discharge or transfer.

Standardized assessment of these data elements will enable or support: Clinical decision-making and early clinical intervention; person-centered, high quality care through, for example, facilitating better care continuity and coordination; better data exchange and interoperability between settings; and longitudinal outcome analysis. Hence, reliable data elements assessing special services, treatments, and interventions are needed to initiate a management program that can optimize a patient or resident’s prognosis and reduce the possibility of adverse events.

We are proposing 15 special services, treatments, and interventions as presented below grouped by cancer treatments, respiratory treatments, other treatments, and nutritional approaches. A TEP convened by the data element standardization contractor provided input on the 15 data elements assessing for Special Services, Treatments, and Interventions. This TEP, held on January 5 and 6, 2017, opined that these data elements are appropriate for standardization because they would provide useful clinical information to inform care planning and care coordination. The TEP affirmed that assessment of these services and interventions is standard clinical practice, and that the collection of these data by means of a list and checklist format would conform with common workflow for PAC providers. A full report of the TEP discussion is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

- Cancer Treatment: Chemotherapy (IV, Oral, Other)

We are proposing that the Chemotherapy (IV, Oral, Other) data elements meet the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. The proposed data elements consist of the principal Chemotherapy data element and three sub-elements: IV Chemotherapy, Oral Chemotherapy, and Other.

For more information on the Chemotherapy data element, we refer readers to the document titled, Proposed Specifications for LTC QRP Quality Measures and Standardized Data Elements, available at: https://

As a result of the comments and input received from clinical and subject matter experts, we are proposing a principal Chemotherapy data element with three sub-elements, including Oral and Other for standardization. Our data element contractor then presented the proposed data elements to the Standardized Patient Assessment Data TEP on January 5 and 6, 2017, who supported these data elements for standardization. A full report of the TEP discussion is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-Downloads-and-Videos.html.

Therefore, we are proposing that the Chemotherapy (IV, Oral, Other) data elements with a principal data element and three sub-elements meet the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. We are proposing to add the Chemotherapy (IV, Oral, Other) data elements to the LCDS, and that LTCHs would be required to report these data for the FY 2020 LTCH QRP with respect to LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting.

We are inviting public comment on these proposals.

- Cancer Treatment: Radiation

We are proposing that the Radiation data element meets the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. The proposed data element consists of the single Radiation data element. For more information on the Radiation data element, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

Radiation is a type of cancer treatment that uses high-energy radioactivity to stop cancer by damaging cancer cell DNA, but it can also damage normal cells. Radiation is an important therapy for particular types of cancer, and the resource utilization is high, with frequent radiation sessions required, often daily for a period of several weeks. Assessing whether a patient or resident is receiving radiation therapy is important to determine resource utilization because PAC patients and residents will need to be transported to and from radiation treatments, and monitored and treated for side effects after receiving this intervention.

Therefore, assessing the receipt of radiation therapy, which would compete with other care processes given the time burden, would be important for care planning and care coordination by PAC providers.

The Radiation data element is currently in use in the MDS 3.0. This data element was not tested in the PAC PRD. However, public comment and other expert input on the Radiation data element supported its importance and clinical usefulness for patients in PAC settings, due to the side effects and consequences of radiation treatment on patients that need to be considered in care planning and care transitions. To solicit additional feedback on the Radiation data element we are proposing, we asked for public comment from August 12 to September 12, 2016. Several commenters provided support for the data element, noting the relevance of this data element to facilitating care coordination and supporting care transitions, the feasibility of the item, and the potential for it to improve quality. A full report of the comments is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.
The proposed data element was presented to and supported by the TEP held by our data element contractor on January 5–6, 2017, which opined that Radiation was important corollary information about cancer treatment to collect alongside Chemotherapy (IV, Oral, Other), and that, because capturing this information is a customary part of clinical practice, the proposed data element would be feasible, reliable, and easily incorporated into existing workflow.

Therefore, we are proposing that the Radiation data element meets the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. We are proposing to add the Radiation data element to the LCDS, and that LTCHs would be required to report these data for the 2020 LTCH QRP with respect to LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting.

We are inviting public comment on these proposals.

Respiratory Treatment: Oxygen Therapy (Continuous, Intermittent)

We are proposing that the Oxygen Therapy (Continuous, Intermittent) data elements meet the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. The proposed data elements consist of the principal data element and two sub-elements, “Continuous” and “As needed”.

Therefore, we are proposing that the Oxygen Therapy (Continuous, Intermittent) data elements to the LCDS, and that LTCHs would be required to report these data for the FY 2020 LTCH QRP with respect to LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting.

We are inviting public comment on these proposals.

Respiratory Treatment: Suctioning (Scheduled, as Needed)

We are proposing that the Suctioning (Scheduled, As needed) data elements meet the definition of standardized patient assessment data element with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. The proposed data elements consist of the principal Suctioning data element, and two sub-elements, “Scheduled” and “As needed.” These sub-elements capture two types of suctioning: “Scheduled” indicates suctioning based on a specific frequency, such as every hour: “As needed” means suctioning only when indicated. For more information on the Suctioning (Scheduled, As needed) data elements, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

Suctioning is a process used to clear secretions from the airway when a person cannot clear those secretions on his or her own. It is done by aspirating secretions through a catheter connected to a suction source. Types of suctioning include oropharyngeal and nasopharyngeal suctioning, nasotracheal suctioning, and suctioning through an artificial airway such as a tracheostomy tube. Oropharyngeal and nasopharyngeal suctioning are a key part of many patients’ care plans, both to prevent the accumulation of secretions that can lead to aspiration pneumonias (a common condition in patients with inadequate gag reflexes), and to relieve obstructions from mucus plugging during an acute or chronic respiratory infection, which often lead to desaturations and increased respiratory effort. Suctioning can be done on a scheduled basis if the patient is judged to clinically benefit from regular interventions; or can be done as needed, such as when secretions become so prominent that gurgling or choking is noted, or a sudden
A TEP convened by the data element contractor provided input on the proposed data elements. This TEP, held on January 5 and 6, 2017, opined that these data elements are appropriate for standardization because they would provide useful clinical information to inform care planning and care coordination. The TEP affirmed that assessment of these services and interventions is standard clinical practice. A full report of the TEP discussion is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

Therefore, we are proposing that the Suctioning (Scheduled, As needed) data elements with a principal data element and two sub-elements meet the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. We are proposing to add the Suctioning (scheduled, as needed) data element to the LCDS, and that LTCHs would be required to report these data for the FY 2020 LTCH QRP with respect to LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting.

We are inviting public comment on these proposals.

- Respiratory Treatment: Tracheostomy Care

We are proposing that the Tracheostomy Care data element meets the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. The proposed data element consists of the single Tracheostomy Care data element. For more information on the Tracheostomy Care data element, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

A tracheostomy provides an air passage to help a patient or resident breathe when the usual route for breathing is impaired or impaired. Generally, in all of these cases, suctioning is necessary to ensure that the tracheostomy is clear of secretions which can inhibit successful oxygenation of the individual. Often, individuals with tracheostomies are also receiving supplemental oxygenation. The presence of a tracheostomy, albeit permanent or temporary, warrants careful monitoring and immediate intervention if the tracheostomy becomes occluded or in the case of a temporary tracheostomy, the device used becomes dislodged. While in rare cases the presence of a tracheostomy is not associated with increased care demands (and in some of those instances, the care of the ostomy is performed by the patient) in general the presence of such a device is associated with increased patient risk, and clinical care services will necessarily include close monitoring to ensure that no life-threatening events occur as a result of the tracheostomy, often considered part of the patient’s life line. In addition, tracheostomy care, which primarily consists of cleansing, dressing changes, and replacement of the tracheostomy cannula (tube), is also a critical part of the care plan. Regular cleansing is important to prevent infection such as pneumonia and to prevent any occlusions with which there are risks for inadequate oxygenation.

The proposed data element is currently in use in the MDS 3.0 (“Tracheostomy Care”). Data elements (“Trach Tube with Suctioning”) that were tested in the PAC PRD included an equivalent principal data element on the presence of a tracheostomy. This data element was found feasible for use in each of the four PAC settings as the data collection aligned with usual work flow.

Clinical and subject matter expert advisors working with our data element contractor agreed that the Tracheostomy Care data element is feasible for use in PAC and that it assesses an important treatment that would be clinically useful both within and across PAC providers. We solicited public comment on the suctioning data element currently included in the MDS 3.0 between August 12 to September 12, 2016. Several commenters wrote in support of this data element, noting feasibility of this item in PAC, and the relevance of this data element to facilitating care coordination and supporting care transitions. We also received comments suggesting that we examine the frequency of suctioning in order to better understand the use of staff time, the impact on a patient or resident’s capacity to speak and swallow, and intensity of care required. Based on these comments, we decided to add two sub-elements (scheduled and as needed) to the suctioning element. The proposed data elements, Suctioning (Scheduled, As needed) includes both the principal suctioning data element that is included on the MDS 3.0 and two sub-elements, “scheduled” and “as needed.” A full report of the comments is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.
proposed data elements. This TEP, held on January 5 and 6, 2017, opined that these data elements are appropriate for standardization because they would provide useful clinical information to inform care planning and care coordination. The TEP affirmed that assessment of these services and interventions is standard clinical practice. A full report of the TEP discussion is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

Therefore, we are proposing that the Tracheostomy Care data element meets the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. We are proposing to add the Tracheostomy Care data element to the LCDS, and that LTCHs would be required to reporting these data for the FY 2020 LTCH QRP with respect to LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting.

We are inviting public comment on these proposals.

- **Respiratory Treatment: Non-invasive Mechanical Ventilator (BiPAP, CPAP)**

  We are proposing that the Non-invasive Mechanical Ventilator [BiPAP, Continuous Positive Airway Pressure [BiPAP], Continuous Positive Airway Pressure [CPAP]] data elements meet the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. The proposed data elements consist of the principal Non-invasive Mechanical Ventilator data element and two sub-elements, BiPAP and CPAP. For more information on the Non-invasive Mechanical Ventilator (BiPAP, CPAP) data element, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

  BiPAP and CPAP are respiratory support devices that prevent the airways from closing by delivering slightly pressurized air via electronic cycling throughout the breathing cycle (Bilevel PAP, referred to as BiPAP) or through a mask continuously (Continuous PAP, referred to as CPAP). Assessment of non-invasive mechanical ventilation is important in care planning, as both CPAP and BiPAP are resource-intensive (although less so than invasive mechanical ventilation) and signify underlying medical conditions about the patient or resident who requires the use of this intervention. Particularly when used in settings of acute illness or progressive respiratory decline, additional staff (for example, respiratory therapists) are required to monitor and adjust the CPAP and BiPAP settings and the patient or resident may require more nursing resources.

  Data elements that assess BiPAP and CPAP are currently included on the OASIS–C2 for HHAs (“Continuous/Bi-level positive airway pressure”), LCDS for the LTCH setting (“Non-invasive Ventilator (BiPAP, CPAP)”), and the MDS 3.0 for the SNF setting (“BiPAP/CPAP”). A data element that focused on CPAP was tested across the four PAC providers in the PAC–PRD study and found to be feasible for standardization. All of these data elements assess BiPAP or CPAP with a single check box, not separately.

  Clinical and subject matter expert advisors working with our data element contractor agreed that the standardized assessment of Non-invasive Mechanical Ventilator (BiPAP, CPAP) data elements would be feasible for use in PAC, and assess an important treatment that would be clinically useful both within and across PAC provider types.

  To solicit additional feedback on the form of the Non-invasive Mechanical Ventilator (BiPAP, CPAP) data elements best suited for standardization, we asked for public comment on a single data element, BiPAP/CPAP, equivalent (but for labeling) to what is currently in use on the MDS, OASIS, and LCDS, from August 12 to September 12, 2016. Several commenters wrote in support of this data element, noting the feasibility of these items in PAC, and the relevance of these data elements for facilitating care coordination and supporting care transitions. In addition, there was support in the public comment responses for separating out BiPAP and CPAP as distinct sub-elements, as they are therapies used for different types of patients and residents. A full report of the comments is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

  A TEP convened by the data element contractor provided input on the proposed data elements. This TEP, held on January 5 and 6, 2017, opined that these data elements are appropriate for standardization because they would provide useful clinical information to inform care planning and care coordination. The TEP affirmed that assessment of these services and interventions is standard clinical practice. A full report of the TEP discussion is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

  Therefore we are proposing that the Non-invasive Mechanical Ventilator (BiPAP, CPAP) data elements with a principal data element and two sub-elements meet the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. We are proposing to expand the existing “Non-invasive Ventilator (BiPAP, CPAP)” data element on the LCDS, by retaining and renaming the main data element to be Non-invasive Mechanical Ventilator and adding two sub-elements for BiPAP and CPAP. For the purposes of reporting for the FY 2020 LTCH QRP, LTCHs would be required to report the Non-invasive Mechanical Ventilator (BiPAP, CPAP) data elements with respect to LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting.

  We are inviting public comment on these proposals.

- **Respiratory Treatment: Invasive Mechanical Ventilator**

  We are proposing that the Invasive Mechanical Ventilator data element meets the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. The proposed data element consists of a single Invasive Mechanical Ventilator data element which, for LTCHs, will be collected from the Invasive Mechanical Ventilator (Weaning) and Invasive Mechanical Ventilator (Non-Weaning) data elements that are already included on the LCDS. For more information on the Invasive Mechanical Ventilator data element, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.
Invasive mechanical ventilation includes ventilators and respirators that ventilate the patient through a tube that extends via the oral airway into the pulmonary region or through a surgical opening directly into the trachea. Thus, assessment of invasive mechanical ventilation is important in care planning and risk mitigation. Ventilation in this manner is a resource-intensive therapy associated with life-threatening conditions without which the patient or resident would not survive. However, ventilator use has inherent risks requiring close monitoring. Failure to adequately care for the patient or resident who is ventilator dependent can lead to iatrogenic events such as death, pneumonia and sepsis. Mechanical ventilation further signifies the complexity of the patient’s underlying medical and or surgical condition. Of note, invasive mechanical ventilation is associated with high daily and aggregate costs.

Data elements that capture invasive mechanical ventilation, but vary in their level of specificity, are currently in use in the MDS 3.0 (“Ventilator or respirator”) and LCDS (“Invasive Mechanical Ventilator: weaning” and “Invasive Mechanical Ventilator: non-weaning”). Data elements that assess invasive ventilator use and weaning status were tested in the PAC PRD (“Ventilator—Weaning” and “Ventilator—Non-Weaning”) and found feasible for use in each of the four PAC settings.

Clinical and subject matter expert advisors working with our data element contractor agreed that assessing Invasive Mechanical Ventilator use is feasible in PAC, and would be clinically useful both within and across PAC providers. To solicit additional feedback on the form of a data element on this topic that would be appropriate for standardization, data elements that assess invasive ventilator use and weaning status that were tested in the PAC PRD (“Ventilator—Weaning” and “Ventilator—Non-Weaning”) were included in a call for public comment that was open from August 12 to September 12, 2016 because it was being considered for standardization. Several commenters wrote in support of this data element, highlighting the importance of this information in supporting care coordination and care transitions. Some commenters expressed concern about the appropriateness for standardization, given the prevalence of ventilator weaning across PAC providers; the timing of administration; how weaning is defined; and how it weaning status in particular relates to quality of care. These comments guided the decision to propose single data element focused on current use of invasive mechanical ventilation only, and does not attempt to capture weaning status. A full report of the comments is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-Downloads-and-Videos.html.

A TEP convened by the data element contractor provided input on the proposed data elements. This TEP, held on January 5 and 6, 2017, opined that these data elements are appropriate for standardization because they would provide useful clinical information to inform care planning and care coordination. The TEP affirmed that assessment of these services and interventions is standard clinical practice. A full report of the TEP discussion is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-Downloads-and-Videos.html.

Therefore, we are proposing that the Invasive Mechanical Ventilator data element that assesses the use of an invasive mechanical ventilator, but does not assess weaning status, meets the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. This principal IV Medications data element (Antibiotics, Anticoagulation, Other) and three sub-elements, Antibiotics, Anticoagulation, and Other. For more information on the IV Medications (Antibiotics, Anticoagulation, Other) data element, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LLTC-Quality-Reporting/LLTC-Quality-Reporting-Measures-Information.html.

IV medications are solutions of a specific medication (for example, antibiotics, anticoagulants) administered directly into the venous circulation via a syringe or intravenous catheter (tube). IV medications are administered via intravenous push (bolus), single, intermittent, or continuous infusion through a tube placed into the vein (for example, commonly referred to as central, midline, or peripheral ports). Further, IV medications are more resource intensive to administer than oral medications, and signify a higher patient complexity (and often higher severity of illness).

The clinical indications for each of the sub-elements of the IV Medication data element (Antibiotics, Anticoagulants, and Other) are very different. IV antibiotics are used for severe infections when: (1) The bioavailability of the medication would be inadequate to kill the pathogen; (2) an oral form of the...
medication does not exist; or (3) the patient is unable to take the medication by mouth. IV anticoagulants refer to anti-clotting medications (that is, “blood thinners”), often used for the prevention and treatment of deep vein thrombosis and other thromboembolic complications. IV anticoagulants are commonly used in patients with limited mobility (either chronically or acutely, in the post-operative setting), who are at risk of deep vein thrombosis, or patients with certain cardiac arrhythmias such as atrial fibrillation. The indications, risks, and benefits of each of these classes of IV medications are distinct, making it important to assess each separately in PAC. Knowing whether or not patients are receiving IV medication and the type of medication provided by each PAC provider will improve quality of care.

The principal IV Medication data element is currently in use on the MDS 3.0 and there is a related data element in OASIS–C2 that collects information on Intravenous and Infusion Therapies. One sub-element of the proposed data element, IV Anti-coagulants, and two other data elements related to IV therapy (IV Vasoactive Medications and IV Chemotherapy), were tested in the PAC PRD and found feasible for use in that the data collection aligned with usual work flow in each of the four PAC settings, demonstrating the feasibility of collecting IV medication information, including type of IV medication, through similar data elements in these settings.

Clinical and subject matter expert advisors working with our data element contractor agreed that standardized collection of information on medications, including IV medications, would be feasible in PAC, and assess an important treatment that would be clinically useful both within and across PAC provider types.

We solicited public comment on a related data element, Vasoactive Medications, from August 12 to September 12, 2016. While commenters supported this data element with one noting the importance of this data element in supporting care transitions, others criticized the need for collecting specifically on Vasoactive Medications, giving feedback that the data element was too narrowly focused. In addition, comment received indicated that the clinical significance of vasoactive medications administration alone was not high enough in PAC to merit mandated assessment, noting that related and more useful information could be captured in an item that assessed all IV medication use.


A TEP convened by the data element contractor provided input on the proposed data elements. This TEP, held on January 5 and 6, 2017, opined that these data elements are appropriate for standardization because they would provide useful clinical information to inform care planning and care coordination. The TEP affirmed that assessment of these services and interventions is standard clinical practice. A full report of the TEP discussion is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

Therefore, we are proposing that the IV Medications (Antibiotics, Anticoagulation, Other) data elements with a principal data element and three sub-elements meet the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. We are proposing to add the IV Medications (Antibiotics, Anticoagulation, Other) data element to the LCDS, and that LTCHs would be required to report these data for the FY 2020 LTCH QRP with respect to LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting.

We are inviting public comment on these proposals.

- Other Treatment: Transfusions

We are proposing that the Transfusions data element meets the definition of standardized patient assessment data element with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. We are proposing to add the Transfusions data element to the LCDS, and that LTCHs would be required to report these data for the FY 2020 LTCH QRP with respect to LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting.

Transfusion refers to introducing blood, blood products, or other fluid into the circulatory system of a person. Blood transfusions are based on specific protocols, with multiple safety checks and monitoring required during and after the infusion in case of adverse events. Coordination with the provider’s blood bank is necessary, as well as documentation by clinical staff to ensure compliance with regulatory requirements. In addition, the need for transfusions signifies underlying patient complexity that is likely to require care coordination and patient monitoring, and impacts planning for transitions of care, as transfusions are not performed by all PAC providers.

The proposed data element was selected from three existing assessment items on transfusions and related services, currently in use in the MDS 3.0 (“Transfusions”) and OASIS–C2 (“Intravenous or Infusion Therapy”), and a data element tested in the PAC PRD (“Blood Transfusions”), that was found feasible for use in each of the four PAC settings. We chose to propose the MDS version because of its greater level of specificity over the OASIS–C2 data element. This selection was informed by expert advisors and reviewed and supported in the proposed form by the Standardized Patient Assessment Data TEP held by our data element contractor on January 5 and 6, 2017. A full report of the TEP discussion is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

Therefore, we are proposing that the Transfusions data element that is currently in use in the MDS meets the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. We are proposing to add the Transfusions data element to the LCDS, and that LTCHs would be required to report these data for the FY 2020 LTCH QRP with respect to LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting.
We are inviting public comment on these proposals.

- Other Treatment: Dialysis (Hemodialysis, Peritoneal Dialysis)

We are proposing that the Dialysis (Hemodialysis, Peritoneal dialysis) data elements meet the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. The proposed data elements consist of the principal Dialysis data element and two sub-elements, Hemodialysis and Peritoneal dialysis. For more information on the Dialysis (Hemodialysis, Peritoneal dialysis) data element, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

Dialysis is a treatment primarily used to provide replacement for lost kidney function. Both forms of dialysis (hemodialysis and peritoneal dialysis) are resource intensive, not only during the actual dialysis process but before, during and following. Patients and residents who need and undergo dialysis procedures are at high risk for physiologic and hemodynamic instability from fluid shifts and electrolyte disturbances as well as infections that can lead to sepsis. Further, patients or residents receiving hemodialysis are often transported to a different facility, or at a minimum, to a different location in the same facility. Close monitoring for fluid shifts, blood pressure abnormalities, and other adverse effects is required prior to, during and following each dialysis session. Nursing staff typically perform peritoneal dialysis at the bedside, and as with hemodialysis, close monitoring is required.

The principal Dialysis data element is currently included on the MDS 3.0 and the LCDS v3.0 and assesses the overall use of dialysis. The sub-elements for Hemodialysis and Peritoneal dialysis were tested across the four PAC providers in the PAC PRD study, and found to be feasible for standardization. Clinical and subject matter expert advisors working with our data element contractor opined that the standardized assessment of dialysis is feasible in PAC, and that it assesses an important treatment that would be clinically useful both within and across PAC providers. As the results of expert and public feedback, described below, we decided to propose a data element that includes both the principal Dialysis data element and the two sub-elements (hemodialysis and peritoneal dialysis).

The Hemodialysis data element, which was tested in the PAC PRD, was included in a call for public comment that was open from August 12 to September 12, 2016. Commenters supported the assessment of hemodialysis and recommended that the data element be expanded to include peritoneal dialysis. Several commenters supported the Hemodialysis data element, noting the relevance of this information for sharing across the care continuum to facilitate care coordination and care transitions, the potential for this data element to be used to improve quality, and the feasibility for use in PAC. In addition, we received comment that the item would be useful in improving patient and resident transitions of care. Several commenters also stated that peritoneal dialysis should be included in a standardized data element on dialysis and recommended collecting information on peritoneal dialysis in addition to hemodialysis. The rationale for including peritoneal dialysis from commenters included the fact that patients and residents receiving peritoneal dialysis will have different needs at post-acute discharge compared to those receiving hemodialysis or not having any dialysis. Based on these comments, the Hemodialysis data element was expanded to include a principal Dialysis data element and two sub-elements, hemodialysis and peritoneal dialysis; these are the same two data elements that were tested in the PAC PRD. This expanded version, Dialysis (Hemodialysis, Peritoneal dialysis), are the data elements being proposed. A full report of the comments is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

We note that the Dialysis (Hemodialysis, Peritoneal dialysis) data elements were also supported by the TEP that discussed candidate data elements for Special Services, Treatments, and Interventions during a meeting on January 5 and 6, 2017. A full report of the TEP discussion is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

Therefore, we are proposing that the Dialysis (Hemodialysis, Peritoneal dialysis) data elements with a principal data element and two sub-elements meet the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. We are proposing to expand the Dialysis data element in current use on the LCDS to include sub-elements for Hemodialysis and Peritoneal dialysis. For the purposes of reporting for the FY 2020 LTCH QRP, LTCHs would be required to report these data with respect to LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting.

We are inviting public comment on these proposals.

- Other Treatment: Intravenous (IV) Access (Peripheral IV, Midline, Central line, Other)

We are proposing that the IV Access (Peripheral IV, Midline, Central line, Other) data elements meet the definition of standardized patient assessment data element with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. The proposed data elements consist of the principal IV Access data element and four sub-elements, Peripheral IV, Midline, Central line, and Other. For more information on the IV Access data element, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

Patients or residents with central lines, including those peripherally inserted or who have subcutaneous central line “port” access, always require vigilant nursing care to keep patency of the lines and ensure that such invasive lines remain free from any potentially life-threatening events such as infection, air embolism, or bleeding from an open lumen. Clinically complex patients and residents are likely to be receiving medications or nutrition intravenously. The sub-elements included in the IV Access data elements distinguish between peripheral access and different types of central access. The rationale for distinguishing between a peripheral IV and central IV access is
that central lines confer higher risks associated with life-threatening events such as pulmonary embolism, infection, and bleeding.

The proposed IV Access (Peripheral IV, Midline, Central line, Other) data elements are not currently included on any of the mandated PAC assessment instruments. However, related data elements (for example, IV Medication in MDS 3.0 for SNF, Intravenous or infusion therapy in OASIS–C2 for HHAs) currently assess types of IV access. Several related data elements that describe types of IV access (for example, Central Line Management, IV Vasoactive Medications) were tested across the four PAC providers in the PAC PRD study, and found to be feasible for standardization.

Clinical and subject matter expert advisors working with our data element contractor agreed that assessing type of IV access would be feasible for use in PAC and that it assesses an important treatment that would be clinically useful both within and across PAC provider types.

We asked for public comment on one of the PAC PRD data elements, Central Line Management, from August 12 to September 12, 2016. A central line is one type of IV access. Commenters supported the assessment of central line management and recommended that the data element be broadened to also include other types of IV access. Several commenters supported the data element, noting feasibility and importance for facilitating care coordination and care transitions. However, a few commenters recommended that the definition of this data element be broadened to include peripherally inserted central catheters (“PICC lines”) and midline IVs. Based on public comment feedback and in consultation with clinical and subject matters experts, we expanded the Central Line Management data element to include more types of IV access (Peripheral IV, Midline, Central line, Other). This expanded version, IV Access (Peripheral IV, Midline, Central line, Other), are the data elements being proposed. A full report of the comments is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

We note that the IV Access (Peripheral IV, Midline, Central line, Other) data elements were supported by the TEP that discussed candidate data elements. Services, Treatments, and Interventions during a meeting on January 5 and 6, 2017. A full report of the TEP discussion is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

Therefore, we are proposing that the IV access (Peripheral IV, Midline, Central line, Other) data elements with a principal data element and four sub-elements meet the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. We are proposing to add the IV Access (Peripheral IV, Midline, Central line, Other) data elements to the LCDS and that LTCHs would be required to report these data for the FY 2020 LTCH QRP with respect to LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting. We are inviting public comment on these proposals.

- Nutritional Approach: Parenteral/IV Feeding

We are proposing that the Parenteral/IV Feeding data element meets the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. The proposed data element consists of the single Parenteral/IV Feeding data element. For more information on the Parenteral/IV Feeding data element, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

Parenteral/IV Feeding refers to a patient or resident being fed intravenously using an infusion pump, bypassing the usual process of eating and digestion. The need for IV/parenteral feeding indicates a clinical complexity that prevents the patient or resident from meeting his/her nutritional needs enterally, and is more resource intensive than other forms of nutrition, as it often requires monitoring of blood chemistries, and maintenance of a central line. Therefore, assessing a patient or resident’s need for parenteral feeding is important for care planning and resource use. In addition to the risks associated with central and peripheral intravenous access, total parenteral nutrition is associated with significant risks such as embolism and sepsis.

The Parenteral/IV Feeding data element is currently in use in the MDS 3.0, and equivalent or related data elements are in use in the LCDS, IRF–PAL, and the OASIS–C2. An equivalent data element was tested in the PAC PRD (“Total Parenteral Nutrition”) and found feasible for use in each of the four PAC settings, demonstrating the feasibility of collecting information about this nutritional service in these settings. Total Parenteral Nutrition (an item with the same meaning as the proposed data element, but with the label used in the PAC PRD) was included in a call for public comment that was open from August 12 to September 12, 2016. Several commenters supported this data element, noting its relevance to facilitating care coordination and supporting care transitions. After the public comment period, the Total Parenteral Nutrition data element was re-named Parenteral/IV Feeding, to be consistent with how this data element is referred to in the MDS. A full report of the comments is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

A TEP convened by the data element contractor provided input on the proposed data elements. This TEP, held on January 5 and 6, 2017, opined that these data elements are appropriate for standardization because they would provide useful clinical information to inform care planning and care coordination. The TEP affirmed that assessment of these services and interventions is standard clinical practice.


Therefore, we are proposing that the Parenteral/IV Feeding data element meets the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. We are proposing to retain and rename the existing and equivalent Total Parenteral Nutrition data element to be Parenteral/IV Feeding on the LCDS, and that LTCHs would be required to report these data for the FY 2020 LTCH QRP.
with respect to LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting.

We are inviting public comment on these proposals.

- **Clinical and subject matter expert advisors working with our data element contractor opined that the Feeding Tube data element is feasible for use in PAC, and supported its importance and clinical usefulness for patients in PAC settings, due to the increased level of nursing care and patient monitoring required for patients who received enteral nutrition with this device.**

We solicited additional feedback on an Enteral Nutrition data element (an item with the same meaning as the proposed data element, but with the label used in the OASIS) in a call for public comment that was open from August 12 to September 12, 2016. Several commenters supported the data element, noting the importance of assessing enteral nutrition status for facilitating care coordination and care transitions. After the public comment period, the Enteral Nutrition data element used in public comment was renamed Feeding Tube, indicating the presence of an assistive device. A full report of the comments is available at: [https://www.cms.gov/Medicare/Quality-Improvement-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html](https://www.cms.gov/Medicare/Quality-Improvement-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html).

We note that the Feeding Tube data element was also supported by the TEP that discussed candidate data elements for Special Services, Treatments, and Interventions during a meeting on January 5 and 6, 2017. A full report of the TEP discussion is available at: [https://www.cms.gov/Medicare/Quality-Improvement-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html](https://www.cms.gov/Medicare/Quality-Improvement-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html).


We are proposing that the Mechanically Altered Diet data element refers to food that has been altered to make it easier for the patient or resident to chew and swallow, and this type of diet is used for patients and residents who have difficulty performing these functions. Patients with severe malnutrition are at higher risk for a variety of complications.425 In PAC settings, there are a variety of reasons that patients and residents may have impairments related to oral feedings, including clinical or cognitive status. The provision of a mechanically altered diet may be resource intensive, and can signal difficulties associated with swallowing/eating safety, including dysphagia. In other cases, it signifies the type of altered food source, such as ground or puree, that will enable the safe and thorough ingestion of nutritional substances and ensure safe and adequate delivery of nourishment to the patient. Often, patients on mechanically altered diets also require additional nursing supports such as individual feeding, or direct observation, to ensure the safe consumption of the food product. Assessing whether a patient or resident requires a mechanically altered diet is therefore important for care planning and resource identification.

The proposed data element for a mechanically altered diet is currently included on the MDS 3.0 for SNFs, and in the OASIS–C2 for HHAs, where it is labeled Enteral Nutrition. A related data element, collected in the IRF–PAI for IRFs (Tube/Parenteral Feeding), assesses use of both feeding tubes and parenteral nutrition. The testing of similar nutrition-focused data elements in the PAC PRD, and the current assessment of feeding tubes and related nutritional services and devices, demonstrates the feasibility of collecting information about this nutritional service in these settings.

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information about independent eating that requires “a liquid, pureed or ground meat diet.” The testing of similar nutrition-focused data elements in the PAC PRD, and the current assessment of various nutritional services across the four PAC settings, demonstrates the feasibility of collecting information about this nutritional service in these settings.

Clinical and subject matter expert advisors working with our data element contractor agreed that the proposed Mechanically Altered Diet data element is feasible for use in PAC, and it assesses an important treatment that would be clinically useful both within and across PAC settings. Expert input on the Mechanically Altered Diet data element highlighted its importance and clinical usefulness for patients in PAC settings, due to the increased monitoring and resource use required for patients on special diets. We note that the Mechanically Altered Diet data element was also supported by the TEP that discussed candidate data elements for Special Services, Treatments, and Interventions during a meeting on January 5 and 6, 2017. A full report of the TEP discussion is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

Therefore, we are proposing that the Mechanically Altered Diet data element meets the definition of standardized patient assessment data with respect to special services, treatments, and interventions under section 1899B(b)(1)(B)(iii) of the Act. We are proposing to add the Mechanically Altered Diet data element to the LCDS and that LTCHs would be required to report these data for the FY 2020 LTCH QRP with respect to LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting.

We are inviting public comment on these proposals.

(4) Medical Condition and Comorbidity Data

We are proposing that the data elements needed to calculate the current measure, Percent of Residents or Patients with Pressure Ulcers That Are New or Worsened (Short Stay) (NQF #0678), and the proposed measure, Changes in Skin Integrity Post-Acute Care: Pressure Ulcer/Injury, meet the definition of standardized patient assessment data with respect to medical conditions and co-morbidities under section 1899B(b)(1)(B)(iv) of the Act, and that the successful reporting of that data under section 1886(m)(5)(F)(ii) of the Act would also satisfy the requirement to report standardized patient assessment data under section 1886(m)(5)(F)(ii) of the Act.

“Medical conditions and comorbidities” and the conditions addressed in the standardized data elements used in the calculation and risk adjustment of these measures, that is, the presence of pressure ulcers, diabetes, incontinence, peripheral vascular disease or peripheral arterial disease, mobility, as well as low body mass index, are all health-related conditions that indicate medical complexity that can be indicative of underlying disease severity and other comorbidities.

Specifically, the data elements used in the measure are important for care planning and provide information pertaining to medical complexity. Pressure ulcers are serious wounds representing poor outcomes, and can result in sepsis and death. Assessing skin condition, care planning for pressure ulcer prevention and healing, and informing providers about their presence in patient transitions of care is a customary and best practice. Venous and arterial disease and diabetes are associated with low blood flow which may increase the risk of tissue damage. These diseases are indicators of factors that may place individuals at risk for pressure ulcer development and are therefore important for care planning. Low BMI, which may be an indicator of underlying disease severity, may be associated with loss of fat and muscle, resulting in potential risk for pressure ulcers. Bowel incontinence, and the possible maceration to the skin associated, can lead to higher risk for
pressure ulcers. In addition, the bacteria associated with bowel incontinence can complicate current wounds and cause local infection. Mobility is an indicator of impairment or reduction in mobility and movement which is a major risk factor for the development of pressure ulcers. Taken separately and together, these data elements are important for care planning, transitions in services and identifying medical complexities. In sections IX.C.7.a and IX.C.10.a of the preamble of this proposed rule, we discuss our rationale for proposing that the data elements used in the measures meet the definition of standardized patient assessment data. In summary, we believe that the collection of such assessment data is important for multiple reasons, including clinical decision support, care planning, and quality improvement, and that the data elements assessing pressure ulcers and the data elements used to risk adjust showed good reliability. We solicited stakeholder feedback on the quality measure, and the data elements from which it is derived, by means of a public comment period and TEPs, as described in section IX.C.7.a of the preamble of this proposed rule.

We are inviting public comment on this proposal.

(5) Impairment Data

Hearing and vision impairments are conditions that, if unaddressed, affect activities of daily living, communication, physical functioning, rehabilitation outcomes, and overall quality of life. Sensory limitations can lead to confusion in new settings, increase isolation, contribute to mood disorders, and impede accurate assessment of other medical conditions. Failure to appropriately assess, accommodate, and treat these conditions increases the likelihood that patients will require more intensive and prolonged treatment. Onset of these conditions can be gradual, so individualized assessment with accurate screening tools and follow-up evaluations are essential to determining which patients need hearing- or vision-specific medical attention or assistive devices and accommodations, including auxiliary aids and/or services, and to ensure that person-directed care plans are developed to accommodate a patient’s needs. Accurate diagnosis and management of hearing or vision impairment would likely improve rehabilitation outcomes and care transitions, including transition from institutional-based care to the community. Accurate assessment of hearing and vision impairment would be expected to lead to appropriate treatment, accommodations, including the provision of auxiliary aids and services during the stay, and ensure that patients continue to have their vision and hearing needs met when they leave the facility.

Accurate individualized assessment, treatment, and accommodation of hearing and vision impairments of patients and residents in PAC would be expected to have a positive impact on the National Quality Strategy’s domains of patient and family engagement, patient safety, care coordination, clinical process/effectiveness, and efficient use of health care resources. For example, standardized assessment of hearing and vision impairments used in PAC will support ensuring patient safety (for example, risk of falls), identifying accommodations needed during the stay, and appropriate support needs at the time of discharge or transfer. Standardized assessment of these data elements will enable or support clinical decision-making and early clinical intervention; person-centered, high quality care (for example, facilitating better care continuity and coordination); better data exchange and interoperability between settings; and longitudinal outcome analysis. Hence, reliable data elements assessing hearing and vision impairments are needed to initiate a management program that can optimize a patient or resident’s prognosis and reduce the possibility of adverse events.

• Hearing

We are proposing that the Hearing data element meets the definition of standardized patient assessment data with respect to impairments under section 1899B(b)(1)B(v) of the Act. The proposed data element consists of the single Hearing data element. This data element assesses level of hearing impairment, and consists of one question. For more information on the Hearing data element, we refer readers to the document titled, Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

Accurate assessment of hearing impairment is important in the PAC setting for care planning and resource use. Hearing impairment has been associated with lower quality of life, including poorer physical, mental, and social functioning, and emotional health.426–427 Treatment and accommodation of hearing impairment led to improved health outcomes, including but not limited to quality of life.428 For example, hearing loss in elderly individuals has been associated with depression and cognitive impairment.429–430 Higher rates of incident cognitive impairment and cognitive decline,431 and less time in occupational therapy.432 Accurate assessment of hearing impairment is important in the PAC setting for care planning and defining resource use. The proposed data element was selected from two forms of the Hearing data element based on expert and stakeholder feedback. We considered the two forms of the Hearing data element, one of which is currently in use in the MDS 3.0 (Hearing) and another data element with different wording and fewer response option categories that is currently in use in the OASIS-C2 (Ability to Hear). Ability to Hear was also tested in the PAC PRD and found to have substantial agreement for inter-rater reliability across PAC settings (kappa of 0.78).433

Several data elements that assess hearing impairment were presented to the Standardized Patient Assessment Data TEP held by our data element contractor. The TEP did not reach consensus on the ideal number of

The Hearing data element would be assessed at admission only due to the relatively stable nature of hearing impairment, making it unlikely that this assessment would change between the start and end of the PAC stay. Assessment at discharge would introduce additional burden without improving the quality or usefulness of the data, and we believe it is unnecessary.

We are inviting public comment on these proposals.

- **Vision**

We are proposing that the Vision data element meets the definition of standardized patient assessment data element with respect to impairments under section 1899b(b)(1)(B)(v) of the Act. The proposed data element consists of the single Vision (Ability To See in Adequate Light) data element that consists of one item with five response categories. For more information on the Vision data element, we refer readers to the document titled, **Proposed Specifications for LTCH QRP Quality Measures and Standardized Data Elements**, available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

Evaluation of an individual’s ability to see is important for assessing for risks such as falls and provides opportunities for improvement through treatment and the provision of accommodations, including auxiliary aids and services, which can safeguard patients and improve their overall quality of life. Further, vision impairment is often a treatable risk factor associated with adverse events and poor quality of life. For example, individuals with visual impairment are more likely to experience falls and hip fracture, have less mobility, and report depressive symptoms.

Individualized initial screening can lead to life-improving interventions such as accommodations, including the provision of auxiliary aids and services, during the stay and/or treatments that can improve vision and prevent or slow further vision loss. For patients with some types of visual impairment, use of glasses and contact lenses can be effective in restoring vision. Other conditions, including glaucoma and age-related macular degeneration, have responded well to treatment. In addition, vision impairment is often a treachable risk factor associated with adverse events which can be prevented and accommodated during the stay. Accurate assessment of vision impairment is important in the LTCH setting for care planning and defining resource use.

The Vision data element that we are proposing for standardization was tested as part of the development of the MDS 3.0 and is currently in use in that assessment. Similar data elements, but with different wording and fewer response option categories, are in use in the OASIS–C2 and was tested in post-acute care providers in the PAC PRD and found to be clinically relevant, meaningful for care planning, reliable (kappa of 0.74) and feasible for use in each of the four PAC settings.

Several data elements that assess vision were presented to the TEP held by our data element contractor. The TEP


did not reach consensus on the ideal number of response categories or phrasing of response options, which are the primary differences between the current MDS and OASIS items; some members preferring more granular response options (for example, mild impairment and moderate impairment) while others were comfortable with collapsed response options (that is, mild/moderate impairment). The Development and Maintenance of Post-Acute Care Cross-Setting Standardized Patient Assessment Data Technical Expert Panel Summary Report is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html. We solicited public comment from August 12 to September 12, 2016, on the Ability to See in Adequate Light data element (version tested in the PAC PRD with three response categories). The data element in public comment differed from the proposed data element, but the comments supported the assessment of vision in PAC settings and the useful information a vision data element would provide. The commenters stated that the Ability to See item would provide important information that would facilitate care coordination and care planning, and consequently improve the quality of care. Other commenters suggested it would be helpful as an indicator of resource use and noted that the item would provide useful information about the abilities of patients and residents to care for themselves. Additional commenters noted that the item could feasibly be implemented across PAC providers and that its kappa scores from the PAC PRD support its validity. Some commenters noted a preference for MDS version of the Vision data element over the form put forward in public comment, citing the widespread use of this data element. A full report of the comments is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/Post-Acute-Care-Quality-Initiatives/IMPACT-Act-of-2014/IMPACT-Act-Downloads-and-Videos.html.

Clinical and subject matter expert advisors working with our data element contractor agreed that assessing vision impairment of patients and residents with a standardized data element is feasible in PAC, that it can reliably and accurately identify adults with objective impaired vision, and that this information about impaired vision would be clinically useful to identify needed accommodations and/or treatment both within and across PAC settings.

Therefore, we are proposing the Vision data element from the MDS. We are proposing to add the Vision data element to the LCDS and that LTCHs would be required to report these data for the FY 2020 LTCH QRP with respect to LTCH admissions that occur between April 1, 2018 and December 31, 2018. Following the initial reporting year for the FY 2020 LTCH QRP, subsequent years for the LTCH QRP would be based on a full calendar year of such data reporting. The Vision data element would be assessed at admission only due to the relatively stable nature of vision impairment, making it unlikely that this assessment would change between the start and end of the PAC stay. Assessment at discharge would introduce additional burden without improving the quality or usefulness of the data, and we believe that it is unnecessary.

We are inviting public comment on these proposals.

11. Proposals Relating to the Form, Manner, and Timing of Data Submission Under the LTCH QRP
a. Proposed Start Date for Standardized Patient Assessment Data Reporting by New LTCHs

In the FY 2016 IPPS/LTCH PPS final rule (80 FR 49749 through 49752), we adopted timing for new LTCHs to begin reporting quality data under the LTCH QRP beginning with the FY 2017 LTCH QRP. We are proposing that new LTCHs will be required to begin reporting standardized patient assessment data on the same schedule.

We are inviting public comment on this proposal.

b. Proposed Mechanism for Reporting Standardized Patient Assessment Data Beginning With the FY 2019 LTCH QRP

Under our current policy, LTCHs report data by completing applicable sections of the LCDS, and submitting the LCDS to CMS through the QIES ASP system. For more information on LTCH QRP reporting through the QIES ASP system, refer to the “Related Links” section at the bottom of: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Technical-Information.html.

The proposed standardized patient assessment data elements are already included on, or would be added to, the LCDS. Details regarding the LCDS with respect to the proposed standardized assessment data are available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html.

We are inviting public comments on this proposal.

c. Proposed Schedule for Reporting Standardized Patient Assessment Data Beginning With the FY 2019 LTCH QRP

We are proposing that the standardized patient assessment data as discussed in section IX.C.10.a. of the preamble of this proposed rule necessary to calculate the quality measure “Percent of Residents or Patients with Pressure Ulcers That Are New or Worsened (Short Stay) (NQF #0678)” would be used for the FY 2019 LTCH QRP. We are also proposing that for purposes of the FY 2019 LTCH QRP program year such data would only include the last three quarters of calendar year 2017 (April 1, 2017 through December 31, 2017). In section IX.C.7.a. of the preamble of this proposed rule, we proposed to adopt the measure, “Changes in Skin Integrity Post-Acute Care: Pressure Ulcer/Injury”’ to replace the current measure, “Percent of Residents or Patients with Pressure Ulcers That Are New or Worsened (Short Stay) (NQF # 0678)” with data collection beginning on April 1, 2018. Should the proposed measure be finalized, the FY 2020 LTCH QRP will be determined using the data from the first quarter of CY 2018 using the current measure, “Percent of Residents or Patients with Pressure Ulcers That Are New or Worsened (Short Stay) (NQF #0678)” and last three quarters of CY 2018 using the data from the proposed measure, “Changes in Skin Integrity Post-Acute Care: Pressure Ulcer/Injury.” In section IX.C.10.b of the preamble of this proposed rule, we discussed the additional standardized patient data proposed beginning with the FY 2020 LTCH QRP. Unless otherwise indicated, under our current policy, except for the first program year for which a measure is adopted, LTCHs must report data on measures with respect to LTCH admissions and discharges that occur during the 12 month calendar year period that applies to the program year. For the first program year for which a measure is adopted, LTCHs are only required to report data for LTCH admissions and discharges that occur during the last three quarters of the calendar year that applies to that program year, as the version of the LTCH CARE Data Set that will contain the new items that allow LTCHs to report a new measure, is routinely released on April 1st of any given year.
For example, for the FY 2018 LTCH QRP, data on measures adopted for earlier program years must be reported with respect to all CY 2016 LTCH admissions and discharges. However, data on new measures adopted for the first time for the FY 2018 LTCH QRP must only be reported with respect to LTCH admissions and discharges that occur during the last three calendar quarters of 2016. This is because the newest iteration of the LTCH CARE Data Set would have been scheduled for release on April 1, 2016. We are proposing to apply this policy to the reporting of standardized patient assessment data beginning with the FY 2020 LTCH QRP. The tables below illustrate this policy using the FY 2020 and FY 2021 LTCH QRP as examples.

**SUMMARY ILLUSTRATION OF INITIAL REPORTING CYCLE FOR NEWLY ADOPTED MEASURE AND STANDARDIZED PATIENT ASSESSMENT DATA REPORTING USING CY QUARTERS 2, 3, AND 4 DATA**

<table>
<thead>
<tr>
<th>Proposed data collection/submission quarterly reporting period</th>
<th>Proposed data submission quarterly deadlines beginning with the FY 2020 LTCH QRP</th>
</tr>
</thead>
</table>

* Applies to data reporting using the LTCH CARE Data Set and data reporting using the National Healthcare Safety Network.

^ The term "FY 2020 LTCH QRP" means the fiscal year for which the LTCH QRP requirements applicable to that fiscal year must be met in order for an LTCH to receive the full annual update when calculating the payment rates applicable to it for that fiscal year.

**SUMMARY ILLUSTRATION OF CALENDAR YEAR QUARTERLY REPORTING CYCLE FOR MEASURE AND STANDARDIZED PATIENT ASSESSMENT DATA REPORTING**

<table>
<thead>
<tr>
<th>Proposed data collection/submission quarterly reporting period</th>
<th>Proposed data submission quarterly deadlines beginning with the FY 2021 LTCH QRP</th>
</tr>
</thead>
</table>

* Applies to data reporting using the LTCH CARE Data Set and data reporting using the National Healthcare Safety Network.

^ The term "FY 2021 LTCH QRP" means the fiscal year for which the LTCH QRP requirements applicable to that fiscal year must be met in order for an LTCH to receive the full annual update when calculating the payment rates applicable to it for that fiscal year.

We are inviting public comment on our proposal for standardized data reporting beginning with the FY 2019 LTCH QRP and to extend our current policy governing the schedule for reporting quality measure data to the reporting of standardized patient assessment data beginning with the FY 2020 LTCH QRP.

d. Proposed Schedule for Reporting the Proposed Quality Measures Beginning With the FY 2020 LTCH QRP

As discussed in section IX.C.7. of the preamble of this proposed rule, we are proposing to adopt three quality measures beginning with the FY 2020 LTCH QRP: Changes in Skin Integrity Post-Acute Care: Pressure Ulcer/Injury, Compliance with SBT by Day 2 of the LTCH Stay, and Ventilator Liberation Rate. We are proposing that LTCHs would report data on these measures using the LTCH CARE Data Set that is submitted through the QIES ASAP system. LTCHs would be required to report these data beginning with LTCH admissions and discharges that occur between April 1, 2018 and December 31, 2018. More information on LTCH reporting using the QIES ASAP system is located at: [https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Assessment-Information.html.](https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Assessment-Information.html)

Under our currently policy, LTCHs would only be required to submit data on the proposed measures for the last three quarters of CY 2018 for purposes of the FY 2020 LTCH QRP. Starting in CY 2019, LTCHs would be required to submit data for the entire calendar year beginning with the FY 2021 LTCH QRP.

We are inviting public comment on this proposal.

e. Proposed Removal of Interrupted Stay Items From the LTCH CARE Data Set

We are proposing to remove the program interruption items from the LTCH CARE Data Set. Specifically, we are proposing to remove the following items: (1) A2500, Program Interruption(s); (2) A2510, Number of Program Interruptions During This Stay in This Facility; and (3) A2525, Program Interruption Dates, because we do not currently utilize this information nor do we have plans to utilize this information for the LTCH QRP. For a detailed discussion of burden related to LTCH CARE Data Set, we refer readers to section XIV.B.9. of the preamble of this proposed rule.

We are inviting public comment on this proposal.

12. Proposed Changes to Previously Codified Participation Requirements Under the LTCH QRP

We are proposing to revise the regulatory text at §412.560(a) to state that an LTCH must begin submitting quality data, including standardized patient assessment data, under the LTCH QRP by no later than the first day of the calendar quarter subsequent to 30 days after the date on its CMS Certification Number (CCN) notification letter.

We are inviting public comments on this proposal.

13. Proposed Changes to Previously Codified Data Submission Requirements Under the LTCH QRP

We are proposing to revise the regulatory text at §412.560(b)(1) to require LTCHs to report both data on measures and standardized patient assessment data under the LTCH QRP in a form and manner, and at a time, specified by CMS.

We are inviting public comments on this proposal.

14. Proposed Changes to Previously Codified Exception and Extension Requirements Under the LTCH QRP

We are proposing to revise the regulatory text at §412.560(c) to extend
these policies to the submission of standardized patient assessment data beginning with the FY 2019 LTCH QRP. We are inviting public comments on this proposal.

15. Proposed Changes to Previously Codified Reconsiderations Requirements Under the LTCH QRP

We are proposing to revise the regulatory text at § 412.560(d) to extend these policies to the submission of standardized patient assessment data beginning with the FY 2019 LTCH QRP. We are inviting public comments on this proposal.

16. Proposal To Apply the LTCH QRP Data Completion Thresholds to the Submission of Standardized Patient Assessment Data Beginning With the FY 2019 LTCH QRP

In the FY 2015 IPPS/LTCH PPS final rule (79 FR 50311 through 50314), we finalized LTCH QRP thresholds for completeness of LTCH data submissions. To ensure that LTCHs are meeting an acceptable standard for completeness of submitted data, we finalized the policy that, beginning with the FY 2016 LTCH QRP, LTCHs must meet or exceed two separate data completeness thresholds: One threshold set at 80 percent for completion of measures data collected using the LTCH CARE Data Set and a second threshold set at 100 percent for data collected and submitted using the CDC NHSN. These thresholds would apply to all measures and data elements adopted into LTCH QRP. An LTCH must meet or exceed both thresholds to avoid receiving a 2 percentage point reduction to their annual payment update for a given fiscal year, beginning with the FY 2016 LTCH QRP for measures data and beginning with the FY 2019 LTCH QRP for standardized patient assessment data elements.

We are inviting public comment on our proposal to extend our current LTCH QRP data completion requirements to the reporting of standardized patient assessment data. We are also inviting public comment on our proposal to codify the LTCH QRP data completion thresholds at § 412.560(f) for measures and standardized patient assessment data elements collected using the LTCH CARE Data Set.

17. Proposals and Policies Regarding Public Display of Measure Data for the LTCH QRP

Section 1886(m)(5)(E) of the Act requires the Secretary to establish procedures for making the LTCH QRP data available to the public after ensuring that an LTCH has the opportunity to review its data prior to public display. Measure data is currently displayed on the Long-Term Care Hospital Compare Web site, which is an interactive web tool that assists individuals by providing information on LTCH quality of care including those who need to select an LTCH. For more information on LTCH Compare, we refer readers to: https://www.medicare.gov/longtermcarehospitalcompare/. In addition, for a more detailed discussion about the provider’s confidential review process prior to public display of quality measures we refer readers to the FY 2017 IPPS/LTCH PPS final rule (81 FR 57231 through 57236).

We also finalized the process we use to publish a list of LTCHs that successfully meet the reporting requirements for the applicable LTCH QRP year on the LTCH QRP Web site in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57231). The list of compliant LTCHs is available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Data-Submission-Deadlines.html.

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57231 through 57236), we finalized the public display of measure data on the LTCH Compare Web site in CY 2017 for the following 4 quality measures pending the availability of data: (1) NHSN Facility-wide Inpatient Hospital-onset MRSA Bacteremia Outcome Measure (NQF #1716); (2) NHSN Facility-wide Inpatient Hospital-onset CDI Outcome Measure (NQF #1717); (3) Influenza Vaccination Coverage Among Healthcare Personnel (NQF #0431) and Percent of Residents or Patients Who Were Assessed and Appropriately Given the Seasonal Influenza Vaccine (NQF #0680).

The public display of NHSN Facility-wide Inpatient Hospital-onset MRSA Bacteremia Outcome Measure (NQF #1716) and NHSN Facility-wide Inpatient Hospital-onset CDI Outcome Measure (NQF #1717) will initially be based on data collected from January 1, 2015 through December 31, 2015 and will be displayed based on 4 rolling quarters. The Influenza Vaccination Coverage Among Healthcare Personnel (NQF #0431) and Percent of Residents or Patients Who Were Assessed and Appropriately Given the Seasonal Influenza Vaccine (NQF #0680) will be based on the influenza vaccination season from October 1, 2015 through March 31, 2016 and will be updated annually. We refer readers to the FY 2017 IPPS/LTCH PPS final rule (81 FR 57231 through 57233) for details on the calculations and display of these quality measures.

In this proposed rule, pending the availability of data, we are proposing to publicly report data in CY 2018 for the following 3 assessment-based measures: (1) Percent of LTCH Patients With an Admission and Discharge Functional Assessment and a Care Plan That Addresses Function (NQF #2631); (2) Application of Percent of LTCH Patients With an Admission and Discharge Functional Assessment and a Care Plan That Addresses Function (NQF #2631); and (3) Application of Percent of Residents Experiencing One or More Falls with Major Injury (NQF #0674). In addition, pending the availability of data, we are proposing to publicly report data in CY 2020 for the assessment-based measure Functional Outcome Measure: Change in Mobility Among Patients Requiring Ventilator Support (NQF #2632). Data collection for these 4 new assessment-based
measures began on April 1, 2016. We are proposing to display data for the assessment-based measures based on four rolling quarters of data and would initially use discharges from January 1, 2017 through December 31, 2017, with the exception of Functional Outcome Measure: Change in Mobility Among Patients Requiring Ventilator Support (NQF #2632) which would be based on eight rolling quarters of data and would initially use discharges from January 1, 2017 through December 31, 2017.

In addition, we are proposing to publicly report 3 claims-based measures: (1) Medicare Spending Per Beneficiary-PAC LTCH QRP; (2) Discharge to Community-PAC LTCH QRP; and (3) Potentially Preventable 30-Day Post-Discharge Readmission Measure for LTCH QRP.

These measures were adopted for the LTCH QRP in the FY 2017 rule to be based on data from 2 consecutive calendar years. As previously adopted in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57233 through 57236), confidential feedback reports for these 3 claims-based measures will be based on calendar years 2015 and 2016 and data collected for discharges beginning January 1, 2015 through December 31, 2016. However, our current proposal revises the dates for public reporting and we are proposing to transition from calendar year to fiscal year to make these measure data publicly available by October 2018. Thus, we are proposing public reporting beginning in CY 2018 for these claims-based measures based on fiscal years 2016 and 2017 and data collected from discharges beginning October 1, 2015 through September 30, 2017.

We are proposing to remove the following claims-based measure “All-Cause Unplanned Readmission Measure for 30 Days Post Discharge from LTCHs’’ from the LTCH QRP and public reporting by October 2018. We refer readers to section IX.C.8. of the preamble of this proposed rule for additional information regarding the proposed removal of this measure from quality reporting and public display. We also are proposing to remove the following assessment-based measure “Percent of Residents or Patients with Pressure Ulcers That Are New or Worsened (Short Stay) (NQF #0678)” and to replace it with a modified version of the measure entitled “Changes in Skin Integrity Post-Acute Care: Pressure Ulcer/Injury” from the LTCH QRP and public reporting by October 2020. We refer readers to section IX.C.7.a. of the preamble of this proposed rule for additional information regarding the proposed replacement of this measure from quality reporting and public display.

For the assessment-based measures: Percent of LTCH Patients With an Admission and Discharge Functional Assessment and a Care Plan That Addresses Function (NQF #2631); Application of Percent of LTCH Patients With an Admission and Discharge Functional Assessment and a Care Plan That Addresses Function (NQF #2631); and Application of Percent of Residents Experiencing One or More Falls with Major Injury (NQF #0674), to ensure the statistical reliability of the measures, we are proposing to assign LTCHs with fewer than 25 eligible cases during a performance period to a separate category: “The number of cases/patient stays is too small to report.” If an LTCH had fewer than 25 eligible cases, the LTCH’s performance would not be publicly reported for the measure for that performance period.

For the claims-based measures: Discharge to Community-PAC LTCH QRP and Potentially Preventable 30-Day Post-Discharge Readmission Measure for LTCH QRP, to ensure the statistical reliability of the measure, we are proposing to assign LTCHs with fewer than 25 eligible cases during a performance period to a separate category: “The number of cases/patient stays is too small to report.” If an LTCH had fewer than 25 eligible cases, the LTCH’s performance would not be publicly reported for the measure for that performance period.

PREVIOUSLY FINALIZED AND PROPOSED MEASURES FOR CY 2018 PUBLIC DISPLAY AND CONFIDENTIAL FEEDBACK REPORTS

Previously Finalized Measures:

- Percent of Residents or Patients with Pressure Ulcers That Are New or Worsened (Short Stay) (NQF #0678).
- NHSN Facility-wide Inpatient Hospital-onset Methicillin-resistant Staphylococcus aureus (MRSA) Bacteremia Outcome Measure (NQF #1716).
- NHSN Facility-wide Inpatient Hospital-onset Clostridium difficile Infection (CDI) Outcome Measure (NQF #1717).
- Influenza Vaccination Coverage Among Healthcare Personnel (NQF #0431).
- Percent of Residents or Patients Who Were Assessed and Appropriately Given the Seasonal Influenza Vaccine (NQF #0680).

Proposed Measures:

- Percent of Long-Term Care Hospital (LTCH) Patients With an Admission and Discharge Functional Assessment and a Care Plan That Addresses Function (NQF #2631).
- Application of Percent of Long-Term Care Hospital (LTCH) Patients With an Admission and Discharge Functional Assessment and a Care Plan That Addresses Function (NQF #2631).
- Application of Percent of Residents Experiencing One or More Falls with Major Injury (NQF #0674).
- Medicare Spending Per Beneficiary-PAC LTCH QRP.
- Discharge to Community-PAC LTCH QRP.
- Potentially Preventable 30-Day Post-Discharge Readmission Measure for LTCH QRP.

PROPOSED ADDITIONAL MEASURE FOR CY 2020 PUBLIC DISPLAY AND CONFIDENTIAL FEEDBACK REPORTS

Functional Outcome Measure: Change in Mobility Among Long-Term Care Hospital (LTCH) Patients Requiring Ventilator Support (NQF #2632).
We are inviting public comment on the proposal for the public display of the four assessment-based measures and three claims-based measures, the removal of the All-Cause Unplanned Readmission Measure for 30 Days Post Discharge from LTCHs from the LTCH QRP and public display, and the replacement of “Percent of Residents or Patients with Pressure Ulcers That Are New or Worsened (Short Stay) (NQF #0678)” with a modified version of the measure entitled “Changes in Skin Integrity Post-Acute Care; Pressure Ulcer/Injury” as described above.

18. Mechanism for Providing Feedback Reports to LTCHs

Section 1899B(f) of the Act requires the Secretary to provide confidential feedback reports to PAC providers on their performance on the measures specified under sections 1899B(c)(1) and (d)(1) of the Act, beginning one year after the specified application date that applies to such measures and PAC providers. In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57233 through 57236), we finalized processes to provide LTCHs the opportunity to review their data and information using confidential feedback reports that will enable LTCHs to review their performance on the measures required under the LTCH QRP. Information on how to obtain these and other reports available to the LTCH can be found at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting.html.

We are not proposing any changes to this policy.

D. Inpatient Psychiatric Facility Quality Reporting (IPFQR) Program

1. Background

a. Statutory Authority

Section 1886(s)(4) of the Act, as added and amended by sections 3401(f) and 10322(a) of the Patient Protection and Affordable Care Act, requires the Secretary to implement a quality reporting program for inpatient psychiatric hospitals and psychiatric units. Section 1886(s)(4)(A)(i) of the Act requires that, for fiscal year (FY) 2014 and each subsequent fiscal year, the Secretary must reduce any annual update to a standard federal rate for discharges occurring during the fiscal year by 2.0 percentage points for any inpatient psychiatric hospital or psychiatric unit that does not comply with quality data submission requirements with respect to an applicable fiscal year.

As provided in section 1886(s)(4)(A)(ii) of the Act, the application of the reduction for failure to report under section 1886(s)(4)(A)(i) of the Act may result in an annual update of less than 0.0 percent for a fiscal year, and may result in payment rates under section 1886(s)(1) of the Act being less than the payment rates for the preceding year. In addition, section 1886(s)(4)(B) of the Act requires that the application of the reduction to a standard Federal rate update be noncumulative across fiscal years. Thus, any reduction applied under section 1886(s)(4)(A) of the Act will apply only with respect to the fiscal year rate involved and the Secretary may not take into account the reduction in computing the payment amount under the system described in section 1886(s)(1) of the Act for subsequent years.

Section 1886(s)(4)(C) of the Act requires that, for FY 2014 (October 1, 2013, through September 30, 2014) and each subsequent year, each psychiatric hospital and psychiatric unit must submit to the Secretary data on quality measures as specified by the Secretary. The data must be submitted in a form and manner at a time specified by the Secretary. Under section 1886(s)(4)(D)(i) of the Act, unless the exception of subsection (ii) applies, measures selected for the quality reporting program must have been endorsed by the entity with a contract under section 1890(a) of the Act. The National Quality Forum (NQF) currently holds this contract.

Section 1886(s)(4)(D)(ii) of the Act provides an exception to the requirement for NQF endorsement of measures: In the case of a specified area or medical topic determined appropriate by the Secretary for which a feasible and practical measure has not been endorsed by the entity with a contract under the annual payment rate update cycle, we revised the regulations at 42 CFR 412.402 to specify that, beginning October 1, 2012, the FY update period would be the 12-month period from October 1 through September 30, which we refer to as a “fiscal year” (FY) (76 FR 26435). Therefore, with respect to the IPFQR Program, the terms “rate year,” as used in the statute, and “fiscal year,” as used in the regulation, both refer to the period from October 1 through September 30. For more information regarding this terminology change, we refer readers to section III. of the FY 2012 IPF PPS final rule (76 FR 26434 through 26435).

The statute uses the term “rate year” (RY). However, beginning with the annual update of the inpatient psychiatric facility prospective payment system (IPF PPS) that took effect on July 1, 2011 (RY 2012), we aligned the IPF PPS update with the annual update of the ICD codes, effective on October 1 of each year. This change allowed for annual payment updates and the ICD coding update to occur on the same schedule and appear in the same Federal Register document, promoting administrative efficiency. To reflect the change to section 1890(a) of the Act, the Secretary may specify a measure that is not so endorsed as long as due consideration is given to measures that have been endorsed or adopted by a consensus organization identified by the Secretary.

Section 1886(s)(4)(E) of the Act requires the Secretary to establish procedures for making public the data submitted by inpatient psychiatric hospitals and psychiatric units under the IPFQR Program. These procedures must ensure that a facility has the opportunity to review its data prior to the data being made public. The Secretary must report quality measures that relate to services furnished by the psychiatric hospitals and units on the CMS Web site.

b. Covered Entities

In the FY 2013 IPPS/LTCH PPS final rule (77 FR 53645), we established that the IPFQR Program’s quality reporting requirements cover those psychiatric hospitals and psychiatric units paid under Medicare’s Inpatient Psychiatric Facility Prospective Payment System (IPF PPS) (42 CFR 412.404(b)). Generally, psychiatric hospitals and psychiatric units within acute care and critical access hospitals that treat Medicare patients are paid under the IPF PPS. Consistent with prior rules, we continue to use the term “inpatient psychiatric facility” (IPF) to refer to both inpatient psychiatric hospitals and psychiatric units. This usage follows the terminology in our IPF PPS regulations at 42 CFR 412.402. For more information on covered entities, we refer readers to the FY 2013 IPPS/LTCH PPS final rule (77 FR 53645).

c. Considerations in Selecting Quality Measures

Our objective in selecting quality measures is to balance the need for information on the full spectrum of care delivery and the need to minimize the burden of data collection and reporting. We have primarily focused on measures that evaluate critical processes of care that have significant impact on patient outcomes and support CMS and HHS priorities for improved quality and efficiency of care provided by IPPs. When possible, we also seek to incorporate measures that directly evaluate patient outcomes. We refer readers to section VIII.F.4.a. of the FY 2013 IPPS/LTCH PPS final rule (77 FR 53645 through 53646) for a detailed discussion of the considerations taken into account in selecting quality measures.
(1) Measure Selection Process

Before being proposed for inclusion in the IPFQR Program, measures are placed on a list of measures under consideration, which is published annually by December 1, on behalf of CMS by the NQF. In compliance with section 1890A(o)(2) of the Act, measures proposed for the IPFQR Program were included in a publicly available document: “List of Measures under Consideration for December 1, 2016” available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/QualityMeasures/Downloads/Measures-under-Consideration-List-for-2016.pdf.

The Measure Applications Partnership (MAP), a multi-stakeholder group convened by the NQF, reviews the measures under consideration for the IPFQR Program, among other Federal programs, and provides input on those measures to the Secretary. The MAP’s 2017 recommendations for quality measures under consideration are captured in the following documents: “Process and Approach for MAP Pre-Rulemaking Deliberations, 2016–2017,” available at: http://www.qualityforum.org/WorkArea/linkit.aspx?LinkIdentifier=id&ItemID=84455 and “2016–2017 Spreadsheet of Final Recommendations to HHS and CMS” available at: http://www.qualityforum.org/WorkArea/linkit.aspx?LinkIdentifier=id&ItemID=84452. We considered the input and recommendations provided by the MAP in selecting all measures for the IPFQR Program, including those discussed below.

(2) Accounting for Social Risk Factors in the IPFQR Program

We understand that social risk factors such as income, education, race and ethnicity, employment, disability, community resources, and social support (certain factors of which are also sometimes referred to as socioeconomic status (SES) factors or socio-demographic status (SDS) factors) play a major role in health. One of our core objectives is to improve beneficiary outcomes including reducing health disparities, and we want to ensure that all beneficiaries, including those with social risk factors, receive high quality care. In addition, we seek to ensure that the quality of care furnished by providers and suppliers is assessed as fairly as possible under our programs while ensuring that beneficiaries have adequate access to excellent care.

We have been reviewing reports prepared by the Office of the Assistant Secretary for Planning and Evaluation (ASPE) and the National Academies of Sciences, Engineering, and Medicine on the issue of measuring and accounting for social risk factors in CMS’ value-based purchasing and quality reporting programs, and considering options on how to address the issue in these programs. On December 21, 2016, ASPE submitted a Report to Congress on a study it was required to conduct under section 2(d) of the Improving Medicare Post-Acute Care Transformation (IMPACT) Act of 2014. The study analyzed the effects of certain social risk factors in Medicare beneficiaries on quality measures and measures of resource use used in one or more of nine Medicare value-based purchasing programs. The report also included considerations for strategies to account for social risk factors in these programs. In a January 10, 2017 report released by the National Academies of Sciences, Engineering, and Medicine, that body provided various potential methods for measuring and accounting for social risk factors, including stratified public reporting.

As noted in the FY 2017 IPPS/LTCH PPS final rule, the NQF has undertaken a 2-year trial period in which new measures, measures undergoing maintenance review, and measures endorsed with the condition that they enter the trial period can be assessed to determine whether risk adjustment for selected social risk factors is appropriate for these measures. This trial entails temporarily allowing inclusion of social risk factors in the risk-adjustment approach for these measures. At the conclusion of the trial, NQF will issue recommendations on the future inclusion of social risk factors in risk adjustment for these quality measures, and we will closely review their findings.

As we continue to consider the analyses and recommendations from these reports and await the results of the NQF trial on risk adjustment for quality measures, we are continuing to work with stakeholders in this process. As we have previously communicated, we are concerned about holding providers to different standards for the outcomes of their patients with social risk factors because we do not want to mask potential disparities or minimize incentives to improve the outcomes for disadvantaged populations. Keeping this concern in mind, while we sought input on this topic previously, we continue to seek public comment on whether we should account for social risk factors in the IPFQR Program, and if so, what method or combination of methods would be most appropriate for accounting for social risk factors.

Examples of methods include: Confidential reporting to providers of measure rates stratified by social risk factors; public reporting of stratified measure rates; and potential risk adjustment of a particular measure as appropriate based on data and evidence.

In addition, we are seeking public comment on which social risk factors might be most appropriate for reporting stratified measure scores and/or potential risk adjustment of a particular measure. Examples of social risk factors include, but are not limited to, dual eligibility/low-income subsidy, race and ethnicity, and geographic area of residence. We are seeking comments on which of these factors, including current data sources where this information would be available, could be used alone or in combination, and whether other data should be collected to better capture the effects of social risk. We will take commenters’ input into consideration as we continue to assess the appropriateness and feasibility of accounting for social risk factors in the IPFQR Program. We note that any such changes would be proposed through future notice-and-comment rulemaking.

We look forward to working with stakeholders as we consider the issue of accounting for social risk factors and reducing health disparities in CMS programs. Of note, implementing any of the above methods would be taken into consideration in the context of how this and other CMS programs operate (for example, data submission methods, availability of data, statistical considerations relating to reliability of data calculations, among others), so we also welcome comment on operational considerations. CMS is committed to ensuring that its beneficiaries have access to and receive excellent care, and that the quality of care furnished by providers and suppliers is assessed fairly in CMS programs.
(3) IPFQR Program Measures Adopted in Previous Payment Determinations

The current IPFQR Program includes 18 mandatory measures. For more information on these measures, we refer readers to the following final rules:

- The FY 2013 IPPS/LTCH PPS final rule (77 FR 53646 through 53652);
- The FY 2014 IPPS/LTCH PPS final rule (78 FR 50889 through 50895);
- The FY 2015 IPPS final rule (79 FR 45963 through 45974);
- The FY 2016 IPPS PPS final rule (80 FR 46694 through 46714); and
- The FY 2017 IPPS/LTCH PPS final rule (81 FR 57236 through 57249).

2. Factors for Removal or Retention of IPFQR Program Measures

a. Background

The Hospital IQR Program adopted formal policies regarding measure retention and removal in the FY 2011 IPPS/LTCH PPS final rule (75 FR 50185). We believe that it is important to be consistent between programs to the extent possible. Therefore, to align with the policies adopted in this and other quality reporting programs, we are proposing to adopt similar policies within the IPFQR Program. In the past, we have retained measures from each previous year’s IPFQR Program measure set for subsequent years’ measure sets, except when we specifically proposed to remove or replace a measure. For example, we removed HBIPS–6 and HBIPS–7 and replaced these measures with Transition Record with Specified Elements Received by Discharged Patients (NQF #0647) and Timely Transmission of Transition Record (NQF #0648) respectively in the FY 2016 IPPS PPS final rule (80 FR 46701 through 46709). In this proposed rule, we are proposing factors to consider in removing or retaining measures effective upon finalization of this proposed rule, anticipated to be effective October 1, 2017 and for subsequent years.

We will continue to use the notice and comment rulemaking process to propose measures for removal or replacement.

b. Proposed Considerations in Removing or Retaining Measures

With respect to measure removal, we believe it is important to be transparent in identifying factors that we would take into consideration on a case-by-case basis as guidelines to evaluate a measure for potential removal from the IPFQR Program. We believe that these factors should be aligned between our programs whenever possible. Therefore, we refer readers to the Hospital IQR Program (80 FR 49641 through 49642) factors we consider in removing or retaining measures. We intend to align our policies in the IPFQR Program with those in the Hospital IQR Program. Thus, in this proposed rule, we are proposing: (1) Measure removal factors; (2) criteria for determining when a measure is “topped-out;” and (3) measure retention factors. These proposals are discussed in more detail below.

We are proposing the following measure removal factors for the IPFQR Program:

- Measure performance among IPFs is so high and unvarying that meaningful distinctions and improvements in performance can no longer be made (“topped-out” measures);
- Measure does not align with current clinical guidelines or practice;
- Measure can be replaced by a more broadly applicable measure (across settings or populations) or a measure that is more proximal in time to desired patient outcomes for the particular topic;
- Measure performance or improvement does not result in better patient outcomes;
- Measure can be replaced by a measure that is more strongly associated with desired patient outcomes for the particular topic;
- Measure collection or public reporting leads to negative unintended consequences other than patient harm; and
- Measure is not feasible to implement as specified.

For the purposes of considering measures for removal from the program, we are also proposing to align our criteria for determining that a measure is “topped-out” with the Hospital IQR Program’s criteria (80 FR 49642), which states that a measure is “topped-out” if there is statistically indistinguishable performance at the 75th and 90th percentiles and the truncated coefficient of variation is less than or equal to 0.10. Furthermore, we recognize that there may be times when measures may meet some of the outlined factors for removal, but continue to bring value to the program. Therefore, we are also proposing the following factors for consideration in determining whether to retain a measure in the IPFQR Program, which also are based on factors established in the Hospital IQR Program (80 FR 49641 through 49642):

- Measure aligns with other CMS and HHS policy goals, such as those delineated in the National Quality Strategy or CMS Quality Strategy;
- Measure aligns with other CMS programs, including other quality reporting programs; and
- Measure supports efforts to move IPFs towards reporting electronic measures.

We reiterate that these removal and retention factors are considerations that we take into account in balancing the benefits and drawbacks of whether or not to remove measures on a case-by-case basis.

We are inviting public comment on our proposals to adopt: (1) Measure removal factors; (2) criteria for determining when a measure is “topped out;” and (3) measure retention factors as discussed above. If finalized, these factors and criteria will become effective upon finalization of this proposed rule, anticipated to be effective October 1, 2017 and for subsequent years; measures identified as appropriate for removal would be proposed through notice-and-comment rulemaking subsequent to that date.

3. Proposed New Quality Measure for the FY 2020 Payment Determination and Subsequent Years—Medication Continuation Following Inpatient Psychiatric Discharge

a. Background

We are proposing one new measure, Medication Continuation following Inpatient Psychiatric Discharge, for the FY 2020 payment determination and subsequent years. The measure uses Medicare fee-for-service (FFS) claims to identify whether patients admitted to IPFs with diagnoses of major depressive disorder (MDD), schizophrenia, or bipolar disorder had filled at least one evidence-based medication within 2 days prior to discharge through 30 days post-discharge. We believe that medication continuation is important for patients discharged from the inpatient psychiatric setting with MDD, schizophrenia, or bipolar disorder because of significant negative outcomes associated with non-adherence to medication regimens. For example, patients with MDD who do not remain on prescribed medications are more likely to have negative health outcomes such as relapse and readmission, decreased quality of life, and increased healthcare costs.451 452 Patients with schizophrenia who do not adhere to their medication regimen are more likely to be hospitalized, use emergency psychiatric services, be arrested, be victims of crimes, and consume alcohol.

Patients with bipolar disorder who do not adhere to their medications have increased suicide risk. For these reasons, guidelines from the American Psychiatric Association (APA) and the Department of Veterans Affairs/Department of Defense (VA/DoD), which are based on extensive literature, recommend pharmacotherapy as the primary form of treatment for patients with these conditions. Interventions that can be applied in the inpatient setting that increase medication compliance and prevent the negative outcomes associated with nonadherence have been identified. These interventions include patient education, enhanced therapeutic relationships, shared decision-making, and text-message reminders, with multidimensional approaches resulting in the best outcomes.

Furthermore, patients and caregivers interviewed during the development of this measure indicated the importance of the facility’s role in communicating information about medications to the patient, pharmacy, and outpatient providers.

b. Appropriateness for the IPFQR Program

In accordance with section 1890a(a)(2) of the Act, this measure was included in a publicly available document: “List of Measures under Consideration for December 1, 2016” available at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/QualityMeasures/Downloads/Measures-under-Consideration-List-for-2016.pdf. The MAP Hospital Workgroup concluded that the measure addressed a critical quality objective, was evidence-based, and would contribute to efficient use of resources. One Workgroup member commented that it was appropriate to hold IPF’s accountable for patients filling a prescription for an evidence-based medication post-discharge, further remarking that the measure was moving in the right direction.

The MAP Hospital Workgroup classified the measure as “Refine and Resubmit Prior to Rulemaking.” The measure received this classification because the MAP recommended that measure testing be completed to demonstrate reliability and validity at the facility level in the hospital setting and that the measure be submitted to NQF for review and endorsement.

The MAP also requested additional details on the measure, such as: (1) The definition of medication dispensation; (2) how does the facility know whether the medication was dispensed; and (3) how the measure would be impacted if Medicare Part D coverage is optional. The MAP also recommended that this measure be submitted to NQF for review and endorsement. The final methodology report includes the results of reliability and validity testing, and additional measure updates that occurred between the MAP review and NQF submission in December 2016. This methodology report also provides the additional details requested by MAP at the December meeting.

Reliability and validity testing completed in 2016 using the final measure specifications demonstrates that the measure, as specified, provides reliable and valid facility-level scores of medication continuation.

Reliability was established using a method of mean denominator and volume categories. Using that approach, a minimum denominator size of 75 discharges was established to attain an overall reliability score of at least 0.7; this reliability score is within acceptable norms and indicates sufficient signal strength to discriminate performance between facilities. This means that it is possible to distinguish good performance from poor performance based on measure scores among facilities with at least 75 cases in the denominator.

Validity was established by evaluating the correlations of medication continuation scores with the conceptually related IPFQR Program measures. The medication continuation scores were moderately correlated with the scores for 7- and 30-day follow-up after hospitalization for mental illness as expected (rho = 0.35 and 0.45, where rho is the Spearman’s rank correlation coefficient). In other words, the positive correlation between scores of these two types of measures is expected because high follow-up rates with mental health providers and high follow-up rates of medication continuation both indicate a high-quality transition from the inpatient to the outpatient setting. The medication continuation scores were negatively correlated with readmission scores as expected (rho = −0.27). This negative correlation is expected because patients that do not continue their medications are more likely to relapse and be
readmitted. All correlations are statistically significant at p-value <0.0001. After reviewing these results and the proposed measure specifications, all of the 10 TEP members who were present for the face validity vote agreed that the measure score had face validity.

This measure was submitted to NQF for endorsement on December 16, 2016, the NQF Standing Committee has recommended the measure for endorsement, and we are currently awaiting NQF’s final decision. Under section 1886(s)(4)(D)(ii) of the Act, measures selected for the IPFQR Program must have been endorsed by the entity with a contract under section 1890(a) of the Act. The NQF currently holds this contract. However, section 1886(s)(4)(D)(ii) of the Act provides that, in the case of a specified area or medical topic determined appropriate by the Secretary for which a feasible and practical measure has not been endorsed by the entity with a contract under section 1890(a) of the Act, the Secretary may specify that a measure that is not so endorsed as long as due consideration is given to measures that have been endorsed or adopted by a consensus organization identified by the Secretary. We have reviewed NQF endorsed measures related to medication continuation in this patient population and did not identify any equivalent measures. We believe this measure is consensus-based because of the extensive measure development process, including the solicitation of expert and patient opinion and public comments (discussed in more detail below).

In addition, the proposed measure addresses several aspects of the CMS Quality Strategy goals and objectives. The measure supports the CMS Quality Strategy Goal to "promote effective prevention and treatment of chronic disease," which includes an objective to improve behavioral health access and quality of care by using evidence-based practices. The measure also supports the CMS Quality Strategy Goal to "promote effective communication and coordination of care." Specifically, the measure addresses three objectives within the goal of "promoting effective communication and coordination of care": (1) "To reduce admissions and readmissions" as patients with these conditions who do not adhere to their medication regimens are at an increased risk of relapse and readmission; (2) "to embed best practices to enable successful transitions between all settings of care." Because ensuring medication continuation following discharge is a critical component of transitioning from the IPF to the home or home health care; and (3) "to enable effective healthcare system navigation," as we believe this measure will encourage IPFs to provide information to patients regarding the importance of medication continuation and guidance on how to fill prescriptions following discharge.

The proposed measure would complement the portfolio of facility-level measures in the IPFQR Program that assess the transition from the inpatient to outpatient setting: Follow-Up After Hospitalization for Mental Illness; Thirty-day All Cause Unplanned Readmission Following Psychiatric Hospitalization in an Inpatient Psychiatric Facility; Transition Record with Specified Elements Received by Discharged Patients; and Timely Transmission of Transition Record.

More detailed information about the development of this measure as well as final measure specifications can be downloaded from the CMS Web site at: https://www.cms.gov/medicare/Quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInits/Measure-Methodology.html. To access the report, click on the zip file titled “Inpatient Psychiatric Facility Medication Continuation Measure.”

c. Measure Calculation

The measure is calculated by dividing the number of admissions that meet the numerator criteria (described below) by the number of admissions that meet the denominator criteria (also described below).

(1) Numerator

The numerator for the measure includes discharges for patients with a principal diagnosis of MDD, schizophrenia, or bipolar disorder in the denominator who were dispensed at least one evidence-based outpatient medication within 2 days prior to discharge through 30 days post-discharge. The evidence-based medications that define the numerator are based on the practice guidelines for each condition from the APA and VA/DoD. Furthermore, we sought to align the medications with evidence-based medications from existing quality measures including the Antidepressant Medication Management measure from the Healthcare Effectiveness Data and Information Set (HEDIS) 2015 for MDD, the Adherence

to Antipsychotic Medications for Individuals with Schizophrenia measure (NQF #1879) for schizophrenia, and the Adherence to Mood Stabilizers for Individuals with Bipolar I Disorder measure (NQF #1880) for bipolar disorder. Staff pharmacists reviewed these lists of medications for completeness and appropriateness in the IPF setting. The finalized lists of evidence-based medications are available in the measure methodology report at: https://www.cms.gov/medicare/Quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInits/Measure-Methodology.html. To access the report, click on the zip file titled “Inpatient Psychiatric Facility Medication Continuation Measure.”

We considered the appropriate number of days prior to discharge and post-discharge to include in the follow-up period for the denominator. Clinical experts noted that discharge planning may start as early as 2 days prior to discharge and that some facilities may help patients fill their outpatient prescriptions prior to discharge. Therefore, the numerator includes outpatient medications filled up to 2 days prior to discharge (Day 2 through Day 1). The follow-up period extends 30 days post-discharge (Day 0 through Day 30) to align with other care coordination measures, such as the 30 day follow-up period in Follow-Up After Hospitalization for Mental Illness (FUH) (NQF #0576) when we finalized for the IPFQR Program in the FY 2014 IPPS/LTCH PPS final rule (76 FR 50894 through 50896). To further support a 30-day follow-up period, we confirmed that over 93 percent of the evidence-based prescriptions filled prior to the admission were for a 30-day supply, which indicates that most patients would need to fill a medication within 30 days of discharge to avoid gaps in treatment even if they had some medications at home.

(2) Denominator

The denominator for the measure includes Medicare FFS beneficiaries aged 18 years and older who were discharged from an IPF to home or home health care 490 with a principal diagnosis of MDD, schizophrenia, or bipolar disorder. The denominator excludes discharges for patients who:

- Received Electroconvulsive Therapy (ECT) during the inpatient stay or follow-up period because some patients who receive ECT during the inpatient stay or follow-up period may have failed pharmacotherapy and would not fill an evidence-based prescription post-discharge;
- Received Transcranial Magnetic Stimulation (TMS) during the inpatient stay or follow-up period because some patients who receive TMS during the inpatient stay or follow-up period may have failed pharmacotherapy and would not fill an evidence-based prescription post-discharge;
- Were pregnant during the inpatient stay because some of the evidence-based medications for the treatment of MDD, schizophrenia, and bipolar disorder are contraindicated during pregnancy;

All patients in the measure denominator are enrolled in Medicare Parts A, B, and D during the measurement and follow-up periods. Therefore, these patients have prescription drug coverage for evidence-based medications in the measure. While patients are responsible for some out-of-pocket medication costs after Part D has been applied, low income patients qualify for additional support through both Medicare and Medicaid to help mitigate the cost of prescriptions and ensure that patients do not face financial barriers to filling necessary medications.

We refer readers to the measure specifications for more details about measure inclusions and exclusions at: https://www.cms.gov/medicare/Quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInits/Measure-Methodology.html. To access the report, click on the zip file titled “Inpatient Psychiatric Facility Medication Continuation Measure.”

The proposed measure would be implemented using Medicare FFS Parts A, B, and D claims data to calculate the measure results. Valid prescription drug claim data from Medicare Parts B and D provide the data necessary to calculate this measure. Therefore, no data collection will be required from IPFs. The measure would be reported as a combined facility-level rate across all three conditions. The measurement period is 2 years to maximize the number of facilities with a minimum of 75 discharges, which is necessary for calculation of reliable facility-level scores.492 If this measure is finalized as proposed, we will inform stakeholders of the claims data collection period through a subregulatory process, such as on a CMS Web site and/or on our applicable listservers.

e. Public Comment

During the measure development process, we solicited public comments on the measure via the CMS Quality Measures Public Comment Page.493 We provided the draft measure information form,494 and draft measure justification form495 to the public for review. We accepted public comments from August 25, 2016 through September 15, 2016. Numerous commenters expressed support for the Medication Continuation following Inpatient Psychiatric Discharge measure (with only 6 of 53 commenters expressing reluctance to support the measure) and commented on the importance of measuring medication continuation as this is an important component of care transitions and reduces the risk of readmissions. We received public comments about denominator specifications, numerator specifications, data collection, and the attribution of the measure to the IPF, and the relevance of the proposed measure. After review and evaluation of all the public comments received, we expanded the follow-up period from day of discharge (Day 0) through 30 days post discharge to include outpatient

d. Data Sources

The proposed measure would be implemented using Medicare FFS Parts A, B, and D claims and enrollment data to calculate the measure results. Valid prescription drug claim data from Medicare Parts B and D provide the data necessary to calculate this measure. Therefore, no data collection will be required from IPFs. The measure would be reported as a combined facility-level rate across all three conditions. The measurement period is 2 years to maximize the number of facilities with a minimum of 75 discharges, which is necessary for calculation of reliable facility-level scores.492 If this measure is finalized as proposed, we will inform stakeholders of the claims data collection period through a subregulatory process, such as on a CMS Web site and/or on our applicable listservers.

490 The measure specifications, as submitted to the MAP, did not include home health care. For details of this addition, please see the measure methodology report: https://www.cms.gov/medicare/Quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInits/Measure-Methodology.html. To access the report, click on the zip file titled “Inpatient Psychiatric Facility Medication Continuation Measure.”


494 Ibid.

495 Ibid.
prescriptions filled up to 2 days prior to discharge as described above. For specific information regarding the comments we received, we refer readers to the public comment summary at: https://www.cms.gov/medicare/quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInits/Measure-Methodology.html. To access

the report, click on the zip file titled "Inpatient Psychiatric Facility Medication Continuation Measure.”

We believe the proposed measure evaluates a process with a demonstrated quality gap and has the potential to benefit patients. For these reasons and the reasons stated above, we are proposing the Medication Continuation following Inpatient Psychiatric Discharge measure described in this section for the FY 2020 payment determination and subsequent years.

In summary, we are proposing one measure for the FY 2020 payment determination and subsequent years, as shown in the table below.

NEWLY PROPOSED IPFQR PROGRAM MEASURE FOR THE FY 2020 PAYMENT DETERMINATION AND SUBSEQUENT YEARS

<table>
<thead>
<tr>
<th>National quality strategy priority</th>
<th>NQF #</th>
<th>Measure ID</th>
<th>Measure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Communication/Care Coordination ...</td>
<td>N/A</td>
<td>N/A</td>
<td>Medication Continuation following Inpatient Psychiatric Discharge.</td>
</tr>
</tbody>
</table>

We welcome public comment on our proposal to adopt the Medication Continuation following Inpatient Psychiatric Discharge measure.

4. Summary of Proposed and Previously Finalized Measures for the FY 2020 Payment Determinations and Subsequent Years

If the Medication Continuation following Inpatient Psychiatric Discharge measure is adopted, the number of measures for the FY 2020 payment determination and subsequent years will total 19 as set forth in the table below.

PROPOSED AND PREVIOUSLY FINALIZED MEASURES FOR THE FY 2020 PAYMENT DETERMINATION AND SUBSEQUENT YEARS

<table>
<thead>
<tr>
<th>NQF #</th>
<th>Measure ID</th>
<th>Measure</th>
</tr>
</thead>
<tbody>
<tr>
<td>640</td>
<td>HBIPS–2</td>
<td>Hours of Physical Restraint Use.</td>
</tr>
<tr>
<td>641</td>
<td>HBIPS–3</td>
<td>Hours of Seclusion Use.</td>
</tr>
<tr>
<td>560</td>
<td>HBIPS–5</td>
<td>Patients Discharged on Multiple Antipsychotic Medications with Appropriate Justification.</td>
</tr>
<tr>
<td>576</td>
<td>FUH</td>
<td>Follow-up After Hospitalization for Mental Illness.</td>
</tr>
<tr>
<td>1661</td>
<td>SUB–1</td>
<td>Alcohol Use Screening.</td>
</tr>
<tr>
<td>1663</td>
<td>SUB–2 and SUB–2a</td>
<td>Alcohol Use Brief Intervention Provided or Offered and SUB–2a Alcohol Use Brief Intervention.</td>
</tr>
<tr>
<td>1664</td>
<td>SUB–3 and SUB–3a</td>
<td>Alcohol and Other Drug Use Disorder Treatment Provided or Offered at Discharge and SUB–3a Alcohol and Other Drug Use Disorder Treatment at Discharge.</td>
</tr>
<tr>
<td>1651</td>
<td>TOB–1</td>
<td>Tobacco Use Treatment Provided or Offered and TOB–1 Tobacco Use Treatment.</td>
</tr>
<tr>
<td>1654</td>
<td>TOB–2 and TOB–2a</td>
<td>Tobacco Use Treatment Provided or Offered at Discharge and Tob–2a Tobacco Use Treatment.</td>
</tr>
<tr>
<td>1656</td>
<td>TOB–3 and TOB–3a</td>
<td>Tobacco Use Treatment Provided or Offered at Discharge and Tob–3a Tobacco Use Treatment at Discharge.</td>
</tr>
<tr>
<td>1659</td>
<td>IMM–2</td>
<td>Influenza Immunization.</td>
</tr>
<tr>
<td>647</td>
<td>N/A</td>
<td>Transition Record with Specified Elements Received by Discharged Patients (Discharges from an Inpatient Facility to Home/Self Care or Any Other Site of Care).</td>
</tr>
<tr>
<td>648</td>
<td>N/A</td>
<td>Timely Transmission of Transition Record (Discharges from an Inpatient Facility to Home/Self Care or Any Other Site of Care).</td>
</tr>
<tr>
<td>N/A</td>
<td>N/A</td>
<td>Screening for Metabolic Disorders.</td>
</tr>
<tr>
<td>431</td>
<td>N/A</td>
<td>Influenza Vaccination Coverage Among Healthcare Personnel.</td>
</tr>
<tr>
<td>N/A</td>
<td>N/A</td>
<td>Assessment of Patient Experience of Care.</td>
</tr>
<tr>
<td>N/A</td>
<td>N/A</td>
<td>Use of an Electronic Health Record.</td>
</tr>
<tr>
<td>2860*</td>
<td>N/A</td>
<td>Thirty-Day All-Cause Unplanned Readmission Following Psychiatric Hospitalization in an Inpatient Psychiatric Facility.</td>
</tr>
<tr>
<td>N/A</td>
<td>N/A</td>
<td>Medication Continuation following Inpatient Psychiatric Discharge.**</td>
</tr>
</tbody>
</table>

* Since this measure was finalized in the FY 2017 IPPS/LTCH PPS final rule (57239 through 57246), NQF endorsement has been received.

** New measure proposed for the FY 2020 payment determination and subsequent years.

5. Possible IPFQR Program Measures and Topics for Future Consideration

As we have previously indicated (79 FR 45974 through 45975), we seek to develop a comprehensive set of quality measures to be available for widespread use for informed decision-making and quality improvement in the IPF setting. Therefore, through future rulemaking, we intend to propose new measures for development or adoption that will help further our goals of achieving better healthcare and improved health for individuals who obtain inpatient psychiatric services through the widespread dissemination and use of quality information. As noted on the "List of Measures under Consideration for December 1, 2016”496 published by the NQF on behalf of CMS, we are considering a measure of Medication Reconciliation on Admission and a measure of Identification of Opioid Use Disorder among Patients Admitted to...

Inpatient Psychiatric Facilities. We welcome comments on these measure concepts for future inclusion in the IPFQR Program. In addition, we have identified several areas which we believe are important to stakeholders, but which are not currently sufficiently covered by IPFQR Program measures. These areas are:

- Family and caregiver engagement;
- Patient experience of care;
- Opioid use and treatment;
- Access to care; and
- Inpatient assaults and violence.

We welcome public comments on possible new measures in these or other areas.

6. Public Display and Review Requirements

We refer readers to the FY 2013 IPPS/LTCH PPS final rule (77 FR 53653 through 53654), in which we finalized that we would publicly display the submitted data on the CMS Web site beginning in the first quarter of the calendar year following the respective payment determination year. We also finalized that IPFs would have the opportunity to preview their data between September 20 and October 19 of the respective payment determination year. In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50899 through 50900), we finalized that the data be publicly displayed and review of data stating that we would publicly display the data in April of the calendar year following the start of the payment determination year and that the preview period would be 30 days approximately twelve weeks prior to the public display of the data. In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57248 through 57249), we finalized changes to how we specify the timeframes for the IPFQR Program, including that we would: (1) No longer specify the exact dates of the preview period or data publication in rulemaking; (2) make the data for the IPFQR Program available as soon as possible; (3) announce the exact timeframes through subregulatory guidance; and (4) continue our policy that the time period for review will be approximately 30 days. In this proposed rule, we are not proposing any changes to the public display and review policies.

7. Form, Manner, and Timing of Quality Data Submission for the FY 2019 Payment Determination and Subsequent Years

a. Procedural Requirements for FY 2019 Payment Determination and Subsequent Years

In the FY 2013 IPPS/LTCH PPS final rule (77 FR 53654 through 77 FR 53655), we finalized procedural requirements for the IPFQR Program, including the requirements that facilities must do the following to participate in the IPFQR Program:

- Register with QualityNet before the IPF begins reporting;
- Identify a QualityNet Administrator who follows the registration process listed on the QualityNet Web site;
- Complete a Notice of Participation (NOP) within a specified time period; and
- Submit aggregate numerator and denominator data for all age groups, for all measures.

In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50901), we clarified that the policy we adopted for the FY 2016 payment determination also applies to the FY 2017 payment determination and subsequent years, unless we change it through future rulemaking. In this proposed rule, we are proposing to make changes related to the Notice of Participation (NOP) and withdrawals for the FY 2019 payment determination and subsequent years.

In the FY 2013 IPPS/LTCH PPS final rule (77 FR 53654), we finalized our policies that IPFs participating in the IPFQR Program must comply with several procedural requirements. In that rule, one of the policies we finalized was that the time frame for completing an online NOP form is between January 1 and August 15 before each respective payment determination year (for example, for the FY 2017 payment determination year, IPFs would be required to submit an NOP between January 1, 2016 and August 15, 2016). Similarly, in the FY 2013 IPPS/LTCH PPS final rule (77 FR 53654), we also finalized that withdrawals from the IPFQR Program will be accepted no later than August 15 before the beginning of each respective payment determination year.

As described in section IX.D.7.b. of the preamble of this proposed rule, there have been times that we have updated the data submission period through subregulatory means; this has led to a data submission period that is not aligned with the submission period for the NOP or program withdrawal. To ensure these dates align, in this proposed rule, we are proposing to change the submission timeframes for both NOPs and withdrawals from between January 1 and August 15 before each respective payment determination year to prior to the end of the data submission period before each respective payment determination year. This means that we are proposing to accept NOPs and withdrawals any time prior to the end of the data submission period before the payment determination year. For example, for the FY 2019 payment determination year, if our proposal in IX.D.7.b. of the preamble of this proposed rule is finalized, the end of the data submission period would be a date on or after June 15, 2018 (which we would announce via subregulatory means). This date would coincide with the deadline to submit an NOP or withdraw from the program.

In addition, we are proposing to provide precise dates that define the end of the data submission period/NOP/withdrawal submission deadline through subregulatory means, such as on a CMS Web site and/or on our applicable listservs, beginning with the FY 2019 payment determination.

We are inviting public comment on our proposals to: (1) Change the submission timeframes for both NOPs and withdrawals to the end of the data submission period before each respective payment determination year; and (2) provide precise dates that define the end of the data submission period/NOP/withdrawal submission deadline through subregulatory means for the FY 2019 payment determination and subsequent years.

b. Data Submission Requirements for the FY 2019 Payment Determination and Subsequent Years

We refer readers to the FY 2013 IPPS/LTCH PPS final rule (77 FR 53655 through 53656) and the FY 2014 IPPS/LTCH PPS final rule (78 FR 50901) for our previously finalized policies regarding quality data submission requirements. In this proposed rule, we are proposing to make changes related to the data submission period for the FY 2019 payment determination and subsequent years.

In the FY 2013 IPPS/LTCH PPS final rule (77 FR 53655) we finalized our policies related to reporting periods and submission timelines for data required by the IPFQR Program. IPFs are required to submit their aggregated data on the measures on an annual basis, beginning in FY 2014 (77 FR 53655). In that rule, we specified that data must be submitted between July 1 and August 15 of the calendar year preceding a given payment determination year (for example, between July 1, 2015 and August 15, 2015 for the FY 2016 payment determination (77 FR 53655 through 53656)). In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50899), we clarified that this policy applies to all future years of data submission for the IPFQR Program unless we change the policy through future rulemaking.

Because there have been times that the submission period has been updated...
through the subregulatory process (for example, due to systems issues impacting data collection in the specified timeframe), in order to avoid contradictory guidance between dates established in the Federal Register and dates established through subregulatory guidance, we are proposing to no longer specify the exact dates of the submission period through rulemaking. We are proposing to provide these exact dates through a subregulatory process instead, beginning with the FY 2019 payment determination. We are proposing to shift to a 45-day submission period beginning at least 30 calendar days following the end of the data collection period. For example, for the FY 2019 payment determination, the latest reporting period for a measure for which facilities must submit data ends on March 31, 2018. In this example, the submission period would begin at least 30 days after March 31, 2018 (that is, no earlier than May 1, 2018). IPFs then would have 45 days from May 1 to submit their data, which would result in a June 15, 2018 submission deadline for this example. Because the exact dates could vary from year to year, for the FY 2019 payment determination and subsequent years, we are also proposing to provide notification of the exact dates of the 45-day submission period through subregulatory means, such as on a CMS Web site and/or on our applicable listservs.

We welcome public comments on our proposals to: (1) Change the specification of the submission deadline from exact dates (that is, July 1–August 15) to a 45-day submission period beginning at least 30 days following the end of the data collection period; and (2) provide notification of the exact dates of the 45-day submission period through subregulatory means for the FY 2019 payment determination and subsequent years.

c. Reporting Requirements for the FY 2019 Payment Determination and Subsequent Years

We refer readers to the FY 2013 IPPS/LTCH PPS final rule (77 FR 53659 through 53660), the FY 2014 IPPS/LTCH PPS final rule (78 FR 50901 through 50902), and the FY 2016 IPPS PPS final rule (80 FR 46715 and 46716), for information about data reporting periods. We are not proposing any changes to these policies in this proposed rule.

d. Population and Sampling

We refer readers to the FY 2013 IPPS/LTCH PPS final rule (79 FR 45973), the FY 2016 IPPS PPS final rule (80 FR 46717 through 46719), for information about population, sampling, and minimum case thresholds. In this proposed rule, we are not proposing any changes to the population and sampling methodology or to the minimum case thresholds.

e. Data Accuracy and Completeness Acknowledgement (DACA) Requirements

We are not proposing any changes to the DACA requirements and refer readers to the FY 2013 IPPS/LTCH PPS final rule (77 FR 53658) for more information on these requirements.

f. Reconsideration and Appeals Procedures

We refer readers to the FY 2013 IPPS/LTCH PPS final rule (77 FR 53658 through 53660), FY 2014 IPPS/LTCH PPS final rule (78 FR 50953), and 42 CFR 412.434 for details on our reconsideration and appeals procedures. We are not proposing any changes to these policies.

9. Extraordinary Circumstances

Exceptions (ECE) Policy for the IPFQR Program

a. Background

In the FY 2013 IPPS/LTCH PPS final rule (77 FR 53659 through 53660), we finalized policies for facilities to request waivers, now called "exceptions" (79 FR 45978), from quality reporting requirements for the FY 2014 payment determination and subsequent years. We stated that in the event of extraordinary circumstances not within the control of IPFs, such as a natural disaster, IPFs may request a reporting extension or a complete waiver of the requirement to submit quality data for one or more quarters for the FY 2014 payment determination and subsequent years. In that rule, we also finalized that facilities would be required to submit a request form with the following information:

- The IPF’s CMS Certification Number (CCN);
- The IPF’s name;
- Contact information for the IPF’s Chief Executive Officer (CEO) and any other designated personnel, including name, email address, telephone number, and mailing address (the address must be a physical address, not a post office box);
- The IPF’s reason for requesting an extension or waiver;
- Evidence of the impact of extraordinary circumstances, including but not limited to photographs, newspaper and other media articles; and
- A date when the IPF will again be able to submit IPFQR Program data, and a justification for the proposed date.

In addition, we finalized that the form must be signed by the IPF’s CEO and submitted within 30 days of the date that the extraordinary circumstance occurred. We also finalized that following the receipt of the request form, we would: (1) Provide a written acknowledgement, using the contact information provided in the request, to the CEO and any additional designated IPF personnel, notifying them that the IPF’s request has been received; and (2) provide a formal response to the CEO and any designated IPF personnel, using the contact information provided in the request, notifying the IPF of our decision. Furthermore, in that rule, we discussed that the above policy does not preclude us from granting waivers or extensions to IPFs that have not requested them when we determine that an extraordinary circumstance has affected an entire region or locale. We stated that if we make the determination to grant such a waiver or extension, we would communicate this decision through routine communication channels (77 FR 53659). In the FY 2014 IPPS/LTCH PPS final rule, we did not make any changes to this policy (78 FR 50903).

In the FY 2015 IPPS PPS final rule (79 FR 45978), we clarified that the term “exception” is synonymous with the term “waiver” used in previous rules and renamed our policy to “Extraordinary Circumstances Exception” in order to align with similar exceptions in other CMS quality reporting programs. In that rule, we also finalized that that we may grant a waiver or extension to IPFs if we determine that a systemic problem with one of our data collection systems directly affects the ability of the IPFs to submit data. We stated that because we do not anticipate that these types of systemic errors will occur often, we do not anticipate granting a waiver or extension on this basis frequently (79 FR 45978). We noted that if we make the determination to grant a waiver or extension, we would communicate this decision through routine communication channels to IPFs, vendors, and quality improvement organizations (QIOs) by means of, for example, memoranda, emails, and notices on the QualityNet Web site (79 FR 45978).

In this proposed rule, we are proposing to modify aspects of our current ECE policy to align with those of other CMS quality reporting programs. Many of our quality reporting and value-based purchasing programs
We believe aligning these five areas across the programs will improve administrative efficiencies for affected facilities or hospitals. We note that, in this FY 2018 IPPS/LTCH PPS proposed rule, we are also proposing to update ECE policies in the Hospital Readmissions Reduction Program (in section V.I.12. of the preamble of this proposed rule), the HAC Reduction Program (in section IX.A.15. of the preamble of this proposed rule), and the PCHQR Program (in section IX.B.10. of the preamble of this proposed rule) in order to align policies. We refer readers to these sections for more details.

b. Proposed ECE Policy Modifications

The IPFQR Program currently includes policies to: (1) Make explicit the ability to grant ECEs specific for systemic issues with CMS data collection systems that directly affect the ability of hospitals/facilities to submit data; and (2) refer to the ECE policy as “extraordinary circumstances exceptions.” Therefore we are not making proposals related to these two items. However, to improve cross-program alignment we are proposing to update the IPFQR Program’s ECE policy by: (1) Allowing designated personnel to sign the ECE request form that IPFs currently submit with contact information for the CEO and designated personnel and the signature from the CEO; (2) extending the deadline from 30 days following the date that the extraordinary circumstance occurred to 90 days following the date the extraordinary circumstance occurred; and (3) specifying that we will strive to provide our formal response to an ECE request within 90 days following the date that the extraordinary circumstance occurred. However, we believe that it may be difficult for some IPFs to timely evaluate the impact of a certain extraordinary circumstance within 30 calendar days. Therefore, we are proposing to change the ECE request form submission deadline from within 30 days of the date that the extraordinary circumstance occurred to within 90 days of the date that the extraordinary circumstance occurred.

We believe that extending the deadline to 90 calendar days would allow IPFs more time to determine whether it is necessary and appropriate to submit an ECE request and to provide a more comprehensive account of the extraordinary circumstance in their ECE request form to CMS. As an example, if an IPF has suffered damage due to a hurricane on October 1, 2017, it would have until December 30, 2017, 90 calendar days after the hurricane, to submit an ECE form via the QualityNet Secure Portal, mail, email, or secure fax as instructed on the ECE form.

(3) Clarification of CMS Response Timeframe

As stated above, in the FY 2013 IPPS/LTCH PPS final rule (77 FR 53659 through 53660), we finalized that ECE requests for the IPFQR Program must be submitted within 30 days of the date that the extraordinary circumstance occurred. However, we believe that the number of requests we receive and the complexity of the information provided affect the timeframe that we need to make ECE determinations. Therefore, in an effort to provide facilities with a predictable timeframe, we are clarifying that we will strive to complete our review of each ECE request as quickly as possible. However, the number of requests we receive and the complexity of the information provided affect the timeframe that we need to make ECE determinations.
We welcome public comments on our proposals to: (1) Specify that ECE forms can be signed by either the CEO or the designated personnel as listed on the ECE form; and (2) change the ECE request form submission deadline to within 90 days of the date that the extraordinary circumstance occurred. We also invite public comments on our intent to clarify that we will strive to complete our review of ECE requests within 90 days of receipt.

E. Clinical Quality Measurement for Eligible Hospitals and Critical Access Hospitals (CAHs) Participating in the EHR Incentive Programs

1. Background

The HITECH Act (Title IV of Division B of the ARRA, together with Title XIII of Division A of the ARRA) authorizes incentive payments under Medicare and Medicaid for the adoption and meaningful use of certified electronic health record (EHR) technology (CEHRT). Incentive payments under Medicare were available to eligible hospitals and CAHs for certain payment years (as authorized under sections 1886(n) and 1814(l) of the Act, respectively) if they successfully demonstrated meaningful use of CEHRT, which includes reporting on clinical quality measures (CQMs or eCQMs) using CEHRT.

Sections 1886(b)(3)(B) and 1814(l) of the Act also establish downward payment adjustments under Medicare, beginning with FY 2015, for eligible hospitals and CAHs that do not successfully demonstrate meaningful use of CEHRT for certain associated reporting periods. Section 1903(a)(3)(F)(i) of the Act establishes 100 percent Federal financial participation (FFP) to States for providing incentive payments to eligible Medicaid providers (described in section 1903(l)(2) of the Act) to adopt, implement, upgrade and meaningfully use CEHRT.

Under sections 1814(l)(3)(A), 1886(n)(3)(A), and 1903(l)(6)(C)(ii) of the Act and the definition of “meaningful EHR user” under 42 CFR 495.4, eligible hospitals and CAHs must report on CQMs selected by CMS using CEHRT, as part of being a meaningful EHR user under the Medicare and Medicaid EHR Incentive Programs.

2. Proposed Modifications to the CQM Reporting Requirements for the Medicare and Medicaid EHR Incentive Programs for CY 2017

a. Background

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57255), we stated the CQM reporting periods in CY 2017 for the Medicare and Medicaid EHR Incentive Programs as outlined below. For the Medicare EHR Incentive Program, we finalized the following submission periods for eligible hospitals and CAHs reporting CQMs by attestation and eligible hospitals and CAHs electronically reporting CQMs (81 FR 57255). In regard to the Medicaid EHR Incentive Program, we provided States with the flexibility to determine the submission periods for reporting CQMs.

- Eligible Hospitals and CAHs Reporting CQMs by Attestation:
  - For eligible hospitals and CAHs demonstrating meaningful use for the first time in 2017, the reporting period is any continuous 90-day period within CY 2017. The submission period for attestation is the 2 months following the close of the calendar year, ending February 28, 2018.
  - For eligible hospitals and CAHs that demonstrated meaningful use in any year prior to 2017, the reporting period is the last 90 days of CY 2017 (consisting of four quarterly data reporting periods). The submission period for attestation is the 2 months following the close of the calendar year, ending February 28, 2018.

- Eligible Hospitals and CAHs Reporting CQMs Electronically: For eligible hospitals and CAHs demonstrating meaningful use for the first time in 2017 or that have demonstrated meaningful use in any year prior to 2017, the reporting period is the full CY 2017 (consisting of four quarterly data reporting periods). The submission period for reporting CQMs electronically begins in late spring 2017 and continues through the 2 months following the close of the calendar year, ending February 28, 2018.

- For Attestation: If only participating in the EHR Incentive Program, report on all 16 available CQMs.
  - For Electronic Reporting: If only participating in the EHR Incentive Program, or participating in both the EHR Incentive Program and the Hospital IQR Program (81 FR 57150 through 57159), report on 8 of the available CQMs.

For further information on the policies applicable for CQM reporting for the EHR Incentive Program in 2017, we refer readers to the discussion in the FY 2017 IPPS/LTCH PPS final rule at 81 FR 57249 through 57257.

Since the publication of the FY 2017 IPPS/LTCH PPS final rule, we have continued to receive frequent feedback from hospitals and EHR vendors about the ongoing challenges of implementing CQM reporting capabilities. A summary of the main concerns identified by these data submitters is as follows:

- The timing of the transition to a new EHR system during 2017 (system upgrades or new EHR vendor) may influence hospitals’ ability to report in a timely manner;
- The current timeframe for the implementation of new EHR requirements presents challenges due to the varying 6 to 24-month cycles needed for vendors to code new measures, test and institute measure updates, train hospital staff, and rollout other upgraded features;
- Hospitals have had difficulty identifying applicable measures that reflect their patient population, given the reduction in the number of available CQMs from 29 to 16 for CY 2017; and
- Hospitals have had challenges with data mapping and workflow because of the need to collect CY 2017 data while still reporting CY 2016 data.

In addition, there have been other recent issues related to the CMS data receiving system not being able to process QRDA Category I files, and as a result, the system is not generating notifications confirming for providers that their files have been received and processed by the system. The aforementioned issues and challenges being experienced by hospitals and vendors are impacting the capability of hospitals to meet the requirements for CY 2017. As a result, we are proposing modifications to the CY 2017 final policies in this proposed rule, which would reduce CQM reporting requirements in order for hospitals and vendors to address these issues.

In this proposed rule, we are proposing two modifications to our CY 2017 electronic CQM reporting policies for the Medicare and Medicaid EHR Incentive Programs. For eligible hospitals and CAHs reporting CQMs electronically in CY 2017, we are proposing to:

1. Decrease the number of calendar quarters for which such hospitals are required to submit data; and
2. decrease the number of CQMs for which such hospitals must submit data (further discussion below). These proposals are made in conjunction with our proposals discussed in sections IX.A.8. and IX.A.10.d. of the preamble of this proposed rule to align requirements for the Medicare and Medicaid EHR Incentive Programs and the Hospital IQR Program. In making these proposals, we believe that eligible...
hospitals and CAHs would have additional time to upgrade their systems and processes in preparation for the transition to electronic reporting on additional CQMs for additional quarters in future years.

As we continue to make strides with electronic reporting, we want to ensure we provide eligible hospitals and CAHs with a robust selection of CQMs. As noted above, hospitals have expressed concerns with identifying applicable measures that reflect their patient population; thus, we believe that the addition of new CQMs in the future will offer more clinically relevant CQMs that facilitate reporting and help drive quality improvement. In section IX.A.9.d. of the preamble of this proposed rule, we discuss and seek feedback on future potential CQMs for the Hospital IQR Program and the Medicare and Medicaid EHR Incentive Programs for eligible hospitals and CAHs.

b. Proposed Changes to Policies Regarding Electronic Reporting of CQMs for CY 2017

In response to concerns from stakeholders, we are proposing to modify the CQM reporting period for eligible hospitals and CAHs reporting CQMs electronically for the Medicare and Medicaid EHR Incentive Programs in CY 2017—for eligible hospitals and CAHs demonstrating meaningful use for the first time in 2017 or that have demonstrated meaningful use in any year prior to 2017, the reporting period would be two self-selected quarters of CQM data in CY 2017.

In addition, we are proposing to modify the reporting criteria regarding the required number of CQMs for eligible hospitals and CAHs that are reporting electronically for the reporting periods in CY 2017 under the Medicare and Medicaid EHR Incentive Programs—if only participating in the EHR Incentive Program, or participating in both the EHR Incentive Program and the Hospital IQR Program, eligible hospitals and CAHs would report on at least 6 (self-selected) of the available CQMs. For a list of the available CQMs for reporting periods in CY 2017, we refer readers to the table in the FY 2017 IPPS/LTCH PPS final rule at 81 FR 57255.

It should be noted that we are not proposing to modify any other aspects of the policies for reporting CQMs electronically for CY 2017, including the submission periods, nor are we proposing any changes to our policies for reporting CQMs by attestation.

Through our proposals for CY 2017, we intend to continue to maintain alignment between the Medicare and Medicaid EHR Incentive Programs and the Hospital IQR Program to reduce confusion and reporting burden among participants in the Medicare and Medicaid EHR Incentive Programs that also participate in the Hospital IQR Program. As noted above, we are retaining the submission period for reporting CQMs electronically under the Medicare EHR Incentive Program, in which such submission period begins in late spring 2017 and continues through the 2 months following the close of the calendar year, ending February 28, 2018. In addition, we are continuing to provide States with the flexibility to determine the submission periods for reporting CQMs under the Medicaid EHR Incentive Program. For more details on the aligned reporting requirements for the Hospital IQR and Medicare and Medicaid EHR Incentive Programs, we refer readers to section IX.A.10.d. of the preamble of this proposed rule.

We believe that reducing the number of CQMs required to be electronically reported from 8 to 6 would ease the burden on data submitters, allowing them to shift resources to support system upgrades, map data, and train staff on CQMs. Reducing the number of data reporting periods to 2 quarters, rather than 4 quarters, and allowing eligible hospitals and CAHs to select which two quarters of CY 2017 to electronically report would offer greater reporting flexibility and allow eligible hospitals, CAHs, and vendors more time to plan for reporting, and account for and schedule hospital-specific scenarios such as EHR upgrades or system transitions. We recognize that eligible hospitals and CAHs are concerned about their capability of meeting the CY 2017 requirements established in the FY 2017 IPPS/LTCH PPS final rule and believe that these modified reporting requirements for CY 2017 account for the challenges stakeholders are experiencing while requiring the electronic reporting on an additional portion of CQMs, which is consistent with our goal to transition to electronic reporting (81 FR 57254).

We are inviting public comment on our proposals to modify the CY 2017 CQM reporting requirements for the Medicare and Medicaid EHR Incentive Programs as described above.

3. CQM Reporting for the Medicare and Medicaid EHR Incentive Programs in 2018

a. Background

In the 2015 EHR Incentive Programs Final Rule (80 FR 62892 through 62893), beginning in CY 2017 and for subsequent years, we established a CQM reporting period of one full calendar year (consisting of four quarterly data reporting periods) for the reporting of CQMs by eligible hospitals and CAHs participating in the Medicare and Medicaid EHR Incentive Programs, with an exception for providers demonstrating meaningful use for the first time under the Medicaid EHR Incentive Program, for whom the CQM reporting period is any continuous 90-day period within the calendar year. In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57250), we noted that one full calendar year of data will result in more complete and accurate data, and hospitals will be able to submit one full calendar year of data for both the Medicare and Medicaid EHR Incentive Programs and the Hospital IQR Program, thereby reducing the reporting burden.

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57250 through 57255), we removed 13 CQMs from the set of CQMs available for eligible hospitals and CAHs to report under the Medicare and Medicaid EHR Incentive Programs, beginning with the reporting periods in CY 2017. All 16 of the remaining measures listed in Table 10 of the EHR Incentive Program Stage 2 final rule (77 FR 54083 through 54087) are available for eligible hospitals and CAHs to report for the Medicare and Medicaid EHR Incentive Programs. The following table lists the 16 CQMs available for eligible hospitals and CAHs to report for the Medicare and Medicaid EHR Incentive Programs beginning in CY 2017 (81 FR 57255).

<table>
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<tr>
<td>ED–3</td>
<td>Median Time from ED Arrival to ED Departure for Discharged ED Patients</td>
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CQMs FOR ELIGIBLE HOSPITALS AND CAHs BEGINNING WITH CY 2017

Electronic Clinical Quality Measures (eCQMs)
For CY 2018 and future calendar years, we plan to continue to align the CQM reporting requirements for the Medicare and Medicaid EHR Incentive Programs and the Hospital IQR Program. As we expect to expand the current measures to align with the National Quality Strategy and the CMS Quality Strategy and incorporate updated standards and terminology in current CQMs, including updating the electronic specifications for these CQMs, and creating de novo CQMs, we plan to expand the set of CQMs available for reporting under the EHR Incentive Programs in future years. We will continue to engage stakeholders to provide input on future proposals for CQMs as well as request comment on future electronic specifications for new and updated CQMs.

b. CQM Reporting Period for the Medicare and Medicaid EHR Incentive Programs in CY 2018

(1) Background

Our goal is to continue to move toward increased electronic reporting while also addressing stakeholder concerns as described above, which we believe will likely continue into CY 2018, but to a lesser extent than in CY 2017. With the CY 2017 proposed policies reducing reporting requirements and providing additional time for eligible hospitals, CAHs, and vendors to make EHR upgrades and system transitions in CY 2017, we believe that stakeholders would be able to address some of the issues and challenges they face prior to CY 2018, but recognize that certain challenges and issues (for example, EHR upgrade and system transition challenges associated with the development cycle of technology and the timeframe to develop and execute work flows and processes and train staff based on EHR upgrades and system transitions) may not be fully resolved and as a result, may persist in CY 2018. As established in the 2015 EHR Incentive Programs Final Rule (80 FR 62894), reporting CQMs by attestation will no longer be an option for eligible hospitals and CAHs starting with the reporting periods in CY 2018, except in circumstances in which electronic reporting is not feasible.

For CY 2018, we are proposing the following CQM reporting period for the Medicare and Medicaid EHR Incentive Programs and the following submission period for the Medicare EHR Incentive Program—for eligible hospitals and CAHs reporting CQMs electronically that demonstrate meaningful use for the first time in 2018 or that have demonstrated meaningful use in any year prior to 2018, the reporting period would be the first 3 quarters of CY 2018, and the submission period would be the 2 months following the close of the calendar year, ending February 28, 2019.

For eligible hospitals and CAHs that report CQMs by attestation under the Medicare EHR Incentive Program as a result of electronic reporting not being feasible, and for eligible hospitals and CAHs that report CQMs by attestation under their state’s Medicaid EHR Incentive Program, we established a CQM reporting period of the full CY 2018 (consisting of 4 quarterly data reporting periods) (80 FR 62893). We have established an exception to this full-year reporting period for eligible hospitals and CAHs demonstrating meaningful use for the first time under their state’s Medicaid EHR Incentive Program; under this exception, the CQM reporting period is any continuous 90-day period within CY 2018 (80 FR 62893).

In this proposed rule, we are proposing the submission period for eligible hospitals and CAHs reporting CQMs by attestation under the Medicare EHR Incentive Program would be the 2 months following the close of the CY 2018 CQM reporting period, ending February 28, 2019.

In regard to the Medicaid EHR Incentive Program, we provide States with the flexibility to determine the method of reporting CQMs (attestation or electronic reporting) and the submission periods for reporting CQMs, subject to prior approval by CMS.

(2) CQM Reporting Criteria for the Medicare and Medicaid EHR Incentive Programs in CY 2018

We are proposing the following reporting criteria under the Medicare and Medicaid EHR Incentive Program for eligible hospitals and CAHs reporting CQMs electronically for the reporting period in CY 2018—eligible hospitals and CAHs participating only in the EHR Incentive Program, or participating in both the EHR Incentive Program and the Hospital IQR Program, report on at least six (self-selected) of the available CQMs from the table in the FY 2017 IPPS/LTCH PPS final rule at 81 FR 57255.

We are proposing the following reporting criteria for eligible hospitals and CAHs that report CQMs by attestation under the Medicare EHR Incentive Program because electronic reporting is not feasible, and for eligible hospitals and CAHs that report CQMs by attestation under their state’s
Medicaid EHR Incentive Program, for the reporting period in CY 2018—report on all 16 available CQMs from the table in the FY 2017 IPPS/LTCH PPS final rule at 81 FR 57255.

In developing these proposals, we considered several alternatives. Specifically, we considered aligning the requirements for CY 2018 with the proposed requirements for CY 2017 outlined in this proposed rule, such that eligible hospitals and CAHs would report on 6 (self-selected) available CQMs for each of the four quarters of data in both CY 2017 and CY 2018. We also considered the final policy in the FY 2017 IPPS/LTCH PPS final rule for the Hospital IQR Program (81 FR 57150 through 57159), which would require hospitals to report one full calendar year of data for at least 8 (self-selected) CQMs out of the available CQMs for both the CY 2017 reporting period/FY 2019 payment determination and the CY 2018 reporting period/FY 2020 payment determination. However, we are proposing changes to this previously adopted policy in the Hospital IQR Program and refer readers to section IX.A.8. of the preamble of this proposed rule for more details. Ultimately, we believe that our proposal balances our goal to shift towards electronic reporting of quality measure data with concerns from stakeholders regarding an increased burden to meet CQM reporting requirements.

In addition, the proposal provides eligible hospitals and CAHs with the opportunity to have several years of experience reporting data electronically for the Hospital IQR and Medicare and Medicaid EHR Incentive Programs. Therefore, we believe that eligible hospitals and CAHs will be better prepared to submit an additional quarter of data for the CY 2018 reporting period compared to the number of quarters we are proposing for the CY 2017 reporting period. This proposal is being made in conjunction with our proposals discussed in section IX.A.10.d. of the preamble of this proposed rule to align requirements for the Medicare and Medicaid EHR Incentive Programs and the Hospital IQR Program.

We are inviting public comment on our proposals regarding the CY 2018 reporting requirements for eligible hospitals and CAHs reporting CQMs under the Medicare and Medicaid EHR Incentive Programs.

c. CQM Reporting Form and Method for the Medicare EHR Incentive Program in 2018

In the FY 2016 IPPS/LTCH PPS final rule (80 FR 49759 through 49760), we removed the QRDA–III as an option for reporting under the Medicare EHR Incentive Program for eligible hospitals and CAHs. For the reporting periods in 2016 and future years, we are requiring QRDA–I for CQM electronic submissions for the Medicare EHR Incentive Program. As noted in the FY 2016 IPPS/LTCH PPS final rule (80 FR 49760), States would continue to have the option, subject to our prior approval, to allow or require QRDA–III for CQM reporting.

As noted in the FY 2016 IPPS/LTCH PPS final rule (80 FR 49760), we encourage health IT developers to test any updates, including any updates to the CQMs and CQS reporting requirements based on the CMS Implementation Guide for Quality Reporting Document Architecture (QRDA) Category I and Category III (CMS Implementation Guide for QRDA) for Hospital Quality Reporting (HQR), on an annual basis.

The form and method of electronic submission are further explained in subregulatory guidance and the certification process. For example, the following documents are updated annually to reflect the most recent CQM electronic specifications: The CMS Implementation Guide for QRDA; program specific performance calculation guidance; and CQM electronic specifications and guidance documents. These documents are located on the eCQI Resource Center Web page at: https://ecqi.healthit.gov/. For further information on CQM reporting, we refer readers to the EHR Incentive Program Web site where guides and tips sheets are located at: http://www.cms.gov/ehrincentiveprograms.

For the CY 2018 reporting period, we are proposing the following for CQM submission under the Medicare EHR Incentive Program:

- Eligible hospital and CAH participating in the Medicare EHR Incentive Program (single program participation)—electronically report CQMs through QualityNet Portal.
- Eligible hospital and CAH options for electronic reporting for multiple programs (that is, EHR Incentive Program and Hospital IQR Program participation)—electronically report through QualityNet Portal.

As noted in the 2015 EHR Incentive Programs Final Rule (80 FR 62894), starting in 2018, eligible hospitals and CAHs participating in the Medicare EHR Incentive Program must electronically report CQMs where feasible; and attestation to CQMs will no longer be an option except in certain circumstances where electronic reporting is not feasible.

For the Medicaid EHR Incentive Program, States continue to be responsible for determining whether and how electronic reporting of CQMs would occur, or if they wish to allow reporting through attestation. Any changes that States make to their CQM reporting methods must be submitted through the State Medicaid Health IT Plan (SMHP) process for CMS review and approval prior to being implemented.

For CY 2018, we are proposing to continue our policy regarding the electronic submission of CQMs, which would require the use of the most recent version of the CQM electronic specification for each CQM to which the EHR is certified. For the CY 2018 electronic reporting of CQMs, this means eligible hospitals and CAHs would be required to use the Spring 2017 version of the CQM electronic specifications and any applicable addenda available on the eCQI Resource Center Web page at: https://ecqi.healthit.gov/. In addition, we are proposing to require that an eligible hospital or CAH would need to have its EHR technology certified to all 16 available CQMs from the table in the FY 2017 IPPS/LTCH PPS final rule at 81 FR 57255 in order to meet the reporting requirements for CY 2018. As described in the 2015 EHR Incentive Programs Final Rule (80 FR 62767), starting in CY 2018, eligible hospitals and CAHs are required to have EHR technology certified to the 2015 Edition.

Starting in CY 2018, we are proposing to require the use of EHR technology certified to the 2015 Edition for CQM reporting. Furthermore, we are proposing that an EHR certified for CQMs under the 2015 Edition certification criteria would not need to be recertified each time it is updated to a more recent version of the CQMs. We believe it is not necessary for an EHR certified for CQMs under the 2015 Edition certification criteria to be recertified each time it is updated to the most recent version of the CQMs because the EHR technology continues to meet the 2015 Edition certification criteria and any updates to the CQM specifications do not impact or change any elements regarding certification and thus, we are proposing that recertification is not necessary. For further discussion regarding EHR certification requirements for 2018, we refer readers to section IX.G.4. of the preamble of this proposed rule. We are inviting public comment on these proposals.
F. Clinical Quality Measurement for Eligible Professionals (EPs) Participating in the Medicaid EHR Incentive Program in 2017

The proposals in this section would apply only to EPs participating in the Medicaid EHR Incentive Program. They would not apply to eligible hospitals or CAHs, or to the Medicare EHR Incentive Program.

1. Proposed Modifications to the CQM Reporting Period for EPs in 2017

In the 2015 EHR Incentive Programs Final Rule (80 FR 62672), we established for the Medicare and Medicaid EHR Incentive Programs a CQM reporting period of the full CY 2017 for EPs who have demonstrated meaningful use in a prior year and a CQM reporting period of any continuous 90 days within CY 2017 for EPs who are demonstrating meaningful use for the first time (80 FR 62891 through 62892). We also noted that we would continue to allow the States to determine the form and manner in which Medicaid EPs should report CQMs, subject to CMS approval (80 FR 62891, 62894).

In the final rule with comment period titled Medicare Program; Merit-Based Incentive Payment System (MIPS) and Alternative Payment Model (APM) Incentive Under the Physician Fee Schedule, and Criteria for Physician-Focused Payment Models (81 FR 77008) (referred to as the “CY 2017 Quality Payment Program final rule with comment period’’), we established at § 414.1320(a), for the 2019 MIPS payment year, a minimum of a continuous 90-day performance period within CY 2017, up to and including the full CY 2017, for the quality performance category of the MIPS. We established at § 414.1320(b), for the 2020 MIPS payment year, a performance period of the full CY 2018.

Following the publication of that final rule with comment period, we received feedback from EPs observing that having CQM reporting or performance periods for Medicare professionals under MIPS that are different from the CQM reporting period for EPs under the Medicaid EHR Incentive Program would create administrative burdens for EPs who wish to participate in both programs and to report CQMs electronically. Our goal has always been to align Medicare and Medicaid reporting and quality improvement programs to the extent possible. In addition, while participation in MIPS is required for professionals who are considered “MIPS eligible clinicians,” participation in the Medicaid EHR Incentive Program is not required. If the CQM reporting periods and MIPS performance periods are not aligned, we believe it is less likely that MIPS eligible clinicians will also participate as EPs in the remaining years of the Medicaid EHR Incentive Program.

Therefore, we are proposing to change the CQM reporting period for EPs who report CQMs electronically in the Medicaid EHR Incentive Program to match the performance period established under MIPS in the quality performance category for MIPS eligible clinicians. We are proposing a minimum of a continuous 90-day period during CY 2017 for EPs electronically reporting CQMs for the Medicaid EHR Incentive Program. We note that we consider the reporting periods established through rulemaking to be minimums and would encourage States to accept data from longer reporting periods. The reporting period for CQMs for EPs who choose to attest rather than report electronically, and who have demonstrated meaningful use in a previous program year, under the EHR Incentive Program would remain one full year (CY 2017), which is in alignment with the requirements for eligible hospitals and CAHs for the Medicare and Medicaid EHR Incentive Programs for 2017 (80 FR 62892 through 62893).

We note that reporting CQMs by attestation is not an option for eligible clinicians under MIPS, so the reason for proposing a shortened reporting period for EPs reporting CQMs electronically, which is to align this reporting period with the MIPS performance period, would not exist for EPs who choose not to report electronically. Nothing in this proposal would change the CQM reporting period for EPs demonstrating meaningful use for the first time, which was established in the 2015 EHR Incentive Programs Final Rule to be any continuous 90 day period regardless of the method of CQM submission (80 FR 62892).

The CQM reporting period for the Medicaid EHR Incentive Program in 2018 for EPs that have demonstrated meaningful use in a previous program year would remain one full year (CY 2018) to align with the corresponding performance period in MIPS for MIPS eligible clinicians. If changes are made to the MIPS performance period through future rulemaking, we will revisit the Medicaid EHR Incentive Program policies to continue our alignment goals.

We intend to reduce EP burden and simplify the program through this proposal, which is intended to better align CQM reporting periods and CQM reporting for the Medicaid EHR Incentive Program with policies under MIPS. Overall, we believe the proposed alignment at the State attestation system and EP levels would both reduce burden associated with reporting on multiple CMS programs and enhance State and CMS operational efficiency.

We are inviting public comment on this proposal, including on whether making the proposed change would create burdens for EPs or States.

2. Proposed Modifications to CQM Reporting Requirements for Medicaid EPs Under the Medicaid EHR Incentive Program

We also are proposing to align the specific CQMs available to EPs participating in the Medicaid EHR Incentive Program with those available to clinicians participating in MIPS who submit CQMs through their EHR. In the final rule titled “Medicare and Medicaid Programs; Electronic Health Record Incentive Program—Stage 2,” we established (77 FR 54069) that EPs are required to report 9 CQMs covering at least 3 of the National Quality Strategy (NQS) domains from a list of 64 CQMs (77 FR 54069, Table 8). Subsequently and in the following years, in general, there has been alignment between the CQMs selected for the Medicaid and Medicare EHR Incentive Programs for EPs and the electronic measures selected for the PQRS program. Updates to the PQRS measure set were proposed and finalized in the annual Physician Fee Schedule (PFS) rule for purposes such as keeping specifications in line with industry standards and clinical guidelines.

In the CY 2017 Quality Payment Program Final rule with comment period (81 FR 77144), we revised the list of CQMs for the 2019 MIPS payment year, based on performance periods within CY 2017, to better reflect updated clinical standards and guidelines. Specifically, we removed a number of CQMs that had not been updated and were no longer clinically relevant (81 FR 77773, Appendix, Table F). Because MIPS is replacing PQRS, in order to keep CQM specifications current, we are proposing to align the CQMs for Medicaid EPs with those updated annually for MIPS. Specifically, we are proposing that the CQMs available for Medicaid EPs in 2017 would consist of those updated for MIPS in 2017, to better reflect updated clinical standards and guidelines.

In the CY 2017 Quality Payment Program Final rule with comment period under Table A, which are denoted with a CMS e-Measure ID number.

In the CY 2017 Quality Payment Program Final rule with comment period...
We propose that for 2017 Medicaid EHR Incentive Programs we would reduce the burden for Medicaid EPs, and that the systems changes that would be needed to implement it would not be significant for either States or EPs. The set of 53 CQMs available to MIPS participants is a subset of the 64 CQMs currently available under the Medicaid EHR Incentive Program. In addition, we believe that if EPs also plan to participate in MIPS, they should already be prepared to report on the 53 CQMs. However, we welcome comments on whether any EPs might be negatively affected by the proposal; for example, on whether any EPs might have EHRs that do not measure enough of the 53 remaining CQMs because they were relying on some of the 11 CQMs that would be removed. We do not anticipate that this would be a common situation because these 11 CQMs are outdated, and the industry is moving away from them as EHRs are upgraded to meet the MIPS requirements.

We anticipate that the proposal to reduce the number of available CQMs would have only a minimal impact on States, which would have to make minor adjustments to State systems to reduce the available measures from 64 to 53. It is our understanding that State systems can turn off or easily exclude CQMs from user visibility on the front end and still easily manage on the back end.

The data submission criteria for the MIPS quality performance category at § 414.1335(a)(1)(i) provide that individual MIPS eligible clinicians and groups who elect to submit data via claims, qualified registry, EHR or qualified clinical data registry must submit data on at least six quality measures, including at least one outcome measure (or, if an applicable outcome measure is not available, one other high priority measure). We refer readers to § 414.1335(a)(2) and (3) for the data submission criteria that apply to individual MIPS eligible clinicians and groups who elect to submit data via other data submission mechanisms.

Instead of requiring MIPS eligible clinicians to report on CQMs across a certain number of NQS domains, MIPS provides individual MIPS eligible clinicians and groups with a variety of alternatives for participating in MIPS, including a data submission mechanisms and scoring criteria. We believe that the burden on EPs and States of adopting all of these MIPS alternatives for the Medicaid EHR Incentive Program would outweigh any benefits gained. The alternative reporting options for MIPS are calibrated as part of an overall quality improvement program beyond what the Medicare and Medicaid EHR Incentive Programs are designed to be. We believe it would be inappropriate to apply all of these new requirements to the Medicaid EHR Incentive Program.

We are proposing to eliminate the requirement to report on CQMs across 3 of the 6 NQS domains that existed in previous years of the Medicaid EHR Incentive Program, for improved alignment with the data submission criteria for the MIPS quality performance category. The removal of this requirement would provide EPs greater flexibility in selecting CQMs to report and would assure that they could report on the same CQMs from their EHR to both MIPS and the Medicaid EHR Incentive Program.

We propose that for 2017 Medicaid EPs would be required to report on any six measures that are relevant to the EP’s scope of practice. This proposal would better align with the data submission criteria for the MIPS quality performance category in 2017.

We note that we would continue our policy on allowing zero denominators to be reported to allow EPs to meet the CQM reporting requirements of the EHR Incentive Programs (80 FR 62889)

Future years’ requirements for reporting CQMs in the Medicaid EHR Incentive Program will be established in future rulemaking, as the policies for MIPS are developed for 2018 and beyond. We will continue to align the quality reporting requirements, as logical and feasible, to reduce EP burden.

We are inviting public comment on these proposals, specifically on whether making these proposed changes to CQM measures and measure reporting effective for 2017 would create burdens on EPs or States. If so, CMS will consider making these proposed changes to the CQM reporting requirements effective beginning with the reporting period in 2018.

G. Changes to the Medicare and Medicaid EHR Incentive Programs

1. Proposed Revisions to the EHR Reporting Period in 2018

We received additional feedback from EPs, hospitals, hospital associations, and other health care providers indicating that additional time may be necessary for testing and implementation of the new application programming interface (API) functionality requirement for Stage 3 citing inability to meet the required timeframe for implementation of Stage 3 and complexity of the new functionality and associated requirements for the Patient Electronic Access to Health Information (80 FR 62841 through 62846) and Coordination of Care Through Patient Engagement (80 FR 62846 through 62852) objectives. The API functionality supports health care providers and patient electronic access to health information, which is key to improving the free flow of health information, quality improvement, and patient engagement. Because this functionality is included as part of the 2015 Edition Base EHR definition (and thus must be part of CEHRT) (80 FR 62675 through 62676), APIs may be enabled by a health care provider or organization for their own use of third party applications with their CEHRT, such as for quality improvement. An API could also be enabled by a health care provider to give patients access to their health information through a third-party application with more flexibility than is often found in many current patient portals. From the health care provider perspective, an API could complement a specific provider branded patient portal or could also potentially make one unnecessary if patients are able to use software applications designed to interact with an API that could support their ability to view, download, and transmit their health information to a third party (80 FR 62842). We want to ensure that health
care providers have the opportunity to thoroughly test their systems and make adjustments in order to successfully attest for the EHR reporting period in CY 2018. In addition, we believe that health care providers may need extra time to fully implement and test workflows with the 2015 Edition of CEHRT, which is required beginning in CY 2018 (80 FR 62874 through 62875). The Office of the National Coordinator for Health Information Technology (ONC) monitors technical development and progress toward certification to evaluate readiness among the health IT industry for implementation of technology certified to 2015 Edition certification criteria. One part of this evaluation involves monitoring products in the certification process, which is supplemented by discussions with health IT developers, ONC-Authorized Testing Laboratory (ONC–ATLs), and ONC- Authorized Certification Body (ONC–ACBs). Health IT developers have conveyed to the ONC that some of the 2015 Edition certification criteria required additional effort, including implementation of new functionalities (including APIs) and facilitation of greater interoperability in comparison to previous Editions. The 2015 Edition health IT certification criteria enables health information exchange through new and enhanced certification criteria standards, and implementation specifications for interoperability while incorporating changes that are designed to spur innovation and provide more choices to health care providers and patients for the exchange of electronic health information including new application access (API) certification criteria. For example, new transitions of care certification criterion rigorously assesses a product’s ability to create and receive an interoperable Consolidated-Clinical Document Architecture (C–CDA). The ONC also adopted certification criteria that both support interoperability in other settings and use cases, such as the Common Clinical Data Set summary record, data segment for privacy, and care plan certification criteria (80 FR 62603). It also indicated that it did not anticipate any significant delays toward the delivery and roll out of products to customers because of this additional effort. This timing was expressed as a goal of developers to enable health care providers to engage in upgrades, training, and other improvements that would be needed to begin using the 2015 Edition CEHRT in 2018.

In addition, the ONC also compares data such as program tracking and projections related to the release of the previous Editions of CEHRT to help inform its evaluation of progress during this current transition period for the 2015 Edition. In particular, ONC has reviewed historical data for actual participation and implementation of technologies certified to the 2014 Edition during the transition year in CY 2014, when users were transitioning from technology certified to the 2011 Edition to technology certified to the 2014 Edition. In 2014, projections indicated expectations of market readiness of greater than 90 percent by the end of CY 2014. However, subsequent analysis found that the actual market coverage for hospitals was approximately 98 percent at the end of CY 2014, meaning that 98 percent of hospitals had implemented the 2014 Edition by the end of CY 2014. However, attestations for the EHR Incentive Program for CY 2014 indicated a potential lag in implementation and variability in the amount of time required by hospitals to complete implementation of the technology and the subsequent training, technical processes, and operational and clinical workflows required for successful use. Attestation data show that 9 percent of eligible hospitals and CAHs used EHR technology certified to the 2011 Edition for part or all of their EHR reporting period. In addition, ONC considers the number of health care providers likely to be covered by the individual developers seeking certification under the ONC Health IT Certification Program. The ONC considers trends within the industry when projecting for 2015 Edition readiness. The market trend of consolidation was considered as part of the projection model and supports an analysis that supports an estimate of greater than 85 percent of hospitals will be ready by the end of CY 2017. However, a more conservative approach—based on the identified variance in implementation timelines for hospitals may be necessary to support the hospitals that may require additional time to successfully implement technology certified to the 2015 Edition.

In addition, the historical data indicates EPs are more likely to use a wider range of products, including products which individually make up a smaller segment of the overall market. Therefore, when market factors are taken into account, there exists a larger proportion of readiness that is unknown due to the wider range of products used by EPs. Therefore, a more conservative approach is necessary and supports an estimate of greater than 74 percent readiness by the end of CY 2017 for EPs. Thus, while we expect a majority of EPs, eligible hospitals and CAHs participating in the EHR Incentive Programs to be ready to begin using 2015 Edition CEHRT in CY 2018, it is reasonable to assume there will still be some who will not be ready and will require a longer timeframe for successful implementation. In addition, it is likely that there will be a proportion of them that have the technology implemented in time for the beginning of CY 2018 who would similarly benefit from additional time to implement new processes and workflows supporting their use of certified EHR technology in the EHR Incentive Program. This is especially important given requirements in the ONC Health IT Certification program which leverage new functionalities such as APIs which also require adherence to existing security and privacy standards. We stated in the 2015 EHR Incentive Programs Final Rule (80 FR 62842 through 62843) that the requirement to conduct and review a security risk analysis in compliance with HIPAA Security Rule would include the certified API enabled as a part of the health care provider’s CEHRT.

For the reasons discussed above, we are proposing to modify the EHR reporting periods in 2018 for new and returning participants attesting to CMS or their State Medicaid agency from the full year (CY 2018) to a minimum of any continuous 90-day period within CY 2018. This would mean that EPs that attest directly to a State for the State’s Medicaid EHR Incentive Program and eligible hospitals and CAHs attesting to CMS or the State’s Medicaid EHR Incentive Program would attest to meaningful use of CEHRT for an EHR reporting period of a minimum of any continuous 90-day period from January 1, 2018 through December 31, 2018. The applicable incentive payment year and payment adjustment years for the EHR reporting period in 2018, as well as the deadlines for attestation and other related program requirements, would remain the same as established in prior rulemaking. We are proposing corresponding changes to the definition of “EHR reporting period” and “EHR reporting period for a payment adjustment year” at 42 CFR 495.4.

We are inviting public comment on our proposal.

2. Exception for Decertified EHR Technology for EPs, Eligible Hospitals, and CAHs Seeking To Avoid the Medicare Payment Adjustment

The 21st Century Cures Act (Pub. L. 114–255) was enacted on December 13, 2016. Section 4002(b)(1)(A) amended
section 1848(a)(7)(B) of the Act to provide that the Secretary shall exempt an eligible professional from the application of the payment adjustment under section 1848(a)(7)(A) of the Act with respect to a year, subject to annual renewal, if the Secretary determines that compliance with the requirement for being a meaningful EHR user is not possible because the certified EHR technology used by such professional has been decertified under ONC’s Health IT Certification Program. Similarly, section 4002(b)(2) of the 21st Century Cures Act amended section 1886(b)(3)(B)(ix)(II) of the Act to provide that the Secretary shall exempt a hospital from the application of the payment adjustment under section 1886(b)(3)(B)(ix)[I] with respect to a fiscal year, subject to annual renewal, if the Secretary determines that compliance with the requirement for being a meaningful EHR user is not possible because the certified EHR technology used by the hospital is decertified under ONC’s Health IT Certification Program. We include proposals below to implement these amendments with respect to EPs, eligible hospitals, and CAHs. We note that sections 1848(a)(7)(B) and 1886(b)(3)(B)(ix)[II] of the Act provide that in no case may an EP, eligible hospital, or CAH be granted an exemption from the payment adjustment based on significant hardship or decertified EHR technology for more than five years. The ONC Health IT Certification Program: Enhanced Oversight and Accountability final rule (“EOA final rule”) (81 FR 72404), effective December 19, 2016, created a regulatory framework for the ONC’s direct review of health information technology (health IT) certified under the ONC Health IT Certification Program, including, when necessary, requiring the correction of non-conformities found in health IT certified under the Program and terminating certifications issued to certified health IT. Prior to the EOA final rule, ONC–Authorized Certification Bodies (ONC–ACBs) had the only authority to terminate or revoke certification of health IT under the program, which they used on previous occasions. On September 23, 2015, we posted an FAQ discussing the requirements for using a decertified CEHRT.498 Once all administrative processes, if any, are complete, then notice of a “termination of certification” is listed on the Certified Health IT Product List (CHPL) Web page.499 As appropriate, ONC will also publicize the termination of certification of health IT through other communication channels (for example, ONC list serve(s)). Further, when ONC terminates the certification of a health IT product, the health IT developer is required to notify all potentially affected customers in a timely manner.

We further note that in comparison to termination actions taken by ONC and ONC–ACBs, a health IT developer may voluntarily withdraw a certification that is in good standing under the ONC Health IT Certification Program. A voluntary withdrawal may be the result of the health IT developer going out of business, the developer no longer supporting the product, or for other reasons that are not in response to ONC–ACB surveillance, ONC direct review, or a finding of non-conformity by ONC or an ONC–ACB.500 In such instances, ONC will list these products on the “Inactive Certificates” Web page of the CHPL.501 We are proposing to revise § 495.102(d) to add a new exception for EPs who demonstrate through an application process that compliance with the requirement for being a meaningful EHR user is not possible because the certified EHR technology used by the EP has been decertified under ONC’s Health IT Certification Program. We are proposing this exception for eligible hospitals that voluntarily withdraw a certification under ONC’s Health IT Certification Program. We are proposing this exception for the CY 2018 payment adjustment year, which is the final year of the payment adjustment for EPs under section 1848(a)(7)(A) of the Act. We considered but are not proposing this exception also for the CY 2017 payment adjustment year because it would require us to reprocess claims for potentially the entire CY 2017, which would be costly and administratively burdensome. ONC provides that there is a 6-step process that usually occurs when implementing a certified EHR technology system.502 We believe that if an EP has to procure new certified EHR technology they will likely have to go through some phases of this cycle again and understand that it would be time consuming and may take up to a year to implement.

We are proposing an EP may qualify for this exception if their certified EHR technology was decertified either before or during the applicable EHR reporting period for the CY 2018 payment adjustment year, which under § 495.4 is any continuous 90-day period in CY 2016 or 2017, depending on whether the EP has successfully demonstrated meaningful use in a prior year. If the certified EHR technology was decertified at any time during the 12-month period preceding the applicable EHR reporting period for the CY 2018 payment adjustment year, or during the applicable EHR reporting period for the CY 2018 payment adjustment year, the EP may qualify for this exception. For example, if an EP intended to attest to meaningful use for a 90-day EHR reporting period beginning on April 1, 2016, the EP could apply for this exception if their certified EHR technology was decertified at any time during the 12-month period beginning on April 1, 2015 and ending on March 31, 2016, or if their certified EHR technology was decertified at any time during their 90-day EHR reporting period beginning on April 1, 2016. We believe a 12-month period is reasonable because we understand the burden placed on EPs related to time and funds needed to purchase and deploy new certified EHR technology including the process that goes along with implementing new certified EHR technology.

In addition, we are proposing that the EP must demonstrate in its application and through supporting documentation if available that the EP intended to attest to meaningful use for a certain EHR reporting period and made a good faith effort to adopt and implement another CEHRT in advance of that EHR reporting period. We are proposing an EP seeking to qualify for this exception would submit an application in the form and manner specified by us by October 1, 2017, or a later date specified by us.

We are proposing to revise § 412.64(d)(4) to add a new category of exception for eligible hospitals that demonstrate through an application process that compliance with the requirement for being a meaningful EHR user is not possible because the certified EHR technology used by the eligible hospital has been decertified under ONC’s Health IT Certification Program. We are proposing this exception would be available beginning with the FY 2019 payment adjustment year. We considered but are not proposing to make this exception available beginning with the FY 2018 payment adjustment year because making this exception available beginning with the FY 2018

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498 The “list can be found at: https://chpl.healthit.gov/#/decertifications/products.
499 For further descriptions of certification statuses, we refer readers to the CHPL Public User Guide.
500 The “Inactive Certificates” Web page can be found at: https://chpl.healthit.gov/#/decertifications/inactive.
502 https://www.healthit.gov/providers-professionals/ehr-implementation-steps.
payment adjustment would be administratively burdensome, since previous guidance at FAQ 12657 indicated that providers could apply for a hardship if their product was decertified prior to the end of the EHR reporting period.\footnote{https://questions.cms.gov/faq.php?id=5005.} Therefore, we believe that an eligible hospital would have already received a hardship exception under this circumstance. We also note that to date no certifications have been terminated under ONC’s direct review authority. We are proposing an eligible hospital may qualify for this exception if their certified EHR technology was decertified either before or during the applicable EHR reporting period for the payment adjustment year. We refer readers to the definition of “EHR reporting period for a payment adjustment year” under § 495.4 for the applicable EHR reporting periods for payment adjustment years for eligible hospitals. For example, under § 495.4, for the FY 2019 payment adjustment year, the EHR reporting period is any continuous 90-day period in CY 2017. If the certified EHR technology was decertified at any time during the 12-month period preceding the applicable EHR reporting period for the payment adjustment year, or during the applicable EHR reporting period for the payment adjustment year, the eligible hospital may qualify for this exception. For example, if an eligible hospital intended to attest to meaningful use for a 90-day EHR reporting period beginning on April 1, 2017, the eligible hospital could apply for this exception if their certified EHR technology was decertified at any time during the 12-month period beginning on April 1, 2016 and ending on March 31, 2017, or if their certified EHR technology was decertified at any time during their 90-day EHR reporting period beginning on April 1, 2017.

We believe a 12-month period is reasonable for the same reasons stated above for EPs. In addition, we are proposing that the eligible hospital must demonstrate in its application and through supporting documentation if available that the eligible hospital intended to attest to meaningful use for a certain EHR reporting period and made a good faith effort to adopt and implement another CEHRT in advance of that EHR reporting period. We are proposing an eligible hospital seeking to qualify for this exception would submit an application in the form and manner specified by us by July 1 of the year before the payment adjustment year (for example, for the FY 2019 payment adjustment year, by July 1, 2018), or a later date specified by us.

We are proposing to revise § 413.70(a)(6) to add a new category of exception for CAHs that demonstrate through an application process that compliance with the requirement for being a meaningful EHR user is not possible because the certified EHR technology used by the CAH has been decertified under ONC’s Health IT Certification Program. We are proposing this exception would be available beginning with the FY 2018 payment adjustment year. We are proposing a CAH may qualify for this exception if their certified EHR technology was decertified either before or during the applicable EHR reporting period for the payment adjustment year. We refer readers to the definition of “EHR reporting period for a payment adjustment year” under § 495.4 for the applicable EHR reporting periods for payment adjustment years for CAHs. For example, under § 495.4, for the FY 2018 payment adjustment year, the EHR reporting period is either CY 2018 or a continuous 90-day period in CY 2018, depending on whether the CAH has successfully demonstrated meaningful use in a prior year. If the certified EHR technology was decertified at any time during the 12-month period preceding the applicable EHR reporting period for the payment adjustment year, or during the applicable EHR reporting period for the payment adjustment year, the CAH may qualify for this exception. For example, if a CAH intended to attest to meaningful use for a 90-day EHR reporting period beginning on April 1, 2018, the CAH could apply for this exception if their certified EHR technology was decertified at any time during the 12-month period beginning on April 1, 2017 and ending on March 31, 2018, or if their certified EHR technology was decertified at any time during their 90-day EHR reporting period beginning on April 1, 2018. We believe a 12-month period is reasonable for the same reasons stated above for EPs. In addition, we are proposing that the CAH must demonstrate in its application and through supporting documentation if available that the CAH intended to attest to meaningful use for a certain EHR reporting period and made a good faith effort to adopt and implement another CEHRT in advance of that EHR reporting period. We are proposing a CAH seeking to qualify for this exception would submit an application in the form and manner specified by us by November 30 after the end of the applicable payment adjustment year (for example, for the FY 2018 payment adjustment year, by November 30, 2018), or a later date specified by us.

We are inviting public comment on these proposals. We considered alternative timeframes for decertification to the proposed 12-month period preceding the applicable EHR reporting period. We are requesting public comment on whether this 12-month timeframe is reasonable or whether another period should be considered.

3. Ambulatory Surgical Center (ASC)-Based Eligible Professionals (EPs)

Section 16003 of the 21st Century Cures Act amended section 1848(a)(7)(D) of the Act to provide that no payment adjustment may be made under section 1848(a)(7)(A) of the Act for 2017 and 2018 in the case of an eligible professional who furnishes substantially all of his or her covered professional services in an ambulatory surgical center (ASC). Section 1848(a)(7)(D)(iii) of the Act provides that determinations of whether an eligible professional is ASC-based may be made based on the site of service as defined by the Secretary or an attestation, but shall be made without regard to any employment or billing arrangement between the eligible professional and any other supplier or provider of services. Section 1848(a)(7)(D)(iv) of the Act provides that the ASC-based exception shall no longer apply as of the first year that begins more than 3 years after the date on which the Secretary determines, through notice-and-comment rulemaking, that certified EHR technology applicable to the ASC setting is available.

The statute refers to an EP who furnishes "substantially all" of his or her covered professional services in an ASC. Therefore, we must identify the minimum percentage of an EP’s covered professional services that must be furnished in an ASC setting in order for the EP to be considered as furnishing “substantially all” of his or her covered professional services in an ASC. To this end, we are proposing two alternative definitions of an ASC-based EP and requesting public comment to determine the final definition.

We are proposing to define an ASC-based EP under § 495.4 as an EP who furnishes 75 percent or more of his or her covered professional services in sites of service identified by the codes contained in the HIPAA standard transaction as an ASC setting in the calendar year that is two years before the payment.
adjustment year. The percentage of covered professional services in this proposed definition is the same as our definition of a hospital-based MIPS eligible clinician under the Quality Payment Program (§ 414.1305 and 81 FR 77238 through 77240). In the alternative, we are proposing to define an ASC-based EP as an EP who furnishes 90 percent or more of his or her covered professional services in sites of service identified by the codes used in the HIPAA standard transaction as an ASC setting in the calendar year that is two years before the payment adjustment year. The percentage of covered professional services in this alternative proposal is the same as our definition of a hospital-based EP for the EHR Incentive Programs (§495.4 and 75 FR 44439 through 44442). Under these proposals, we would use claims for services furnished in CY 2015 to determine whether an EP is ASC-based for the CY 2017 payment adjustment year, and we would use claims for services furnished in CY 2016 to determine whether an EP is ASC-based for the CY 2018 payment adjustment year. We are also proposing to use Place of Service (POS) code 24 to identify services furnished in an ASC and are requesting public comment on whether other POS codes or mechanisms to identify sites of service should be used in addition to or in lieu of POS code 24.

We analyzed claims data from CYs 2015 and 2016 to estimate how many EPs would be considered ASC-based under our proposal and alternative proposal. Under our proposed definition of “substantially all,” for CY 2015, we found that 380 EPs billed at least 75 percent of their covered professional services in POS 24, out of 523,000 Medicare EPs, which equals approximately .07 percent of Medicare EPs. For CY 2016, we found that 404 EPs billed at least 75 percent of their covered professional services in POS 24, out of 508,575 Medicare EPs, which equals approximately .08 percent of Medicare EPs.

Under our alternative proposed definition of “substantially all,” for CY 2015, we found that 176 EPs billed at least 90 percent of their covered professional services in POS 24, out of 523,000 Medicare EPs, which equals approximately .03 percent of Medicare EPs. For CY 2016, we found that 197 EPs billed at least 90 percent of their covered professional services in POS 24, out of 508,575 Medicare EPs, which equals approximately .04 percent of Medicare EPs.

We are inviting public comment on these proposals.

4. Certification Requirements for 2018

In the 2015 EHR Incentive Program final rule (80 FR 62871 through 62875), we adopted a final policy regarding which Edition of CEHRT must be used by EPs, eligible hospitals, and CAFs for the EHR Incentive Program, which is reflected in the definition of CEHRT §495.4. At a minimum, EPs, eligible hospitals, and CAFs would be required to use EHR technology certified to the 2014 Edition certification criteria for their respective EHR reporting periods in 2015 through 2017. They may also upgrade to the 2015 Edition to meet the required certified EHR technology definition for the EHR reporting periods in 2015, 2016, or 2017, or they may use a combination of 2014 and 2015 Editions if they have modules from both editions that meet the requirements for the meaningful use objectives and measures or if they fully upgrade during an EHR reporting period. Starting with 2018, all EPs, eligible hospitals, and CAFs would be required to use technology certified to the 2015 Edition to demonstrate meaningful use for an EHR reporting period in 2018 and subsequent years (80 FR 62873 through 62875). We received comments on the Stage 3 proposed rule requesting that we allow health care providers to use the 2014 and 2015 Editions of CEHRT in 2018 (80 FR 62874 through 62875). We also received feedback from EPs, eligible hospitals and hospital associations after the 2015 EHR Incentive Program final rule was published. The feedback expressed concerns regarding the burden that will likely occur as a result of the new functionalities required in the implementation of the Stage 3 requirements including an increase in the cost of care without better patient outcomes.

Based on our past experience with the transition from the 2011 Edition to the 2014 Edition and concerns expressed by stakeholders, we understand that transitioning to technology certified to a new Edition can be complex and can require more resources and time than anticipated, including the time necessary to effectively deploy the upgraded system and make the necessary patient safety, staff training and workflow investments. We understand and appreciate these concerns, and are working in cooperation with our Federal partners at ONC to monitor progress on the 2015 Edition upgrade. Furthermore, we believe that there are many benefits for switching to EHR technology certified to the 2015 Edition. At this time, our analysis shows that progress toward certification and upgrade of systems should enable EPs that attest directly to a State for the State’s Medicaid EHR Incentive Program and eligible hospitals and CAHs attesting to CMS or the State’s Medicaid EHR Incentive Program to upgrade systems to the 2015 Edition and successfully attest for an EHR reporting period in 2018.

We will work with ONC to monitor the deployment and implementation status of EHR technology certified to the 2015 Edition. If we identify a change in the current trends and significant issues with the certification and deployment of the 2015 Edition, we will consider flexibility in 2018, for those EPs that attest directly to a State for the State’s Medicaid EHR Incentive Program and eligible hospitals and CAHs attesting to CMS or the State’s Medicaid EHR Incentive Program that are not able to implement 2015 Edition CEHRT to attest for an EHR reporting period in 2018.

One possibility is the flexibility to use technology certified to the 2014 Edition on the 2015 Edition for an EHR reporting period in 2018. Another option is allowing a combination of EHR technologies certified to the 2014 Edition and 2015 Edition to be used for an EHR reporting period in 2018, for those EPs, eligible hospitals, and CAHs that are not able to fully implement EHR technology certified to the 2015 Edition.

We are inviting public comment on these options for offering flexibility in CY 2018 with regard to EHR certification requirements.

X. Proposed Revisions of Medicare Cost Reporting and Provider Requirements

A. Electronic Signature and Submission of the Certification and Settlement Summary Page of the Medicare Cost Report

1. Background

Sections 1815(a) and 1833(e) of the Act provide that no payments will be made to a provider unless it has furnished such information, as may be requested by the Secretary, to determine the amount of payments due the provider under the Medicare program. In general, providers submit this information through annual cost reports that cover a 12-month period of time. Under the provisions of 42 CFR 413.20(b) and 413.24(f), providers are required to submit cost reports annually, with the reporting period based on the provider’s accounting year. For cost reporting periods beginning on or after October 1, 1989, section 1886(f)(1) of the Act and §413.24(f)(4) of the regulations require hospitals to submit cost reports in a standardized electronic format, and the same
requirement was later imposed for other types of providers.

Currently, under §413.24(f)(4)(ii), hospitals, skilled nursing facilities, home health agencies, hospices, end-stage renal disease facilities, organ procurement organizations, histocompatibility laboratories, rural health clinics, Federally qualified health centers, and community mental health centers are required to file Medicare cost reports in a standardized electronic format. When preparing the cost report, the provider’s electronic program must produce the CMS standardized output file in a form that can be read by the contractor’s automated system. This electronic file, also known as the electronic cost report, is forwarded to the contractor for processing through its system. (42 CFR 413.24(f)(4)(ii) and (iii))

Although the Medicare cost report is forwarded to the contractor in electronic format, certain hard copy portions must be separately submitted by the provider to its contractor. Specifically, under §413.24(f)(4)(iv), the provider is required to submit a hard copy of the settlement summary, if applicable, which is a statement of certain worksheet totals, and a certification statement containing a signature by the provider’s administrator or chief financial officer certifying the accuracy of the electronic file. The certification statement and the settlement summary both appear together on the “Certification and Settlement Summary” page of the Medicare cost report for all providers that are required to file a cost report. By signing the certification statement, the provider is certifying, among other things, to the accuracy of the electronic file, and also that it has read the statement that misrepresentation or falsification of information contained in the cost report may be punishable by criminal, civil or administrative action.

This certification statement signed by the provider’s administrator or chief financial officer was incorporated into §413.24(f)(4) of the regulations in a final rule with comment period (59 FR 26964 through 26965) issued in response to public comments received following the Uniform Electronic Cost Reporting System for Hospitals proposed rule (56 FR 41110). Currently, this certification statement is required to have an original signature. This original signature requirement is also set forth in Chapter 1 of the Provider Reimbursement Manual (CMS Pub. 15–2), which explains that a facsimile or stamped copy of the signature is unacceptable. Due to this signature requirement, the Certification and Settlement Summary page containing the original signature is required to be mailed by the provider to the contractor. As set forth in §413.24(f)(4)(iv) and (5)(i) and (ii), an acceptable cost report submission must include the electronic cost report, along with a hard copy of the Certification and Settlement Summary page with an original signature, the Provider Cost Reimbursement Questionnaire, if applicable, and the supporting documentation required from teaching hospitals (the Intern and Resident Information System diskette).

2. Proposed Changes Relating to Electronic Signature on the Certification and Settlement Summary Page of the Medicare Cost Report

In this proposed rule, in lieu of requiring the provider to sign the certification statement with an original signature on a hard copy of the Medicare cost report’s Certification and Settlement Summary page, we are proposing to revise §413.24(f)(4)(iv) to allow providers to use an electronic signature. For Medicare cost reporting purposes, we are proposing that this electronic signature be placed on the signature line of the certification statement and may be (1) any format of the original signature that contains the first and last name of the provider’s administrator or chief financial officer (for example, photocopy or stamp) or (2) an electronic signature that must be the first and last name of the provider’s administrator or chief financial officer entered in the provider’s electronic program. An electronic signature for this purpose cannot be a symbol, numerical characters, or codes. We believe that allowing providers to utilize an electronic signature would afford providers greater flexibility in signing the certification statement and allow a faster and more efficient submission of the Medicare cost report.

To indicate the provider’s election to sign the certification statement with an electronic signature, we are proposing to add an electronic signature checkbox placed immediately after the certification statement and above the signature line on the Certification and Settlement Summary page of the Medicare cost report. The checkbox electing the electronic signature would read: “I have read and agree with the above certification statement. I certify that I intend my electronic signature on this certification statement to be the legally binding equivalent of my original signature.” We are proposing that the checkbox must be checked to signify that the certification statement has been read and that an electronic signature will be placed on the signature line by the provider.

Only when the checkbox is checked would the signature line be accepted with an electronic signature. Completion of both the electronic signature checkbox and the electronic signature, placed on the signature line by the provider’s administrator or chief financial officer under the certification statement, would together constitute an accepted electronic signature of the provider’s administrator or chief financial officer on the certification statement. By signing the certification statement with an electronic signature on the Certification and Settlement Summary page, the signatory would be attesting that its electronic signature was executed with the intent to sign the certification statement, that the electronic signature is being submitted in lieu of an original signature, and additionally that the electronic signature has the same legal effect as an original signature. Because we are proposing that it would be optional for providers to utilize an electronic signature on the certification statement, providers would continue to be able to sign the certification statement with an original signature on a hard copy of the Certification and Settlement Summary page.

We are inviting public comments on our proposals.


In section X.A.2. of the preamble of this proposed rule, we are proposing to allow providers to use an electronic signature on the certification statement of the Certification and Settlement Summary page of the Medicare cost report. We are further proposing that if the provider signs the certification statement with an electronic signature in the manner proposed in section X.A.2. of the preamble of this proposed rule and checks the electronic signature checkbox, the provider also may submit the Certification and Settlement Summary page electronically to the contractor at the same time and in the same manner in which the Medicare cost report is submitted. For example, if the provider submits the electronic cost report file via electronic mail to the contractor, the provider may also include the Certification and Settlement Summary page signed with an electronic signature.

Under our proposal, a provider could sign the certification statement with an original signature on the Certification and Settlement
Summary page. However, if the provider chooses to do so, this page would have to be mailed to its contractor. We believe this proposal, to allow the electronic submission of the Certification and Settlement Summary page, would reduce the need for and storage of paper documents. Under our proposal, providers would have the option to submit the entire cost report electronically, in lieu of the previous requirement to mail a hard copy of the Certification and Settlement Summary page of the Medicare cost report to the contractor. We believe this proposed option would improve the capability of providers to efficiently transmit the Medicare cost report and save providers an appreciable amount of time as well as the cost of separately mailing a hard copy of the Certification and Settlement Summary page of the Medicare cost report to the contractor.

We are inviting public comments on this proposal.

4. Clarifications Relating to the Items Required To Be Submitted by Providers With the Medicare Cost Report

a. Settlement Summary and Certification Statement

In this proposed rule, we are clarifying the portion of the language in §413.24(f)(4)(iv) that describes the items a provider is required to submit along with the electronically filed cost report. Specifically, we are proposing to remove the language in §413.24(f)(4)(iv) which sets forth this expired transition period. Specifically, we are proposing to remove the language that specifies that, during a transition period (first two cost-reporting periods on or after December 31, 2004 for hospices and end-stage renal disease facilities, and the first two cost-reporting periods on or after March 31, 2005 for organ procurement organizations, histocompatibility laboratories, rural health clinics, Federally qualified health centers, and community mental health centers), providers must submit a hard copy of the completed cost report forms in addition to the electronic file. Because the transition period has expired and these providers are no longer required to submit a hard copy of the completed cost report forms in addition to the electronic file, this language in §413.24(f)(4)(iv) is no longer necessary.

b. Removal of the Transition Period Language

Following the effective dates for which certain providers were required to submit cost reports in a standardized electronic format under §413.24(f)(4)(iv), a transition period was implemented when certain providers were required to submit a hard copy of the completed cost report forms in addition to the electronic file. In this proposed rule, we are proposing to remove the language in §413.24(f)(4)(iv) which sets forth this expired transition period. Specifically, we are proposing to remove the language that specifies that, during a transition period (first two cost-reporting periods on or after December 31, 2004 for hospices and end-stage renal disease facilities, and the first two cost-reporting periods on or after March 31, 2005 for organ procurement organizations, histocompatibility laboratories, rural health clinics, Federally qualified health centers, and community mental health centers), providers must submit a hard copy of the completed cost report forms in addition to the electronic file. Because the transition period has expired and these providers are no longer required to submit a hard copy of the completed cost report forms in addition to the electronic file, this language in §413.24(f)(4)(iv) is no longer necessary.

5. Proposed Revisions to 42 CFR 413.24(f)(4)(iv)

In this proposed rule, to reflect our proposals discussed earlier, we are proposing to revise §413.24(f)(4)(iv) to clarify whether any information contained in this cost report may be punishable by criminal, civil, and administrative action, fine and/or imprisonment under Federal law. Furthermore, all caps and informs the provider that “Misrepresentation or falsification of any information contained in this cost report may be punishable by criminal, civil, and administrative action, fines, and/or imprisonment may result.” This language has appeared on the Certification and Settlement Summary page for many years. Because the certification section of the Medicare cost report refers to it as having been read by the provider, incorporation of it into the
regulation text would provide completeness and clarification of the certification statement.

We are inviting public comments on these proposals.

B. Clarification of Limitations on the Valuation of Depreciable Assets Disposed of On or After December 1, 1997

In this section of this proposed rule, we are proposing revisions to the Medicare provider reimbursement policy pertaining to allowable costs and the limits on the valuation of a depreciable asset that may be recognized in establishing an appropriate allowance for depreciation for assets disposed of on or after December 1, 1997. Questions have arisen with regard to whether this limitation on the valuation of depreciable assets depends on the manner in which a provider disposes of an asset. In this proposed rule, we are clarifying that the elimination of the gain or loss for depreciable assets applies to assets a provider disposes of by sale or scrapping on or after December 1, 1997, regardless of whether the asset is scrapped, sold as an individual asset of a Medicare participating provider, or sold incident to a provider’s change of ownership.

Reasonable cost is defined at section 1861(v)(1)(A) of the Act and in the implementing regulations at 42 CFR part 413. Since the inception of the Medicare program, allowable costs under Medicare have included a provider’s direct and indirect costs necessary for the provision of patient care, including the cost of using assets in patient care. Depreciation of these assets is an allowable cost under Medicare and the allowance is computed using the depreciable basis and estimated useful life of the assets (§ 413.134). Under Medicare’s reasonable cost reimbursement system, the appropriate allowance for depreciation and for interest on capital indebtedness on buildings and equipment used in the provision of patient care is based in part on the historical cost of the asset (§ 413.134(a) and (b)). When an asset is disposed of, no further depreciation may be taken on it. Gains and losses on the disposition of depreciable assets may be includable, as applicable, either in computing allowable cost or in computing the adjustment to Medicare reimbursable cost, depending upon the manner of disposition of the asset, the date of scrapping, and the amount of the depreciation adjustment (§ 413.134 and Part 1, Chapter 1 of the Provider Reimbursement Manual (CMS Pub. 15–1)).

Prior to the enactment of the Balanced Budget Act of 1997 (Pub. L. 105–33), when a Medicare certified provider’s capital asset was disposed of through sale or scrapping, Medicare shared in any gain or loss from the transaction. In this regard, if a provider realized a gain or loss from the sale or scrapping of an asset, an adjustment to the provider’s allowable costs was necessary so that Medicare paid its share of the actual cost the provider incurred in using the asset for patient care. Generally, when a provider sold its depreciable assets at more than the net book value, Medicare shared in the gain. If the provider sold its depreciable assets at less than the net book value, Medicare shared in the loss. The amount of a gain was limited to the amount of depreciation previously included in Medicare allowable costs. The amount of a loss was limited to the undepreciated basis of the asset permitted under the program.

In the Balanced Budget Act of 1997, Congress eliminated Medicare’s recognition of gains or losses on a provider’s disposition of assets on or after December 1, 1997. Section 4404 of the Balanced Budget Act of 1997 (Pub. L. 105–33) amended section 1861(v)(1)(O)(i) of the Act to state that, in establishing an appropriate allowance for depreciation and for interest on capital indebtedness with respect to an asset of a provider of services which has undergone a change of ownership, such regulations shall provide, except as provided in clause (iii), that the valuation of the asset after such change of ownership shall be the historical cost of the asset, as recognized under the Medicare program, less depreciation allowed, to the owner of record as of August 5, 1997 (or, in the case of a provider undergoing a change of ownership, such asset permitted under the program. The amount of a loss to be included is limited to the amount of depreciation previously included in Medicare allowable costs. The amount of a gain included in the determination of allowable costs was limited to the amount of depreciation previously included in Medicare allowable costs. The amount of a loss included is limited to the undepreciated basis of the asset permitted under the program. The treatment of the gain or loss depends upon the manner of disposition of the asset, as specified in paragraphs (f)(2) through (6) of [§ 413.134]. The gain or loss on the disposition of depreciable assets has no retroactive effect on a proprietary provider’s equity capital for years prior to the year of disposition.”

In the January 1998 final rule with comment period, we amended by section 4404 of Public Law 105–33. In that rule, we stated that, under the provisions of section 4404 of Public Law 105–33, “when a depreciable asset of a provider undergoes a change of ownership, the valuation of the asset, for purposes of establishing a Medicare allowance for depreciation and interest, will be the historical cost of the asset to the owner of record, less depreciation allowed. Thus, when a depreciable asset is sold, the value of the asset to the seller will be the historical cost (as recognized under Medicare) to the owner of record as of August 5, 1997, less depreciation allowed. In this case, there will be no adjustment for gain or loss on the sale. For the buyer, the value of the asset will also be the historical cost (as recognized under Medicare) to the owner of record as of August 5, 1997, less depreciation allowed. Accordingly, the new owner’s allowance for depreciation and interest will be based on this value. Stated simply, the asset moves from the hands of the seller to the hands of the buyer at the asset’s net book value defined in § 413.134(b)(9) (63 FR 1381).

Our policy referenced the asset of a provider undergoing a change of ownership, meaning the asset itself changing owners, regardless of whether the provider changes ownership. In conforming the regulations to the new statutory provision, we revised the regulations at § 413.134(f)(1) to specify that “[d]epreciable assets may be disposed of through sale, scrapping, trade-in, exchange, demolition, abandonment, condemnation, fire, theft, or other casualty. If disposal of a depreciable asset, including the sale or scrapping of an asset before December 1, 1997, results in a gain or loss, an adjustment is necessary in the provider’s allowable cost. (No gain or loss is recognized on either the sale or the scrapping of an asset that occurs on or after December 1, 1997.) The amount of a gain included in the determination of allowable cost is limited to the amount of depreciation previously included in Medicare allowable costs. The amount of a loss included is limited to the undepreciated basis of the asset permitted under the program. The treatment of the gain or loss depends upon the manner of disposition of the asset, as specified in paragraphs (f)(2) through (6) of [§ 413.134]. The gain or loss on the disposition of depreciable assets has no retroactive effect on a proprietary provider’s equity capital for years prior to the year of disposition.”
scraping of an asset that occurs on or after December 1, 1997) to § 413.134(f)(1). This parenthetical was intended to implement section 4404 of the BBA of 1997 by disallowing the gain or loss when a provider sells or scraps an asset.

We believe that, under section 4404 of the BBA of 1997, Medicare’s nonrecognition of a loss or gain with respect to an asset a provider disposes of by sale or scrapping applies, regardless of whether the sale of the asset occurs incident to a provider’s change of ownership or whether the asset is otherwise sold or scrapped by a currently participating Medicare provider.

We note that following the enactment of the Deficit Reduction Act of 1984 (Pub. L. 98–369, section 2314), in which Congress amended section 1861(v)(1) of the Act by adding new subparagraph (O) concerning the valuation and determination of historical costs of assets after July 18, 1984, we stated that the new provisions applied “not only to the sale or purchase of groups of assets, but also to the sale or purchase of individual assets” (57 FR 43913).

Similarly, we believe section 4404 of the BBA of 1997 applies to a provider’s disposition of assets through sale or scrapping, including the sale or scrapping of individual provider assets and assets sold or scrapped incident to a provider’s change of ownership. Accordingly, we are proposing to revise the regulation text at § 413.134(f)(1) to clarify our longstanding policy that Medicare does not recognize a provider’s gain or loss on the sale or scrapping of an asset that occurs on or after December 1, 1997, regardless of whether the asset is sold incident to a provider’s change of ownership or is otherwise sold or scrapped as an asset of a Medicare participating provider.

XL Proposed Changes Relating to Survey and Certification Requirements

A. Proposed Revisions to the Application and Re-Application Procedures for National Accrediting Organizations. Provider and Supplier Conditions, and Posting of Survey Reports and Acceptable Plans of Corrections (PoCs)

1. Background

Health care facilities must demonstrate compliance with the Medicare conditions of participation (CoPs), conditions for coverage (CfCs), or conditions for certification (depending on the type of facility) to be eligible to receive Medicare payments. Section 1865 of the Act allows health care facilities that are “provider entities” to demonstrate this compliance through accreditation by an accreditation program of a private, national accrediting organization (AO) that is approved by the Secretary. An AO must demonstrate the ability to effectively evaluate a facility’s compliance using accreditation standards that meet or exceed the applicable Medicare conditions, as well as survey processes that are comparable to those survey methods, procedures, and forms required by CMS for conducting Federal surveys for the same health care facility type, which are generally outlined in regulations and specified in the State Operations Manual (SOM).

Section 1865(a)(2) of the Act requires that the Secretary base its decision to approve or deny the Medicare accreditation program application of an accrediting organization after considering at least the following factors: (a) Program requirements for the accreditation program to meet or exceed Medicare requirements; (b) survey procedures that are comparable to those of Medicare; (c) the ability to provide adequate resources for conducting surveys; (d) the capacity to furnish information for use by CMS in enforcement activities; (e) monitoring procedures for providers or suppliers identified as being out of compliance with conditions or requirements; and (f) the ability to provide the necessary data for validation surveys to the Secretary. In addition, section 1865(a)(2) of the Act specifies that the Secretary shall consider other factors with respect to determining the AOs ability to meet or exceed applicable conditions, therefore meaning that CMS has the ability to determine “other factors” when considering an AO for deemed status. CMS has responsibility for oversight and approval of AO accreditation programs used for Medicare certification purposes, and for ensuring that providers and suppliers that are accredited under an approved AO accreditation program meet the quality and patient safety standards required by the Medicare conditions and requirements. The Medicare regulations at 42 CFR 488.5 set forth the detailed requirements that a national AO must satisfy in order to receive approval, and maintain recognition, of a Medicare accreditation program. Section 488.5 also details the procedures that CMS follows in reviewing applications from AOs.

The results of surveys conducted by State Survey Agencies of a facility’s compliance with Medicare conditions and requirements of CMS-certified facilities are reported using the CMS Form 2567, “Statement of Deficiencies and Plan of Correction” (OMB No. 0938–0391). These reports describe any findings of noncompliance with Federal requirements (also referred to as “deficiencies”) that the surveyors may have found. If there are cited deficiencies, a facility must submit an acceptable plan of correction (PoC) for achieving compliance to CMS describing how and when, within a reasonable timeframe, it will correct them. Failure to correct deficiencies will lead to the facility’s termination from Medicare participation.

CMS makes survey reports and acceptable PoCs publicly available through a variety of settings as part of the Department’s commitment to transparency, and to providing all health care consumers and the general public with access to quality and safety information. CMS began posting redacted CMS Form 2567 survey data for skilled nursing facilities and nursing facilities on its Nursing Home Compare Web site in July 2012. In March 2013, CMS began posting on its Web site the CMS Form-2567 surveys reports based on complaint investigations for short-term acute care hospitals and critical access hospitals (CAHs). In addition, two Web sites owned by private entities also publish the public CMS survey data of nursing homes, short-term acute care hospitals, and CAHs, based on the CMS survey information. The ProPublica Web site and the Association for Health Care Journalist (AHJC) Web sites, respectively, provide search engines that refer back to the CMS Form-2567 data that CMS has made available. These Web sites enable all health care consumers and the general public across the country to learn about the performance of these providers in order to make more informed decisions about where to get health care. We also believe that release of this information encourages these health care providers to improve the quality of care and services they provide. Such information can also be obtained by the public directly from State Survey Agencies. AOs perform their own accreditation surveys and issue their own survey reports which provide information on accredited facilities’ compliance with Federal standards. These facilities include: Hospitals, psychiatric hospitals, CAHs, home health agencies (HHAs), hospices, ambulatory surgery


centers (ASCs), outpatient physical therapy and speech-language pathology services (OPTs), and rural health clinics (RHCs). These facilities participate in Medicare based on their accreditation from a CMS-approved AO and are not subject to routine surveys from State survey agencies. By contrast, AOs currently do not make their survey reports and accompanying PoCs publicly available. We believe it is important to continue to lead the effort to make information regarding a health care facility’s compliance with health and safety requirements found in survey reports publicly available through our various provider and supplier Compare sites, including hospital and home health Compare sites to increase transparency. CMS recognizes, based on the above references to CMS Compare sites and other resources which make survey reports publicly available, that these survey reports vary in the type of information accessible to the public (complaint) based on the provider or supplier type. For example, the current CMS Survey and Certification site for hospital 2567 downloads (https://www.cms.gov/Medicare/Provider-Enrollment-and-Certification/CertificationandCompliance/Hospitals.html) only contains complaint surveys; no recertification survey reports are posted. In addition, there has been an increasing concern in terms of AO disparity rates based on the AO deficiency findings compared to serious, condition-level deficiencies found by the State Survey Agencies. For example, in FY 2015, the disparity rates increased by 1 percent to 39 percent for hospitals and decreased by 6 percent to 69 percent for psychiatric hospitals, from FY 2014. This continued trend of high disparity rates from FY 2012 to FY 2015 raises serious concerns regarding the AOs’ ability to appropriately identify and cite health and safety deficiencies during the survey process. Therefore, we believe that posting AO survey reports and acceptable PoCs would address some of the concerns of reporting hospital information from both CMS and AOs, as well as the disparity between serious deficiency findings, and provide a more comprehensive picture to health care consumers and the public in general.

As the number of health care facilities participating in Medicare by virtue of their accreditation and deemed status increases, the number of survey reports and acceptable PoC available to health care consumers decreases. The table below illustrates that 40 percent of Medicare-participating providers or suppliers with an accreditation option participate in Medicare via accreditation and deemed status. In addition, 89 percent of hospitals and psychiatric hospitals across the country participate in Medicare via accreditation and deemed status. This represents a significant number of hospital and other health care facility survey reports and acceptable PoCs that are currently not available to health care consumers. This information is not available to assist health care consumers in their decision making when selecting a health care facility in which to receive care for themselves or a loved one. Therefore, we believe that it is critical that accrediting organizations with CMS-approved accreditation programs make available publicly all survey reports and acceptable plans of correction on their Web sites.

### TOTAL MEDICARE PARTICIPATING FACILITIES—FY 2015 DEEMED VERSUS NON-DEEMED

<table>
<thead>
<tr>
<th>Program type</th>
<th>Deemed* (percentage)</th>
<th>Non-deemed** (percentage)</th>
<th>Total</th>
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<tr>
<td></td>
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</tr>
<tr>
<td>Hospital</td>
<td>3,500 (89)</td>
<td>432 (11)</td>
<td>3,932</td>
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<tr>
<td>Psychiatric Hospital</td>
<td>424 (89)</td>
<td>53 (11)</td>
<td>477</td>
</tr>
<tr>
<td>CAH</td>
<td>420 (32)</td>
<td>887 (68)</td>
<td>1,307</td>
</tr>
<tr>
<td>HHA</td>
<td>4,450 (47)</td>
<td>5,008 (53)</td>
<td>9,458</td>
</tr>
<tr>
<td>Hospice</td>
<td>1,694 (40)</td>
<td>2,573 (60)</td>
<td>4,267</td>
</tr>
<tr>
<td>ASC</td>
<td>1,499 (27)</td>
<td>3,973 (73)</td>
<td>5,472</td>
</tr>
<tr>
<td>OPT</td>
<td>175 (8)</td>
<td>1,957 (92)</td>
<td>2,132</td>
</tr>
<tr>
<td>RHC</td>
<td>253 (6)</td>
<td>3,862 (94)</td>
<td>4,115</td>
</tr>
<tr>
<td>Total</td>
<td>12,415 (40)</td>
<td>18,745 (60)</td>
<td>31,160</td>
</tr>
</tbody>
</table>

* As reported by accrediting organizations.
** Surveyed by a State survey agency for compliance with Medicare conditions.

### 2. Proposed Regulation Changes

In an effort to increase transparency, in this proposed rule, we are proposing to require AOs with CMS-approved accreditation programs to post final accreditation survey reports and acceptable PoCs on public facing Web site designated by the AO. All current AOs with CMS-approved accreditation programs have Web sites that inform the general public about their organization. Therefore, we are proposing to require AOs to have their final accreditation survey reports and acceptable PoCs available on their Web sites.

Establishing the standard for posting both accredited and nonaccredited provider and supplier survey reports, which would include initial and recertification surveys, and acceptable PoCs would expand transparency even further. Disclosure of survey findings protects both patient health and safety, in which public disclosure of findings currently only shows the subset of complaint activity. Expanding these requirements through the posting of all survey reports and acceptable PoCs would allow for a more comprehensive way to show a provider’s or supplier’s compliance with all health and safety requirements.

Therefore, we are proposing to revise §488.5 of the regulations to incorporate this proposed requirement. We are proposing to add a new standard at §488.5(a)(21) to require that each national AO applying or reapplying for CMS-approval of its Medicare provider or supplier accreditation program provide a statement acknowledging that it agrees to make all Medicare provider or supplier final accreditation survey reports (including statements of deficiency findings) as well as acceptable PoCs publicly available on their Web sites.

Note that other types of facilities may also participate in Medicare via an approved accreditation program, but to date, no AO has sought and received approval for any of these additional facility types.
its Web site within 90 days after such information is made available to those facilities for the most recent 3 years. This provision would include all triennial, full, follow-up, focused, and complaint surveys, whether they are performed onsite or offsite.

In addition, pursuant to section 1834(e) of the Act, State Survey Agencies do not evaluate suppliers of the technical component of advanced diagnostic imaging services. CMS-approved advanced diagnostic imaging AOs are the only source of compliance data for suppliers of the technical component of advanced diagnostic imaging services. Therefore, we believe it is critical that these AOs also be required to post survey reports and acceptable PoCs on their Web sites. Otherwise, it will not be possible to provide health care consumers with compliance information about Medicare-participating suppliers of advanced diagnostic imaging services.

We are proposing to amend our regulations at 42 CFR 414.68 governing imaging accreditation under Medicare by redesignating paragraphs (c)(7) through (c)(14) as paragraphs (c)(6) through (c)(15), respectively, and adding a new paragraph (c)(7) to require that each national advanced diagnostic imaging AO that applies or reapplies for CMS approval of its Medicare advanced diagnostic imaging accreditation program must provide a statement acknowledging that it agrees to make all Medicare advanced diagnostic imaging final accreditation survey reports as well as acceptable PoCs publicly available on its Web site within 90 days after such information is made available to the supplier of advanced diagnostic imaging services for the most recent 3 years. This provision would apply to all full, follow-up, focused, and complaint surveys, regardless of whether they are performed onsite or offsite.

We are inviting public comments on these proposals.

B. Proposed Changes to Termination Public Notice Requirements for Certain Providers and Suppliers

1. Background

Under the provisions of sections 1866(b)(2) of the Act and implementing regulations at 42 CFR 489.53, the Secretary may terminate an agreement with a provider of services if it is determined that the provider is not in substantial compliance with applicable requirements governing provider agreements. For instance, CMS must determine that the provider:

- Is not complying substantially with the terms of the agreement, the provisions of title XVIII, or regulations promulgated thereunder;
- Has failed to supply information necessary to determine whether payments are or were due and the amounts of such payments;
- Refuses to permit examination of fiscal and other records (including medical records) necessary for the verification of information furnished as a basis for claiming payment under the Medicare program; or
- Refuses to permit photocopying of any records or other information necessary to determine or verify compliance with participation requirements.

Sections 1866(b)(1) and (2) of the Act require reasonable public notice, as prescribed in regulations, of both voluntary and involuntary terminations of Medicare and Medicaid participating providers and suppliers. Various existing regulations specify the requirements of public notice for voluntary and involuntary terminations prior to termination of a provider or supplier agreement. Specifically, for voluntary terminations, providers at 42 CFR 405.2404(d), ASCs at 42 CFR 405.2442, ASCs at 42 CFR 416.35(d), and OPOs at 42 CFR 486.312(e) are required to publish termination notices in the local public newspaper.

2. Basis for Proposed Changes

The existing regulations requiring termination notices to be published in local newspapers have been outdated over time as the public and beneficiaries increasingly turn to the Internet and other electronic forums for information. Currently, rural health centers (RHCs), Federally qualified health centers (FQHCs), ambulatory surgical centers (ASCs), and organ procurement organizations (OPOs) are required to publish public notices of voluntary and involuntary termination of participation in the Medicare and Medicaid programs in one or more local newspapers. Providers and suppliers that voluntarily terminate their participation agreement must give notice to the public at least 15 days before the effective date of termination and the notice must be published in one or more local newspapers. The use of hard copy local newspaper through time has become less effective, as a large majority of the public uses alternate sources such as Web sites or other online news and resources.

According to national studies, approximately 23 percent of the general public continues to read print newspapers. Many individuals have turned to digital platforms to read news rather than print news, which continues to decline on an annual basis, therefore, limiting the effectiveness of publishing termination notices in local newspapers. In light of the public’s increased access to the Internet and other electronic forums for information and the decline of print newspaper readership, in this proposed rule, we are proposing changes in the existing regulations noted earlier regarding newspaper publication of termination notices to allow CMS Regional Offices and providers and suppliers more media platforms in which to publish termination notices, both voluntary and involuntary, with the intent of making these notices more visible and effective.

3. Proposed Changes to Regulations

In this proposed rule, we are proposing to remove the regulatory language specifying public notice of terminations for FQHCs, RHCs, ASCs, and OPOs to be exclusively in newspapers to allow for more flexibility for both the CMS Regional Offices and providers and suppliers. Specifically, we are proposing changes to the regulations for RHCs at 42 CFR 405.2404(d), for FQHCs at 42 CFR 405.2442(a) and (b), for ASCs at 42 CFR 416.35(d), and for OPOs at 42 CFR 486.312(e) to remove the reference to publication in newspapers as the means for notifying the community of involuntary and voluntary terminations from participation in Medicare and Medicaid programs. This proposal for termination notices to the public for RHCs, FQHCs, ASCs, and OPOs would align with the termination notices CMS currently has set forth for all other providers and suppliers. For example, under 42 CFR 405.2404(d) (enforcement procedures for long-term care facilities), CMS must notify the public of a termination of a nursing home's provider agreement, but the regulation does not specify through which public forum this notice is to be given. Similarly, 42 CFR 489.53(d)(5) also does not specify the method of public notification required for terminations. Through this proposed change, RHCs, FQHCs, ASCs, and OPOs would have the same requirement for the notice to the public as under 42 CFR 405.2404(d)(5), where there is a termination by CMS in which public notice is required but the method for these providers or suppliers
for providing public notice is not specified, to allow for flexibility.

In addition, we are proposing to revise 42 CFR 489.52(c)(2) to remove the requirement to publish notice in one or more local newspapers in circumstances of the termination of a provider agreement by a provider and instead to allow providers to inform the community via public notice, without specifying the method used for public notice. We believe that these proposed changes will ensure that the community continues to be aware of terminations of Medicare and Medicaid participating providers and suppliers.

The method for delivering the required public notice is no longer being specified by removing the word “newspaper” from the regulations for RHCS, FQHCs, ASCs, and OPOs. Instead, we are proposing to allow for flexibility for the CMS Regional Offices and the providers or suppliers to post public notices through a manner in which the maximum number of community individuals and beneficiaries would be informed. This may include, but is not limited to State Web site postings, facility Web sites, or local news and social media channels. It also would not preclude publication in local newspapers. Through this proposed rule, we will continue to fulfill the regulatory requirement to publically post involuntary termination notices. We are also operationally considering allowing voluntarily terminating providers and suppliers the same public notice platform used for involuntary notices in order to meet their regulatory public notice requirements. This could include media venues such as Web site postings and press releases through the use of CMS Regional press officers.

We are inviting public comments on our proposals. In addition, we are seeking suggestions from the public on sufficient mechanisms to provide public information, other than local newspapers, for posting Medicare and Medicaid participating provider and supplier termination notices.

XII. MedPAC Recommendations

Under section 1886(e)(4)(B) of the Act, the Secretary must consider MedPAC’s recommendations regarding hospital inpatient payments. Under section 1886(e)(5) of the Act, the Secretary must publish in the annual proposed and final IPPS rules the Secretary’s recommendations regarding MedPAC’s recommendations. We have reviewed MedPAC’s March 2017 “Report to Congress: Medicare Payment Policy” and have given the recommendations in the report consideration in conjunction with the proposed policies set forth in this proposed rule. MedPAC recommendations for the IPPS for FY 2018 are addressed in Appendix B to this proposed rule.

For further information relating specifically to the MedPAC reports or to obtain a copy of the reports, contact MedPAC at (202) 653–7226, or visit MedPAC’s Web site at: http://www.medpac.gov.

XIII. Other Required Information

A. Publicly Available Data

IPPS-related data are available on the Internet for public use. The data can be found on the CMS Web site at: http://www.cms.hhs.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html. Following is a listing of the IPPS-related files that are available.

1. CMS Wage Data Public Use File

This file contains the hospital hours and salaries from Worksheet S–3, Parts II and III from FY 2014 Medicare cost reports used to create the proposed FY 2018 IPPS wage index. Multiple versions of this file are created each year. For a complete schedule on the release of different versions of this file, we refer readers to the wage index schedule in section III.M. of the preamble of this proposed rule.

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<thead>
<tr>
<th>Processing year</th>
<th>Wage data year</th>
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Media: Internet at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Wage-Index-Files.html.


2. CMS Occupational Mix Data Public Use File

This file contains the CY 2013 occupational mix survey data to be used to compute the occupational mix adjustment wage indexes. Multiple versions of this file are created each year. For a complete schedule on the release of different versions of this file, we refer readers to the wage index schedule in section III.M. of the preamble of this proposed rule.

Media: Internet at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Wage-Index-Files.html.


3. Provider Occupational Mix Adjustment Factors for Each Occupational Category Public Use File

This file contains each hospital’s occupational mix adjustment factors by occupational category. Two versions of these files are created each year to support the rulemaking.

Media: Internet at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Wage-Index-Files.html.


4. Other Wage Index Files

CMS releases other wage index analysis files after each proposed and final rule.

Media: Internet at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Wage-Index-Files.html.


5. FY 2018 IPPS SSA/FIPS CBSA State and County Crosswalk

This file contains a crosswalk of State and county codes used by the Social Security Administration (SSA) and the Federal Information Processing Standards (FIPS), county name, and a list of Core-Based Statistical Areas (CBSAs).

Media: Internet at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Acute-Inpatient-Files-for-Download.html.


6. HCRIS Cost Report Data

The data included in this file contain cost reports with fiscal years ending on or after September 30, 1996. These data files contain the highest level of cost report status.


(We note that data are no longer offered on a CD. All of the data collected are now available free for download from the cited Web site.)
7. Provider-Specific File

This file is a component of the PRICER program used in the MAC’s system to compute DRG/MS–DRG payments for individual bills. The file contains records for all prospective payment system eligible hospitals, including hospitals in waiver States, and data elements used in the prospective payment system recalibration processes and related activities. Beginning with December 1988, the individual records were enlarged to include pass-through per diems and other elements.

Media: Internet at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/ProspectivePayment/ProspectivePaymentSystem/Standardizing%20File/Standardizing%20File.htm.
Period Available: Quarterly Update.

8. CMS Medicare Case-Mix Index File

This file contains the Medicare case-mix index by provider number as published in each year’s update of the Medicare hospital inpatient prospective payment system. The case-mix index is a measure of the costliness of cases treated by a hospital relative to the cost of the national average of all Medicare hospital cases, using DRG/MS–DRG weights as a measure of relative costliness of cases. Two versions of this file are created each year to support the rulemaking.

Media: Internet at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Acute-Inpatient-Files-for-Download.html.

9. MS–DRG Relative Weights (Also Table 5—MS–DRGs)

This file contains a listing of MS–DRGs, MS–DRG narrative descriptions, relative weights, and geometric and arithmetic mean lengths of stay for each fiscal year. Two versions of this file are created each year to support the rulemaking.

Media: Internet at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Acute-Inpatient-Files-for-Download.html.

10. IPPS Payment Impact File

This file contains data used to estimate payments under Medicare’s hospital inpatient prospective payment systems for operating and capital-related costs. The data are taken from various sources, including the Provider-Specific File, HCRIS Cost Report Data, MedPAR Limited Data Sets, and prior impact files. The data set is abstracted from an internal file used for the impact analysis of the changes to the prospective payment systems published in the Federal Register. Two versions of this file are created each year to support the rulemaking.

Media: Internet at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Historical-Impact-Files-for-FY–1994-through-Present.html.

11. AOR/BOR Tables

This file contains data used to develop the MS–DRG relative weights. It contains mean, maximum, minimum, standard deviation, and coefficient of variation statistics by MS–DRG for length of stay and standardized charges. The BOR tables are “Before Outliers Removed” and the AOR is “After Outliers Removed.” (Outliers refer to statistical outliers, not payment outliers.) Two versions of this file are created each year to support the rulemaking.

Media: Internet at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Acute-Inpatient-Files-for-Download.html.

12. Prospective Payment System (PPS) Standardizing File

This file contains information that standardizes the charges used to calculate relative weights to determine payments under the hospital inpatient operating and capital prospective payment systems. Variables include wage index, cost-of-living adjustment (COLA), case-mix index, indirect medical education (IME) adjustment, disproportionate share, and the Core-Based Statistical Area (CBSA). The file supports the rulemaking.

Media: Internet at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Acute-Inpatient-Files-for-Download.html.

13. Hospital Readmissions Reduction Program Supplemental File

This file contains information on the calculation of the Hospital Readmissions Reduction Program (HRRP) payment adjustment. Variables include the proxy excess readmission ratios for acute myocardial infarction (AMI), pneumonia (PN) and heart failure (HF), coronary obstruction pulmonary disease (COPD), total hip arthroplasty (THA)/total knee arthroplasty (TKA), and coronary artery bypass grafting (CABG) and the proxy readmissions payment adjustment for each provider included in the program. In addition, the file contains information on the number of cases for each of the applicable conditions excluded in the calculation of the readmission payment adjustment factors. It also contains MS–DRG relative weight information to estimate the payment adjustment factors. The file supports the rulemaking.

Media: Internet at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Acute-Inpatient-Files-for-Download.html.

14. Medicare Disproportionate Share Hospital (DSH) Supplemental File

This file contains information on the calculation of the uncompensated care payments for FY 2018. Variables include the data used to determine a hospital’s share of uncompensated care payments, total uncompensated care payments and estimated per claim uncompensated care payment amounts. The file supports the rulemaking.

Media: Internet at: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/Acute-Inpatient-Files-for-Download.html.

B. Collection of Information Requirements

1. Statutory Requirement for Solicitation of Comments

Under the Paperwork Reduction Act of 1995, we are required to provide 60-day notice in the Federal Register and solicit public comment before a collection of information requirement is submitted to the Office of Management and Budget (OMB) for review and approval. In order to fairly evaluate whether an information collection should be approved by OMB, section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995 requires that we solicit comment on the following issues:

- The need for the information collection and its usefulness in carrying out the proper functions of our agency.
- The accuracy of our estimate of the information collection burden.
- The quality, utility, and clarity of the information to be collected.

Recommendations to minimize the information collection burden on the
affected public, including automated collection techniques.

In this proposed rule, we are soliciting public comment on each of these issues for the following sections of this document that contain information collection requirements (ICRs).

2. ICRs for Add-On Payments for New Services and Technologies

Section II.H.1. of the preamble of this proposed rule discusses add-on payments for new services and technologies. Specifically, this section states that applicants for add-on payments for new medical services or technologies for FY 2019 must submit a formal request. A formal request includes a full description of the clinical applications of the medical service or technology and the results of any clinical evaluations demonstrating that the new medical service or technology represents a substantial clinical improvement. In addition, the request must contain a significant sample of the data to demonstrate that the medical service or technology meets the high-cost threshold.

We believe the burden associated with this requirement is exempt from the PRA under 5 CFR 1320.3(c), which defines the agency collection of information subject to the requirements of the PRA as information collection imposed on 10 or more persons within any 12-month period. This information collection does not impact 10 or more entities in a 12-month period. For FY’s 2008, 2009, 2010, 2011, 2012, 2013, 2014, 2015, 2016, and 2017, we received 1, 4, 5, 3, 3, 5, 7, 9, 9, and 9 applications, respectively. We note that 3 of the 9 applications for FY 2018 were withdrawn prior to the publication of the proposed rule as indicated in section II.H.6. of the preamble of this proposed rule.

3. ICRs for the Occupational Mix Adjustment to the Proposed FY 2018 Wage Index (Hospital Wage Index Occupational Mix Survey)

Section II.E. of the preamble of this proposed rule discusses the occupational mix adjustment to the proposed FY 2018 wage index. While the preamble does not contain any new ICRs, we note that there is an OMB approved information collection request associated with the hospital wage index.

Section 304(c) of Public Law 106–554 amended section 1886(d)(3)(E) of the Act to require us to collect data at least once every 3 years on the occupational mix of employees for each short-term, acute care hospital participating in the Medicare program in order to construct an occupational mix adjustment to the wage index. We collect the data via the occupational mix survey.

The burden associated with this information collection requirement is the time and effort required to collect and submit the data in the Hospital Wage Index Occupational Mix Survey to CMS. The aforementioned burden is subject to the PRA; it is currently approved under OMB control number 0938–0907.

4. Hospital Applications for Geographic Reclassifications by the MGCRB

Section III.I.2. of the preamble of this proposed rule discusses proposed changes to the wage index based on hospital reclassifications. As stated in that section, under section 1886(d)(10) of the Act, the MGCRB has the authority to accept short-term IPPS hospital applications requesting geographic reclassification for wage index and to issue decisions on these requests by hospitals for geographic reclassification for purposes under the IPPS.

The burden associated with this application process is the time and effort necessary for an IPPS hospital to complete and submit an application for reclassification to the MGCRB. The burden associated with this requirement is subject to the PRA. The current information collection requirement for this application process is approved under OMB Control Number 0938–0573 in 2014 but expired on February 28, 2017. A request for an extension of this currently approved collection requirement under OMB control number 0938–0573 is currently awaiting OMB approval and can be accessed at: https://www.reginfo.gov/public/do/PRAViewICR?ref_nbr=201612-0938-023.

5. ICRs for Temporary Exception to the LTCH PPS Site Neutral Payment Rate for Certain Spinal Cord Specialty Hospitals

In section VIII.E. of the preamble of this proposed rule, we discuss the proposed implementation of section 15009 of Public Law 114–255, which provides for a temporary exception to the site neutral payment rate for certain spinal cord specialty hospitals under section 1886(m)(6)(F)(iii) of the Act. Under this provision, discharges occurring in cost reporting periods beginning during FY 2018 and FY 2019 for LTCHs that meet the specified statutory criteria are excluded from the site neutral payment rate (that is, all discharges from such LTCHs during this period would be paid at the LTCH PPS standard Federal payment rate). In order for an LTCH to qualify for this temporary exception, the LTCH must, among other things, meet the “significant out-of-state admissions criterion” at section 1886(m)(6)(F)(iii) of the Act. To meet the significant out-of-state admissions criterion, an LTCH must have discharged inpatients (including both individuals entitled to, or enrolled for, Medicare Part A benefits and individuals not so entitled or enrolled) during FY 2014 who had been admitted from at least 20 of the 50 States, determined by the States of residence of such inpatients and based on such data submitted by the hospital to the Secretary as the Secretary may require. The statute further provides authority for the Secretary to implement the significant out-of-state admissions criterion at section 1886(m)(6)(F)(iii) of the Act by program instruction or otherwise, and exempts the policy initiatives from any information collection requirements under the Paperwork Reduction Act. As such, the burden associated with the data submitted by the hospital to meet the significant out-of-State admissions criteria is not subject the PRA. However, our estimate of the burden associated with this data submission is discussed in section J.f. of Appendix A of this proposed rule.

6. ICRs for the Hospital Inpatient Quality Reporting (IQR) Program

a. Background

The Hospital IQR Program (formerly referred to as the Reporting Hospital Quality Data for Annual Payment (RHQDAPU) Program) was originally established to implement section 501(b) of the MMA, Public Law 108–173. This program expanded our voluntary Hospital Quality Initiative. The Hospital IQR Program originally consisted of a “starter set” of 10 quality measures. The collection of information associated with the original starter set of quality measures was previously approved under OMB control number 0938–0918.

All of the information collection requirements previously approved under OMB control number 0938–0918 have been combined with the information collection request currently approved under OMB control number 0938–1022. We no longer use OMB control number 0938–0918. OMB has currently approved 3,681,023 hours of burden and approximately $121 million for purposes of payment under the IPPS. OMB has currently approved collection techniques.

In section IX.A. of the preamble of this proposed rule, we are making the following proposals that we expect to
affect our burden estimates: (1) Updates to the electronic clinical quality measure (eCQM) reporting requirements with regard to the number of eCQMs and quarters of data for the FY 2019 and FY 2020 payment determinations; (2) updates to our previously finalized eCQM validation procedures for the FY 2020 payment determination and subsequent years; and; (3) begin voluntary reporting on the new Hybrid Hospital-Wide 30-Day Readmission measure for the CY 2018 reporting period. Details on these proposals, as well as the expected burden changes, are discussed below.

This proposed rule also includes proposals to: (1) Update the eCQM certification requirements for the FY 2019 and FY 2020 payment determinations; (2) allow hospitals to use an educational review process to correct incorrect validation results for the first three quarters of validation for chart-abstracted measures beginning with the FY 2020 payment determination and for subsequent years; (3) refine the Hospital Consumer Assessment of Healthcare Providers and Systems (HCAHPS) Survey measure (NQF #0166 and 0228) to replace the questions on pain management for the FY 2020 payment determination and subsequent years; (4) refine the Hospital 30-Day, All-Cause, Risk-Standardized Mortality Rate Following Acute Ischemic Stroke Measure to include the National Institutes of Health (NIH) Stroke Scale data for the FY 2023 payment determination and subsequent years; (5) provide confidential reports of measure data stratified by dual eligible status for the Hospital 30-day, All-Cause, Risk-Standardized Readmission Rate Following Pneumonia Hospitalization and Hospital 30-day, All-Cause, Risk Standardized Mortality Rate (RSRR) for Pneumonia measures; and; (6) align the naming of the Extraordinary Circumstances Exceptions (ECE) Policy for the FY 2020 payment determination and subsequent years. As discussed further below, we do not expect these proposals to affect our burden estimates.

In the FY 2017 IPPS/LTCH PPS final rule, we finalized policies to require hospitals to submit a full year (four quarters) (81 FR 57159) of data for at least eight eCQMs of the available eCQMs (81 FR 57157) for both the FY 2019 and FY 2020 payment determinations. In section IX.A.8. of the preamble of this proposed rule, we are proposing the following changes to these finalized policies: (1) Revise the CY 2017 reporting period/FY 2019 payment determination eCQM reporting requirements, such that hospitals are required to report six eCQMs and to submit two, self-selected, calendar quarters of data; and (2) revise the CY 2018 reporting period/FY 2020 payment determination eCQM reporting requirements such that hospitals are required to report six eCQMs for the first three quarters of CY 2018.

(1) Calculations for the CY 2017 Reporting Period/FY 2019 Payment Determination

As in previous years, we believe the total burden associated with the eCQM reporting policy will be similar to that previously outlined in the Medicare EHR Incentive Program Stage 2 final rule (77 FR 54126 through 54133). Under that program, the burden estimate for a hospital to report one eCQM is 10 minutes per record per quarter. We believe this estimate is accurate and appropriate to apply to the Hospital IQR Program because we align the eCQM reporting requirements between both programs. Therefore, using the estimate of 10 minutes per record per quarter, we anticipate our proposal to require: (1) Reporting on at least six of the available eCQMs; and (2) submission of two self-selected quarters of eCQM data, would result in a burden reduction of 3 hours and 20 minutes (200 minutes) per hospital for the FY 2019 payment determination. This estimate was calculated by considering the burden difference between the updated eCQM reporting requirements proposed for the FY 2019 payment determination (10 minutes per record ¥ 6 eCQMs ¥ 2 quarters = 120 minutes for 2 quarters of reporting) and the eCQM reporting requirements previously finalized in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57157 through 57159) (10 minutes per record ¥ 8 eCQMs ¥ 4 quarters = 320 minutes for 4 quarters of reporting). Through these calculations (120 minutes-320 minutes), we arrived at a reduction of 200 minutes per hospital per year, or 3 hours and 20 minutes per hospital per year, for the FY 2019 payment determination.

In total, for the FY 2019 payment determination, we expect our proposal to require hospitals to report data on six eCQMs for two quarters (as compared to our previously finalized requirements to report data on eight eCQMs for four quarters) to represent an annual burden reduction of 11,000 hours across all 3,300 IPPS hospitals participating in the Hospital IQR Program (−200 minutes per hospital/60 minutes per hour × 3,300 hospitals = −11,000 hours). Using the wage estimate described above, we expect this to represent a cost reduction of $361,240 ($32.84 hourly wage × 11,000 annual hours reduction) across all 3,300 IPPS hospitals participating in the Hospital IQR Program.

(2) Calculations for the CY 2018 Reporting Period/FY 2020 Payment Determination

Using the same estimate as described above of 10 minutes per record per quarter, if our proposed updates to the CY 2018 reporting period/FY 2020 payment determination are finalized, we anticipate our proposal to require: (1) Reporting on at least 6 of the available eCQMs; and; (2) submission of the first three quarters of CY 2018 eCQM data, would result in a burden reduction of 2 hours and 20 minutes (140 minutes) per hospital for the FY 2020 payment determination as compared to the previously finalized requirements to report eight eCQMs for four quarters for the FY 2020 payment determination (81 FR 57157 through 57159). This estimate was calculated by considering the burden difference between the updated eCQM reporting requirements proposed for the FY 2020 payment determination (10 minutes per record ¥ eCQMs ¥ 3 quarters = 180 minutes for 3 quarters of reporting) and the eCQM reporting

requirements previously finalized for the FY 2020 payment determination (10 minutes per record × 8 eCQMs × 4 quarters = 320 minutes for 4 quarters of reporting). Through these calculations (180 minutes–320 minutes), we arrived at a reduction of 140 minutes per hospital per year, or 2 hours and 20 minutes per hospital per year, for the FY 2020 payment determination as compared to the previously finalized requirements for the FY 2020 payment determination. In total, this would represent an annual burden reduction of 7,700 hours across all 3,300 IPPS hospitals participating in the Hospital IQR Program ( 9,100 minutes per hospital/60 minutes per hour × 3,300 hospitals) and a cost reduction of $252,868 ($32.84 hourly wage × 7,700 annual hours reduction) across all 3,300 IPPS hospitals.

b. Burden Estimate for Proposed Modifying to eCQM Certification Requirements for the FY 2019 and FY 2020 Payment Determinations and Subsequent Years

In section IX.10.d of the preamble of this proposed rule, we discuss our proposed changes to the Hospital IQR Program eCQM submission requirements to align them with the Medicare EHR Incentive Program for eligible hospitals and CAHs. Specifically, for the CY 2017 reporting period/FY 2019 payment determination, we are proposing that: (1) A hospital using EHR technology certified to the 2014 or 2015 Edition, but such EHR technology is not certified to all available eCQMs, would be required to have its EHR technology certified to all eCQMs that are available to report; and (2) EHR technology that is certified to all available eCQMs would not need to be recertified each time it is updated to a more recent version of the eCQM specifications. For the CY 2018 reporting period/FY 2020 payment determination, we are proposing that: (1) A hospital using EHR technology certified to the 2015 Edition, but such EHR technology is not certified to all available eCQMs, would be required to have its EHR technology certified to all eCQMs that are available to report; and (2) an EHR certified for all available eCQMs under the 2015 Edition of CEHRT would not need to be recertified each time it is updated to a more recent version of the eCQM specifications.

Further, we are proposing that: (1) For the CY 2017 reporting period, hospitals would be required to use the most recent version of the CQM electronic specifications; Spring 2016 version of the eCQM specifications and any applicable addenda. For eCQM specifications, we refer readers to the eCQI Resource Center Web site at: https://ecqi.healthit.gov/. Because the use of certified EHR technology is already required for the Medicare EHR Incentive Program, we believe that these proposals will have no effect on burden for hospitals under the Hospital IQR Program.

c. Burden Estimate for Proposed Modifications to the Existing Validation Processes

In section IX.A.11. of the preamble of this proposed rule, we discuss our proposal to adopt a modification to the existing eCQM data validation process for the Hospital IQR Program data beginning with validation for the FY 2020 payment determination. First, we are proposing to require eight cases to be submitted per quarter for eCQM validation for the FY 2020 payment determination and subsequent years. We are making this proposal in conjunction with our proposal to require two quarters of data for the CY 2017 eCQM reporting period and our proposal to require three quarters of data for the CY 2018 eCQM reporting period. Accordingly, if those eCQM reporting proposals are finalized, we are proposing that the number of required case files for validation would be 16 records (eight cases per quarter over two quarters) for the FY 2020 payment determination and 24 records (eight cases per quarter over three quarters) for the FY 2021 payment determination. We note that, as discussed in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57176), CY 2017 eCQM data will be validated beginning in CY 2018 for the FY 2020 payment determination and subsequent years. Therefore, CY 2018 data will be validated beginning in CY 2019 for the FY 2021 payment determination. Second, we are proposing to add additional exclusion criteria to our hospital and case selection process for eCQM validation for the FY 2018 reporting period/FY 2020 payment determination and subsequent years. Third, we are proposing to continue our previously finalized medical record submission requirements for the FY 2021 payment determination and subsequent years as well as to provide clarification to our finalized policy. We believe the updates to the exclusions and maintaining previously finalized medical record submission requirements will have no effect on burden for hospitals. We discuss the burden associated with the proposed eCQM validation process in more detail below.

In previous years (79 FR 50347), we estimated a burden of 16 hours (960 minutes) for the submission of 12 records, which will equal 1 hour and 20 minutes (or 80 minutes) per record (960 minutes/12 records) for validation of eCQM data. Applying the time per individual submission of 1 hour and 20 minutes (or 80 minutes) per record for the 16 records we are proposing that hospitals submit for validation for the FY 2020 payment determination, we estimate a total burden of approximately 21 hours (80 minutes × 16 records/60 minutes per hour) for each hospital selected for participation in eCQM validation for the FY 2020 payment determination. We estimate that the total burden would be approximately 4,200 hours across the 200 hospitals selected for eCQM validation (21 hours per hospital × 200 hospitals = 4,200 hours). As compared to our total burden estimate of 8,533 hours previously estimated in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57261), this represents a burden reduction of approximately 4,333 hours across up to 200 hospitals selected for eCQM validation (4,200 hours estimated in this proposed rule – 8,533 hours estimated in the FY 2017 IPPS/LTCH final rule = – 4,333 hours). Using the estimated hourly labor cost of $32.84, we estimate an annual cost reduction of $142,296 (4,333 hours × $32.84 per hour) across the 200 hospitals selected for eCQM validation due to our proposal to decrease the number of records collected for validation from 32 records to 16 records for the FY 2020 payment determination.

(2) Calculations for Proposed Modifications to the Validation of eCQM Data for the FY 2021 Payment Determination and Subsequent Years

Applying the time per individual submission of 1 hour and 20 minutes (or 80 minutes) per record for the 24 records we are proposing that hospitals submit for eCQM validation for the FY 2021 payment determination, we estimate a total burden of approximately 32 hours (80 minutes × 24 records/60 minutes per hour) for each hospital selected for participation in eCQM validation. We estimate that the total burden would be approximately 6,400 hours across the 200 hospitals selected...
for eCQM validation (32 hours per hospital \times 200 hospitals = 6,400 hours). We note that compared to our total burden estimate of 8,533 hours previously estimated in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57261) for the FY 2020 payment determination and subsequent years, this would represent a burden reduction of approximately 2,133 hours across up to 200 hospitals selected for eCQM validation for the FY 2021 payment determination (6,400 hours estimated for the FY 2021 payment determination in this proposed rule—8,533 hours estimated in the FY 2017 IPPS/LTCH PPS final rule = −2,133 hours). Using the estimated hourly labor cost of $32.84, we estimate an annual cost reduction of $70,048 (2,133 hours × $32.84 per hour) across the 200 hospitals selected for eCQM validation due to our proposal to reduce the number of records collected from 32 records as finalized in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57178) to 24 records for the FY 2021 payment determination.

(3) Calculations for Proposed Modifications to the eCQM Validation Exclusions for the FY2020 Payment Determination and Subsequent Years

In section IX.A.11.b. of the preamble of this proposed rule, we are proposing a new eCQM validation exclusion criterion. Specifically, hospitals that do not have at least five discharges for at least one reported eCQM (among the six required eCQMs proposed for the CY 2017 and CY 2018 eCQM reporting periods) included in their QRDA I file submissions would be excluded from the random sample of up to 200 hospitals selected for eCQM validation for the FY 2020 payment determination and subsequent years. In summary, for the FY 2020 payment determination and subsequent years, we would exclude hospitals meeting the newly proposed exclusion criterion discussed above and/or either of the two exclusion criteria finalized in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57178). Lastly, we are proposing that the three exclusions would be applied before the random selection of 200 hospitals for eCQM validation, such that hospitals meeting any of these exclusions would not be eligible for selection.

In section IX.A.11.b. of the preamble of this proposed rule, we are proposing to exclude the following cases from validation for those hospitals selected to participate in eCQM validation: (1) Episodes of care that are longer than 120 days; and (2) cases with a zero denominator for each measure, for the FY 2020 payment determination and subsequent years.

We do not believe that these proposals will impact the burden experienced by hospitals because, while they influence which hospitals and cases would be selected, they would not change the number of hospitals that must participate in eCQM validation, the number of records that would be collected for validation, or the validation reporting requirements for the hospitals selected.

(4) Calculations for the Proposed Modifications to the Medical Record Submission Requirements for the FY 2021 Payment Determination and Subsequent Years

In section IX.A.11.b. of the preamble of this proposed rule, we are proposing for the FY 2021 payment determination and subsequent years, to apply the medical record submission requirements that were finalized in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57179) only for the FY 2020 payment determination. Specifically, we are proposing that for hospitals participating in eCQM validation we: (1) Would require submission of at least 75 percent of sampled eCQM measure medical records in a timely and complete manner; and (2) would maintain the previously finalized policy that the accuracy of eCQM data submitted for validation would not affect a hospital’s validation score (81 FR 57180). We do not expect these proposals to influence our burden estimates, as we are proposing to continue existing policies.

(5) Calculations for the Proposed Educational Review Process for Chart-Abstracted Measures for the FY 2020 Payment Determination and Subsequent Years

In section IX.A.11.c. of the preamble of this proposed rule, we are proposing to formalize the process of allowing hospitals to use an educational review process to correct incorrect validation results for the first three quarters of validation for chart-abstracted measures. Secondly, we are proposing to update the process to specify that if the results of an educational review indicate that we incorrectly scored a hospital, the corrected score would be used to compute the hospital’s final validation score whether or not the hospital submits a reconsideration request. Under this proposal, the educational review request process, as well as CMS’ procedures for responding to requests, remain the same for the FY 2020 payment determination and subsequent years, except that revised scores identified through an educational review would be used to correct a hospital’s validation score. As stated in the FY 2016 IPPS/LTCH PPS final rule (80 FR 49762), we estimate a burden of 15 minutes per hospital to report structural measure data and to complete all forms, including the reconsideration request form and the educational review form. We refer readers to the FY 2017 IPPS/LTCH PPS final rule for more detailed information on the burden associated with the chart-abstracted validation requirements (81 FR 57260). Although this proposal may allow hospitals to avoid the formal reconsideration process, we do not expect this proposal to influence our burden estimates for the chart-abstracted measures validation process as it would not change the requirements for selecting hospitals for validation of chart-abstracted measures nor change the chart-abstracted validation reporting requirements for the selected hospitals.

e. Burden Estimate for the Proposed Voluntary Reporting on the Hybrid Hospital-Wide 30-Day Readmission Measure for the CY 2018 Reporting Period

In section IX.A.7.a. of the preamble of this proposed rule, we are proposing voluntary reporting on the Hybrid Hospital-Wide 30-Day Readmission measure for CY 2018 reporting period. This measure uses both claims-based data as well as a set of 13 core clinical data elements from patient electronic health records (EHRs). We do not expect any additional burden to hospitals to report the claims-based portion of this measure because these data are already reported to the Medicare program for payment purposes.

As described in section IX.A.7.b. of the preamble of this proposed rule, we are proposing that hospitals submit the 13 core clinical data elements and the six data elements required for linking with claims data for this measure using the same submission process required for eCQM reporting, specifically, that these data be reported using QRDA I files submitted to the CMS data receiving system. Accordingly, we expect the burden associated with voluntarily reporting this measure to be similar to our estimates for eCQM reporting (that is 10 minutes per measure, per quarter). We anticipate that approximately 100 hospitals would voluntarily report the Hybrid Hospital-Wide 30-Day Readmission measure. Therefore, using the estimate of 10 minutes per measure per quarter, we estimate that our proposal would result in a burden increase of 0.67 hours (40 minutes) per participating hospital for
the one year (4 quarters) during which this pilot would take place (10 minutes per record × 1 measure × 4 quarters/60 minutes per hour = 0.67 hours). In total, for the one year duration of voluntary reporting the Hybrid Hospital-Wide 30-Day Readmission measure, we estimate an annual burden increase of 67 hours across up to 100 hospitals voluntarily participating (40 minutes per hospital/60 minutes per hour × 100 hospitals = 67 hours). Using the wage estimate described above, we estimate this to represent a cost increase of $2,200 ($32.84 hourly wage × 67 annual hours) across up to 100 hospitals voluntarily participating in the pilot. We note that the claims-based version of the Hospital-Wide All-Cause Unplanned Readmission (HWR) measure is currently a part of the Hospital IQR Program measure set, as adopted in the FY 2013 IPPS/LTCH PPS final rule (77 FR 53350).

f. Burden Estimate for the Proposed Refinement of the HCAHPS Survey Measure for the FY 2020 Payment Determination and Subsequent Years

In section IX.A.6.a. of the preamble of this proposed rule, we are proposing to update the HCAHPS Survey measure by replacing the current Pain Management questions (HCAHPS Q12, Q13, and Q14) with new questions referred to collectively as the “Communication About Pain” composite measure beginning with the FY 2020 payment determination. There is no additional information collection burden associated with the refinement of these questions because we are wording the existing questions and not changing the total number of questions. In addition, consistent with previous years (81 FR 57261), the burden estimate for the Hospital IQR Program excludes the burden associated with the HCAHPS survey measure, which is submitted under a separate information collection request and approved under OMB control number 0938–0981.

g. Burden Estimate for the Proposed Refinement of the Hospital 30-Day, All-Cause, Risk-Standardized Mortality Rate Following Acute Ischemic Stroke Measure for the FY 2023 Payment Determination and Subsequent Years

In section IX.A.6.b. of the preamble of this proposed rule, we are proposing to update the Hospital 30-Day, All-Cause, Risk-Standardized Mortality Rate Following Acute Ischemic Stroke measure to include the use of NIH stroke scale claims data for risk adjustment beginning with the FY 2023 payment determination. Because this proposed update would result only in the inclusion of additional claims-based data that are already reported to the Medicare program for payment purposes, we believe no additional burden on hospitals will result from the update to the stroke mortality measure.

h. Burden Estimate for the Confidential and Potential Future Public Reporting of Readmission Measure Data Stratified by Social Risk Factors

In section IX.A.13 of the preamble of this proposed rule we discuss our intent to provide confidential reports to hospitals that include measure data stratified by dual eligible status for the Hospital 30-day, All-Cause, Risk-Standardized Readmission Rate Following Pneumonia Hospitalization and Hospital 30-day, All-Cause, Risk-Standardized Mortality Rate (RSRR) for Pneumonia measures. In addition to confidential reporting, we are seeking comment on options for public display of measure data stratified by social risk factor indicators on the Hospital Compare Web site. Because this proposal is related to the way we display data, and not the methods of data collection implemented by the hospitals, we believe no additional burden on hospitals would result from confidential reporting of stratified measure data using social risk factor indicators. We note that all measures for which we might consider confidential reporting or public display of stratified measure data would already be included in the Hospital IQR Program, and as claims-based measures, we do not expect any additional burden because these data are already reported to the Medicare program for payment purposes.

i. Burden Estimate for the Proposed Changes to the Hospital IQR Program Extraordinary Circumstances Exceptions (ECE) Policy for the FY2020 Payment Determination and Subsequent Years

In section IX.A.15.b. of the preamble of this proposed rule we discuss our intent to align the naming of this exception policy and update CFR 412.140 to reflect our current ECE policies. We also are clarifying the timing of CMS’ response to ECE requests. Because we are not seeking any new or additional information in our ECE proposals, we believe the updates will have no effect on burden for hospitals.

j. Summary of Burden Estimates for the Hospital IQR Program

In summary, under OMB control number 0938–1022, we estimate: (1) A total burden reduction of 11,000 hours (−11,000 hours due to the proposed updates to the CY 2017 eCQM reporting requirements) and a total cost reduction of $361,240 (−11,000 hours × $32.84 per hour) for the FY 2019 payment determination; (2) a total burden reduction of 11,966 hours (−7,700 hours due to the proposed updates to the CY 2018 eCQM reporting requirements) and a total cost reduction of $392,963 (−7,700 hours × $32.84 per hour) for the FY 2019 payment determination; (3) a total burden reduction of 2,133 hours (−2,133 hours due to the proposed updates to eCQM validation procedures for the FY 2020 payment determination) and a total cost reduction of $70,048 (−2,133 hours × $32.84 per hour) for the FY 2021 payment determination. These are the burden estimate totals for which we are requesting OMB approval under OMB number 0938–1022.
<table>
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<th>Activity</th>
<th>Estimated time per record (minutes) FY 2019</th>
<th>Number reporting quarters per year FY 2019</th>
<th>Number of IPPS hospitals reporting</th>
<th>Average number records per hospital per quarter</th>
<th>Annual burden (hours) per hospital</th>
<th>Newly proposed annual burden (hours) across IPPS hospitals for FY 2019 payment determination</th>
<th>Previously finalized annual burden (hours) across IPPS hospitals for FY 2019 payment determination per the FY 2017 IPPS/LTCH PPS final rule</th>
<th>Net difference in annual burden hours</th>
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<tr>
<td>Reporting on 6 eCQMs for 2 Quarters.</td>
<td>80 (10 minutes × 6 measures).</td>
<td>2</td>
<td>3,300</td>
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<th>Number of IPPS hospitals reporting</th>
<th>Average number records per hospital per quarter</th>
<th>Annual burden (hours) per hospital</th>
<th>Newly proposed annual burden (hours) across IPPS hospitals for FY 2020 payment determination</th>
<th>Previously finalized annual burden (hours) across IPPS hospitals per the FY 2017 IPPS/LTCH PPS final rule</th>
<th>Net difference in annual burden hours</th>
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<th>Average number records per hospital per quarter</th>
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<th>Previously finalized annual burden (hours) across IPPS hospitals per the FY 2017 IPPS/LTCH PPS final rule</th>
<th>Net difference in annual burden hours</th>
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<td>Total Cost Estimate: Hourly Wage ($32.84) × Change in Burden Hours (−2,133)</td>
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7. ICRs for PPS-Exempt Cancer Hospital Quality Reporting (PCHQR) Program

As discussed in sections IX.B. of the preamble of this proposed rule, section 1866(k)(1) of the Act requires, for purposes of FY 2014 and each subsequent fiscal year, that a hospital described in section 1886(d)(1)(B)(v) of the Act (a PPS-exempt cancer hospital, or a PCH) submit data in accordance with section 1866(k)(2) of the Act with respect to such fiscal year. There is no financial impact to PCH Medicare reimbursement if a PCH does not participate.

We refer readers to the FY 2014 IPPS/LTCH PPS final rule (78 FR 50957 through 50959), the FY 2015 IPPS/LTCH PPS final rule (79 FR 50347 through 50348), the FY 2016 IPPS/LTCH PPS final rule (80 FR 49764), and the FY 2017 IPPS/LTCH PPS final rule (81 FR 57182), as well as to OMB Control Number 0938–1175, for a detailed discussion of the burden for the program requirements that we have previously adopted. Below we discuss only changes in burden that would result from the proposals in this proposed rule.
a. Estimated Hourly Labor Cost

Previously, we used $66 as our hourly labor cost in calculating the burden associated with chart-abstraction activities in the PCHQR Program. However, our experience working with our data analysis contractors and those performing chart abstraction indicates that this work is performed by a different labor category than we previously thought. In addition, our previous labor cost is different from those used in other quality reporting and value-based purchasing programs, and we do not believe there is a justification for these different values given the similarity in quality measures and required staff. Therefore, to align the estimated hourly labor costs (hourly wage plus overhead, as discussed below) used to calculate burden in the PCHQR Program with those used in other CMS quality reporting programs, including the Hospital IQR Program, we are proposing to revise our hourly labor cost estimate to $32.84.509

This labor cost is based on the BLS wage for a Medical Records and Health Information Technician. The BLS is “the principal Federal agency responsible for measuring labor market activity, working conditions, and price changes in the economy.”510 The BLS describes Medical Records and Health Information Technicians as those responsible for organizing and managing health information data; therefore, we believe it is reasonable to assume that these individuals would be tasked with abstracting clinical data for submission for the PCHQR Program. According to the BLS, the median pay for Medical Records and Health Information Technicians is $16.42 per hour, before inclusion of overhead and fringe benefits.

Obtaining data on overhead costs is challenging because overhead costs vary across PCHs, and cost elements assigned to overhead are subjective to interpretation at the facility level. Therefore, we are proposing to calculate the cost of overhead, including fringe benefits, at 100 percent of the mean hourly wage, as is currently done in other CMS quality reporting programs.511 This is necessarily a rough adjustment, both because fringe benefits and overhead costs vary significantly from employer to employer and because methods of estimating these costs vary widely from study to study. Nonetheless, we believe that doubling the hourly wage rate ($16.42 × 2 = $32.84) to estimate total cost is a reasonably accurate estimation method. Accordingly, we are proposing to use an hourly labor cost estimate of $32.84 ($16.42 base salary + $16.42 fringe and overhead) for calculation of burden forthwith. We note that more recent wage data has become available, and we intend to update the wage rate used in these calculations in the FY 2018 IPPS/LTCH PPS Final rule. We are inviting public comment on this proposal.

b. Estimated Burden of PCHQR Program Proposals for the FY 2020 Program Year

In section IX.B.4. of the preamble of this proposed rule, we are proposing to adopt four claims-based measures beginning with the FY 2020 program: (1) Proportion of Patients Who Died from Cancer Receiving Chemotherapy in the Last 14 Days of Life (NQF #0210); (2) Proportion of Patients Who Died from Cancer Admitted to the ICU in the Last 30 Days of Life (NQF #0213); (3) Proportion of Patients Who Died from Cancer Not Admitted to Hospice (NQF #0215); and (4) Proportion of Patients Who Died from Cancer Admitted to Hospice for Less Than Three Days (NQF #0216). In conjunction with our proposal in section IX.B.3. of the preamble of this proposed rule, we are proposing to remove three existing chart-abstracted measures beginning with the FY 2020 program—(1) Adjuvant Chemotherapy is Considered or Administered Within 4 Months (120 Days) of Diagnosis to Patients Under the Age of 80 with AJCC III (Lymph Node Positive) Colon Cancer (PCH–01/NQF #0223); (2) Combination Chemotherapy is Considered or Administered Within 4 Months (120 Days) of Diagnosis for Women Under 70 with AJCC T1c, or Stage II or III Hormone Receptor Negative Breast Cancer (PCH–02/NQF #0559); and (3) Adjuvant Hormonal Therapy (PCH–03/ NQF #0220). If all of these proposals are finalized, the PCHQR Program measure set would consist of 18 measures for the FY 2020 program.

Our proposal to remove the three chart-abstracted measures would reduce the burden associated with quality data reporting on PCHs. Based on the FY 2013 IPPS/LTCH final rule (77 FR 53667) finalized estimates of the burden of collecting measure information, submitting measure information, and training personnel, we estimate the reduction in burden for collecting measure information, submitting measure information, and training personnel provided by the proposed removal of the three measures to be approximately 3,776 hours per year for each PCH, or an average reduction in burden of 315 hours per month per PCH. Therefore, we estimate a reduction in hourly burden of chart abstraction and data submission of approximately 41,536 hours per year across the 11 PCHs.512

We do not anticipate any increase in burden on the PCHs corresponding to our proposal to adopt four claims-based measures into the PCHQR Program beginning with the FY 2020 program year. The four measures are claims-based and therefore do not require facilities to report any additional data. Because these measures do not require facilities to submit any additional data, we do not believe that there is any increase in burden associated with this proposal.

In summary, as a result of our proposals, we estimate a reduction of 40,910 hours of burden per year associated with the proposals above for all 11 PCHs beginning with the FY 2020 program. Coupled with our revised estimated salary costs, we estimate that these proposed changes would result in a reduction in annual labor costs of $1,364,078 (41,537.1 hours × $32.84 hourly labor cost) across the 11 PCHs beginning with the FY 2020 PCHQR Program. The burden associated with these reporting requirements is currently under OMB Control Number 0938–1175. The information collection will be revised and submitted to OMB.

8. ICRs for the Hospital Value-Based Purchasing (VBP) Program

In section V.J. of the preamble of this proposed rule, we discuss proposed requirements for the Hospital VBP Program. Specifically, in this proposed rule, with respect to quality measures, we are proposing to: (1) Remove the current PSI 90 measure beginning with the FY 2019 program year; (2) adopt the Hospital-Level, Risk-Standardized Payment Associated with a 30-Day Episode-of-Care for Pneumonia PN Payment measure beginning with the FY 2022 program year; and (3) adopt the Patient Safety and Adverse Events

511 See, e.g., FY2016 IPPS/LTCH Final Rule at 80 FR 49764 FN 154.
As required under section 1886(o)(2)(A) of the Act, Hospital VBP Program measures, including the proposed additional and updated measures, are used in the Hospital IQR Program. Therefore, their inclusion in the Hospital VBP Program does not result in any additional burden because the Hospital VBP Program uses data that are required for and collected under the Hospital IQR Program. Therefore, the burden associated with these reporting requirements is currently approved under OMB Control Number 0938–1022.

LTCH QRP QUALITY MEASURES PROPOSED IN THIS FY 2018 IPPS/LTCH PPS PROPOSED RULE BEGINNING WITH THE FY 2020 LTCH QRP

The LTCH QRP measure set also currently includes claims-based measures that are calculated based on data that LTCHs are already required to report to the Medicare program for payment purposes. In this proposed rule, we are proposing to remove the All-Cause Unplanned Readmission Measure for 30 Days Post-Discharge from LTCHs (NQF #2512) from the LTCH QRP measure set, beginning with the FY 2019 LTCH QRP. However, because LTCHs will still be required to report data on this measure for payment purposes, we believe that the removal of this measure will not affect the burden estimate for the LTCH QRP.

The LTCH CARE Data Set Version 3.00 was implemented April 1, 2016 and is approved under OMB control number 0938–1163. The LTCH CARE Data Set Version 3.00 is available on the LTCH-QRP Web site at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-
to a decrease of 8,796 hours in burden for all LTCHs (0.06 hours × 146,592 discharges). Given 3.6 minutes of RN time at $69.40 per hour completing an average of 344 sets of LTCH CARE Data Set assessments per provider per year, we estimated the total cost would be reduced by $1,432.89 per LTCH annually, or $610,409 for all LTCHs annually. This decrease in burden will be accounted for in the information collection under OMB control number (0938–1163).

Also, in section IX.C.10.b. of the preamble of this proposed rule, we are proposing requirements related to the reporting of standardized patient assessment data beginning with the FY 2019 LTCH QRP. Some of the proposed data elements are already included on the LTCH CARE Data Set, and our proposal to characterize those data elements as standardized patient assessment data will not result in an additional reporting burden for LTCHs. However, we are proposing to adopt 25 new standardized patient assessment data elements with respect to LTCH admissions and 17 new standardized patient assessment data elements with respect to LTCH discharges. We estimate that it will take an LTCH’s clinical staff 7.5 minutes to report the data elements required with respect to admissions and 5.1 minutes to report the data elements required with respect to discharges, for a total of additional 12.6 minutes. This equates to an increase of 30,784 hours in burden for all LTCHs (0.21 hours × 146,592 discharges). We believe that the additional LTCH CARE Data Set data elements we are proposing will be completed by registered nurses (approximately 45 percent of the time), licensed vocational nurses (approximately 45 percent of the time) and respiratory therapists (approximately 10 percent of the time). We estimate 146,592 discharges from 426 LTCHs annually.

We obtained mean hourly wages for these staff from the U.S. Bureau of Labor Statistics’ May 2016 National Occupational Employment and Wage Estimates (https://www.bls.gov/oes/current/oes_nat.htm). The mean hourly wage for a RN (BLS occupation code: 29–1141) is $34.70, for a respiratory therapist (BLS occupation code: 29–1126) $29.15, and for a licensed vocational nurse (BLS occupation code: 29–2061) $21.56. Individual providers determine the staffing resources necessary. However, to account for overhead and fringe benefits, we have doubled the mean hourly wage, making it $69.40 for an RN, $58.30 for a respiratory therapist, and $43.12 for a licensed vocational nurse. Given the clinician times and wages above completing an average of 344 sets of LTCH CARE Data Set assessments per provider per year, the total cost related to the additional standardized patient assessment data elements is estimated at $4,080.30 per LTCH annually, or $1,738,206 for all LTCHs annually.

In summary, the 4.5-minute increase in burden for the two proposed ventilator weaning quality measures is offset with the 3 minute reduction in burden for the proposed pressure ulcer quality measure and the 3.6 minute reduction in burden for the program interruption items. This results in a net reduction in burden of 2.1 minutes. In addition, we are proposing that data for the new standardized data elements will be collected by LTCHs and reported to CMS using the LTCH CARE Data Set (LTCH CARE Data Set Version 4.00, effective April 1, 2018) for the purpose of fulfilling the requirements of the IMPACT Act. This results in an additional 12.6 minutes of burden for the proposed standardized data elements, with a net burden of 10.5 minutes. Overall, the cost associated with the proposed changes to the LTCH QRP is estimated at an additional $3,187.15 per LTCH annually, or $1,357,726 for all LTCHs annually.

The proposed LTCH CARE Data Set Version 4.00 is available on the LTCH QRP Web site at: https://www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/LTCH-Quality-Reporting/LTCH-Quality-Reporting-Measures-Information.html. For a discussion of burden related to LTCH CARE Data Set Version 4.00, we refer readers to section I.M. of Appendix A of the preamble of this proposed rule.

10. ICRs for the Inpatient Psychiatric Facility Quality Reporting (IPFQR) Program

We refer readers to the FY 2015 IPF PPS final rule (79 FR 45978 through 45980), the FY 2016 IPF PPS final rule (80 FR 46720 through 46721), and the FY 2017 IPPS/LTC PPS (81 FR 57265 through 57266) final rule for a detailed discussion of the burden for the program requirements that we have previously adopted. Additional information on the full burden of existing requirements can also be found in the materials associated with OMB 0938–1171, the OMB Paperwork Reduction Act materials for this Program. We are proposing provisions that affect the FY 2019 payment determination (through procedural requirements that occur in FY 2018) and the FY 2020 payment determination and subsequent years. ICRs associated with proposals for each period are discussed in more detail below.

a. Burden Associated With Procedural Proposals for the FY 2019 Payment Determination and Subsequent Years

For FY 2018 and subsequent years we are proposing: (1) Updates to the Extraordinary Circumstances Exception (ECE) process (affecting submission of ECE requests in FY 2018, which would impact payment determination year FY 2019 and subsequent years); (2) to adopt measure removal factors, including criteria for determining when a measure is “tapped-out,” and measure retention factors (which could affect measures for the FY 2020 payment determination and subsequent years); and (3) changes associated with procedural deadlines (which affects FY 2019 payment determination and subsequent years).

For the ECE proposals, we are specifically proposing to: (1) Specify that ECE forms can be signed by either the CEO or the designated personnel as listed in the contact information section of the form; (2) change the ECE request form submission deadline to within 90 days of the date that the extraordinary circumstance occurred; and (3) we will strive to complete our review of ECE requests within 90 days of receipt. These changes to the ECE process would not change data submission requirements for facilities requesting ECEs, but update procedural requirements related to ECE requests instead. Therefore, we do not expect any changes to burden associated with these proposals.

Second, the proposal to adopt measures removal and retention factors does not affect the data submission requirements. These factors are intended to improve transparency of our measure review and evaluation process. Third, for the procedural deadlines, we are proposing to: (1) Change the submission deadline such that facilities have a 45-day submission period beginning at least 30 days following the end of the data collection period for a measure; (2) change the submission timeframes for both NOPs and withdrawals to the end of the data submission period before each respective payment determination year; and (3) provide exact dates that define the end of the data submission period/ NOP/withdrawal submission deadline through subregulatory means. These proposals do not affect the data that a facility must submit, instead these proposals affect the specification of timeframes.

Because none of the policies that we are proposing for FY 2018 and subsequent years affects the data that
IPFs are required to submit, we do not believe there will be any change in burden.

b. Burden Associated With Proposal for the FY 2020 Payment Determination and Subsequent Years

For FY 2020 and subsequent years, we are proposing one measure, Medication Continuation following Inpatient Psychiatric Discharge. This measure is claims based and therefore does not require facilities to report any additional data. Because this measure does not require facilities to submit any additional data, we do not believe that there is any associated burden associated with this proposal.

11. ICRs for the Electronic Health Record (EHR) Incentive Programs and Meaningful Use

In section IX.E. of the preamble of this proposed rule, we discuss proposed policies for eligible hospitals and CAHs reporting CQMs electronically under the Medicare and Medicaid EHR Incentive Programs. As outlined in this proposed rule, we are proposing the following modifications to the CY 2017 final CQM policies: (1) Revise the CY 2017 reporting period for eligible hospitals and CAHs reporting CQMs electronically to require the submission of 2 self-selected quarters of data; and (2) revise the number of CQMs eligible hospitals and CAHs are required to report electronically for CY 2017 to 6 (self-selected) available CQMs. In addition, we are proposing the following CQM reporting requirements for CY 2018: (1) Eligible hospitals and CAHs reporting CQMs electronically that demonstrate meaningful use for the first time in 2018 or that have demonstrated meaningful use in any year prior to 2018, the reporting period would be the first 3 quarters of CY 2018 with a submission period (Medicare EHR Incentive Program only) consisting of the 2 months following the close of the calendar year, ending on February 28, 2019; (2) eligible hospitals and CAHs reporting CQMs electronically would be required to report at least 6 (self-selected) of the available CQMs; (3) eligible hospitals and CAHs that report CQMs by attestation under the Medicare EHR Incentive Program because electronic reporting is not feasible, and eligible hospitals and CAHs that report CQMs by attestation under their State’s Medicaid EHR Incentive Program, would be required to report on all 16 available CQMs; and (4) eligible hospitals and CAHs reporting CQMs by attestation under the Medicare EHR Incentive Program would have a submission period that would be the 2 months following the close of the CY 2018 CQM reporting period, ending February 28, 2019.

Because the proposed reporting requirements for data collection regarding the reporting of CQMs electronically under the Medicare and Medicaid EHR Incentive Programs would align with the reporting requirements under the Hospital IQR Program, we do not believe that there is any additional burden for the collection of such information. We are not proposing modifications for the CQMs reporting requirements by attestation. Therefore, there would be no change in burden associated with attestation of CQMs.

In section IX.F. of the preamble of this proposed rule, we discuss proposed policies regarding clinical quality measurement for EPs participating in the Medicaid EHR Incentive Program. We note that there may be costs incurred by States associated with systems development as a result of the proposed policies. State attestation systems would likely require minor updates, which may be eligible for support through enhanced Federal funding, subject to CMS prior approval, if outlined in an updated Implementation Advanced Planning Document (IAPD). We anticipate that EPs may also face minor burden and incremental capital cost for updating clinical quality measures and reporting capabilities in the EHR. We intend to reduce EP burden and simplify the program through these proposals, which are intended to better align CQM reporting periods and CQM reporting for the Medicaid EHR Incentive Program with policies under MIPS. Overall, we believe the proposed CQM alignment at the State attestation system and EP levels would both reduce burden associated with reporting on multiple CMS programs and enhance state and CMS operational efficiency.

In section IX.G.1. of the preamble of this proposed rule, we discuss our proposals to change the EHR reporting period in 2018 from the full CY 2018 to any continuous 90-day period within CY 2018 for all returning EPs, eligible hospitals and CAHs in the Medicare and Medicaid EHR Incentive Programs. We do not believe that modifying the EHR reporting period would cause an increase in burden as the reporting requirements for a 90 day reporting period are virtually the same for a full calendar year reporting period and the same objectives and measures will be used for reporting for a full calendar year reporting or a 90 day reporting period.

In section IX.G.2. of the preamble of this proposed rule, as required by the 21st Century Cures Act (Pub. L. 114–255), we are proposing an exemption from the payment adjustments under sections 1848(a)(7)(A), 1886(b)(3)(B)(ix)(I), and 1814(l)(4) of the Act for EPs, eligible hospitals and CAHs, respectively, that demonstrate through an application process that compliance with the requirement for being a meaningful EHR user is not possible because their certified EHR technology has been decertified under ONC’s Health IT Certification Program. The application process involves participants completing an application form for an exception. While the form is standardized, we believe it is exempt from the PRA. The form is structured as an attestation. Therefore, we believe it is exempt under 5 CFR 1320.3(h)(1) of the implementing regulations of the PRA. The form is an attestation that imposes no burden beyond what is required to provide identifying information and to attest to the applicable information.

In section IX.G.3. of the preamble of this proposed rule, as required by the 21st Century Cures Act, we are proposing to exempt ambulatory surgical center-based EPs from the 2017 and 2018 payment adjustments under section 1848(a)(7)(A) of the Act if they furnish substantially all of their covered professional services in an ambulatory surgical center. We do not believe this requirement would cause an increase in burden as CMS would identify the EPs who might meet this requirement. For the expected effects relating to the above proposals, we refer readers to section I.O. of Appendix A of this proposed rule.

We are requesting public comments on these information collection and recordkeeping requirements.

12. ICRs Relating to Proposed Electronic Signature and Electronic Submission of the Certification and Settlement Summary Page of Medicare Cost Reports

In section X.A. of the preamble of this proposed rule, we discuss our proposal to allow providers to use an electronic signature on the certification statement of the Certification and Settlement Summary page of the Medicare cost report and submit it electronically. The Certification and Settlement Summary page, which contains the required provider signature line, currently exists in the Medicare cost report and is mailed to the contractor from the provider. We are proposing to allow providers the option to sign and submit this page electronically. The signature from the provider’s administrator or chief financial officer is an existing data
collection requirement. There would be no new data collection from providers resulting from our proposal. The proposal to allow providers to sign this page electronically is not a substantive change to the existing data collection instrument and would have a minimal impact on providers to complete.

13. ICRs Relating to Survey and Certification Requirements

a. Proposed Transparency in Survey Reports and Plans of Correction

In section XI.A. of the preamble of this proposed rule, we are proposing to require accrediting organizations (AO) to post survey results and findings (that is, statements of deficiency findings) as well as any associated acceptable plans of correction (PoCs), and make this information publicly available on its Web site within 90 days after such information is made available to those facilities, for the most recent 3 years.

According to data and information available to us, as of September 30, 2016, there are approximately 12,434 deemed facilities (providers and suppliers) across all CMS-approved programs that have surveys (other initial and renewal, including complaints) for which an AO would be required to make survey reports and associated PoCs publicly available under our proposal.\(^{513}\)\(^{514}\)\(^{515}\) The CY 2016 Advanced Diagnostic Imaging (ADI) AO annual data submission lists approximately 16,873 ADI suppliers and locations that have surveys (initial, renewal, complaint, and mid-cycle) (approximately 2,128) for which AOs of ADI suppliers would be required to make survey reports and PoCs publicly available on their Web site under our proposal. Unlike Medicare- and Medicaid-certified providers and suppliers, there are no prescriptive statutory, regulatory, or policy requirements regarding the frequency of ADI AOs surveys.

We do not have sufficient data to determine the burden associated with the information collection requirements under our proposal. Therefore, we are requesting public comments on the potential costs and burden associated with our proposal on AOs regarding modifying their existing public Web sites and uploading of survey reports and PoCs.

b. Proposed Changes in Public Notices of Terminations

In section XI.B. of the preamble of this proposed rule, we are proposing to no longer require the posting of voluntary and involuntary termination public notice in newspapers for RHCs, FQHCs, ASCs, and OPFs. These providers and suppliers would be permitted to use other methods of notification in light of the expanded use of information technology. We also are proposing to change the regulations regarding termination of provider agreements by CMS (that is, involuntary termination) or providers or suppliers to remove the provision for public notice through “newspapers” to allow flexibility in the method of public notice.

We believe none of the proposed provisions would have a financial burden as we are only eliminating the specification which requires newspaper hard print to be the notice source.

We refer readers to the economic impact provisions of section I.P. of Appendix A of this proposed rule for additional information.

If you comment on these information collection and recordkeeping requirements, please do either of the following:

1. Submit your comments electronically as specified in the ADDRESSES section of this proposed rule; or

2. Submit your comments to the Office of Information and Regulatory Affairs, Office of Management and Budget.

Attention: CMS Desk Officer, CMS–1677–P
Fax: (202) 395–6974; or Email: OIRA_submission@omb.eop.gov.models.

C. Request for Information on CMS Flexibilities and Efficiencies

CMS is committed to transforming the health care delivery system—and the Medicare program—by putting an additional focus on patient-centered care and working with providers, physicians, and patients to improve outcomes. We seek to reduce burdens for hospitals, physicians, and patients, improve the quality of care, decrease costs, and ensure that patients and their providers and physicians are making the best health care choices possible. These are the reasons we are including this Request for Information in this proposed rule.

As we work to maintain flexibility and efficiency throughout the Medicare program, we would like to start a national conversation about improvements that can be made to the health care delivery system that reduce unnecessary burdens for clinicians, other providers, and patients and their families. We aim to increase quality of care, lower costs improve program integrity, and make the health care system more effective, simple and accessible.

We would like to take this opportunity to invite the public to submit their ideas for regulatory, subregulatory, policy, practice, and procedural changes to better accomplish these goals. Ideas could include payment system redesign, elimination or streamlining of reporting, monitoring and documentation requirements, aligning Medicare requirements and processes with those from Medicaid and other payers, operational flexibility, feedback mechanisms and data sharing that would enhance patient care, support of the physician-patient relationship in care delivery, and facilitation of individual preferences. Responses to this Request for Information could also include recommendations regarding when and how CMS issues regulations and policies and how CMS can simplify rules and policies for beneficiaries, clinicians, physicians, providers, and suppliers. Where practicable, data and specific examples would be helpful. If the proposals involve novel legal questions, analysis regarding CMS’ authority is welcome for CMS’ consideration. We are particularly interested in ideas for incentivizing organizations and the full range of relevant professionals and paraprofessionals to provide screening, assessment and evidence-based treatment for individuals with opioid use disorder and other substance use disorders, including reimbursement methodologies, case coordination, systems and services integration, use of paraprofessionals including community paramedics and other strategies. We are requesting commenters to provide clear and concise proposals that include data and specific examples that could be implemented within the law.

We note that this is a Request for Information only. Respondents are encouraged to provide complete but concise responses. This Request for Information is issued solely for information and planning purposes; it does not constitute a Request for Proposal (RFP), applications, proposal abstracts, or quotations. This Request for Information does not commit the U.S. Government to contract for any supplies or services or make a grant award.

\(^{513}\) FY 2016 Report to Congress (RTC); Review of Medicare’s Program Oversight of Accrediting Organizations (AOs) and the Clinical Laboratory Improvement Amendments of 1988 (CLIA) Validation Program—Section 2, Table 5.

\(^{514}\) CMS Survey and Certification Web site for hospital Form CMS–2567 (Statement of Deficiencies and Plan of Correction) downloads: https://www.cms.gov/Medicare/Provider-Enrollment-and-Certification/CertificationandCompliance/Hospitals.html.

Further, CMS is not seeking proposals through this Request for Information and will not accept unsolicited proposals. Responders are advised that the U.S. Government will not pay for any information or administrative costs incurred in response to this Request for Information; all costs associated with responding to this Request for Information will be solely at the interested party’s expense. We note that not responding to this Request for Information does not preclude participation in any future procurement, if conducted. It is the responsibility of the potential responders to monitor this Request for Information announcement for additional information pertaining to this request. In addition, we note that CMS will not respond to questions about the policy issues raised in this Request for Information. CMS will not respond to comment submissions in response to this Request for Information in the FY 2018 IPPS/LTCH PPS final rule. Rather, CMS will actively consider all input as we develop future regulatory proposals or future subregulatory policy guidance. CMS may or may not choose to contact individual responders. Such communications would be for the sole purpose of clarifying statements in the responders’ written responses. Contractor support personnel may be used to review responses to this Request for Information. Responses to this notice are not offers and cannot be accepted by the Government to form a binding contract or issue a grant. Information obtained as a result of this Request for Information may be used by the Government for program planning on a nonattribution basis. Respondents should not include any information that might be considered proprietary or confidential. This Request for Information should not be construed as a commitment or authorization to incur cost for which reimbursement would be required or sought. All submissions become U.S. Government property and will not be returned. CMS may publically post the public comments received, or a summary of those public comments.

D. Response to Public Comments

Because of the large number of public comments we normally receive on Federal Register documents, we are not able to acknowledge or respond to them individually. We will consider all public comments we receive by the date and time specified in the DATES section of this preamble, and, when we proceed with a subsequent document, we will respond to the public comments in the preamble of that document.

List of Subjects

42 CFR Part 405

Administrative practice and procedure, Health facilities, Health professions, Kidney diseases, Medicare, Reporting and recordkeeping, Rural areas, X-rays.

42 CFR Part 412

Administrative practice and procedure, Health facilities, Medicare, Puerto Rico, Reporting and recordkeeping requirements.

42 CFR Part 413

Health facilities, Kidney diseases, Medicare, Puerto Rico, Reporting and recordkeeping requirements.

42 CFR Part 414

Administrative practice and procedures, Biologics, Drugs, Health facilities, Health professions, Kidney diseases, Medicare, Reporting and recordkeeping requirements.

42 CFR Part 416

Health facilities, Health professions, Medicare, Reporting and recordkeeping requirements.

42 CFR Part 486

Grant programs—health, Health facilities, Medicare, Reporting and recordkeeping requirements, X-ray.

42 CFR Part 488

Administrative practice and procedure, Health facilities, Medicare, Reporting and recordkeeping requirements.

42 CFR Part 489

Health facilities, Medicare, Reporting and recordkeeping requirements.

42 CFR Part 495

Administrative practice and procedure, Electronic health records, Health facilities, Health professions, Health maintenance organizations (HMO), Medicaid, Medicare, Penalties, Privacy, Reporting and recordkeeping requirements.

For the reasons set forth in the preamble of this proposed rule, the Centers for Medicare and Medicaid Services is proposing to amend 42 CFR Chapter IV as set forth below:

PART 405—FEDERAL HEALTH INSURANCE FOR THE AGED AND DISABLED

1. The authority citation for part 405 is revised to read as follows:

Authority: Secs. 205(a), 1102, 1142, 1861, 1862(a), 1869, 1871, 1874, 1881, and 1886(k) of the Social Security Act (42 U.S.C. 405(a), 1302, 1320b–12, 1395x, 1395y(a), 1395ff, 1395hh, 1395kk, 1395rr, and 1395ww(k)), and sec. 353 of the Public Health Service Act (42 U.S.C. 263a).

2. Section 405.2404 is amended by revising paragraph (d) introductory text to read as follows:

§ 405.2404 Termination of rural health clinic agreements.

(d) Notice to the public. Prompt notice of the date and effect of termination must be given to the public by either of the following:

3. Section 405.2442 is amended by revising paragraph (a) introductory text and paragraph (b) to read as follows:

§ 405.2442 Notice to the public.

(a) When the FQHC voluntarily terminates the agreement and an effective date is set for the termination, the FQHC must notify the public in the area serviced by the FQHC prior to a prospective effective date or on the actual day that business ceases, if no prospective date of termination has been set. The notice must include—

(b) When CMS terminates the agreement, CMS will notify the public in the area serviced by the FQHC.

PART 412—PROSPECTIVE PAYMENT SYSTEMS FOR INPATIENT HOSPITAL SERVICES

4. The authority citation for part 412 is revised to read as follows:


5. Section 412.22 is amended by revising paragraph (e) introductory text and paragraph (e)(1)(v) introductory text to read as follows:

§ 412.22 Excluded hospitals and hospital units: General rules.

(e) Hospitals-within-hospitals. A hospital-within-a-hospital is a hospital that occupies space in a building also used by another hospital, or in one or more separate buildings located on the same campus as buildings used by another hospital. Prior to October 1, 2017, except as provided in paragraphs (e)(1)(vi) and (l) of this section, a hospital-within-a-hospital must meet the following criteria in order to be excluded from the prospective payment systems specified in §412.1(a)(1). On or
after October 1, 2017, except as provided in paragraphs (e)(1)(vi) and (f) of this section, a hospital-within-hospital that is excluded from the prospective payment systems specified in §412.1(a)(1) that occupies space in a building also used by a hospital which is not excluded from the prospective payment systems specified in §412.1(a)(1), or in one or more separate buildings located on the same campus as buildings used by a hospital not excluded from the prospective payment systems specified in §412.1(a)(1) must meet the following criteria in order to be excluded from the prospective payment systems specified in §412.1(a)(1).  

1. * * * * *  
(v) Performance of basic hospital functions. Prior to October 1, 2017, the hospital meets one of the following criteria:  

* * * * *

6. Section 412.23 is amended by revising paragraphs (e)(2)(ii), (e)(3)(vi), and (e)(7)(iii) and adding paragraph (j) to read as follows:

§ 412.23 Excluded hospitals: Classifications.  

* * * * *  
(e) * * *  
(2) * * *  
(ii) For cost reporting periods beginning on or after August 5, 1997 and on or before December 31, 2014, a hospital that was first excluded from the prospective payment system under this section in 1986 meets the length-of-stay criterion if it has an average inpatient length of stay for all patients, including both Medicare and non-Medicare inpatients, of greater than 20 days and demonstrates that at least 60 percent of its annual Medicare inpatient discharges in the 12-month cost reporting period ending in fiscal year 1997 have a principal diagnosis that reflects a finding of neoplastic disease as defined in paragraph (f)(1)(iv) of this section.  

(2) Payment. Payment for inpatient operating costs for hospitals classified under paragraph (j)(1) of this section is made as set forth in §412.526(c)(3). Payment for capital costs for hospitals classified under paragraph (j)(1) of this section is made as set forth in §412.526(c)(4).

7. Section 412.64 is amended by—  

a. Revising paragraph (d)(1)(vii);  

b. Adding paragraph (d)(4)(iii); and  

c. Revising paragraph (i)(3)(iii).  

The revisions and addition read as follows:

§ 412.64 Federal rates for inpatient operating costs for Federal fiscal year 2005 and subsequent fiscal years.  

* * * * *  
(d) * * *  
(1) * * *  
(vii) For fiscal years 2017 and 2018, the percentage increase in the market basket index (as defined in §13.40(a)(3) of this chapter) for prospective payment hospitals, subject to the provisions of paragraphs (d)(2) and (3) of this section, less a multifactor productivity adjustment (as determined by CMS) and less 0.75 percentage point.  

(4) * * *  
(iii) Exception for decertified EHR technology. Beginning with the fiscal year 2019 payment adjustment year, the Secretary shall exempt an eligible hospital that is not a qualifying eligible hospital from the application of the reduction under paragraph (d)(3) of this section if the Secretary determines that compliance with the requirement for being a meaningful EHR user is not possible because the certified EHR technology used by the eligible hospital has been decertified under ONC’s Health IT Certification Program. To be considered for an exception, an eligible hospital must submit an application, in the manner specified by CMS, demonstrating that the certified EHR technology was decertified during the 12-month period preceding the applicable EHR reporting period for the payment adjustment year, or during the applicable EHR reporting period for the payment adjustment year, and that the eligible hospital made a good faith effort to obtain another certified EHR technology for that EHR reporting period. (See §495.4 for definitions of payment adjustment year, EHR reporting period, and meaningful EHR user.) Applications requesting this exception must be submitted by July 1 of the year before the applicable payment adjustment year, or a later date specified by CMS. This exception is subject to annual renewal, but in no case may an eligible hospital be granted an exception under paragraph (d)(4) of this section for more than 5 years.  

* * * * *  
(i) * * *  
(3) * * *  
(iii) Any wage index adjustment made under this paragraph (i) is effective for a period of 3 fiscal years, except that hospitals in a qualifying county may elect to waive the application of the wage index adjustment. A hospital may waive the application of the wage index adjustment by notifying CMS in writing within 45 days of the date of public display of the annual notice of proposed rulemaking for the hospital inpatient prospective payment system at the Office of the Federal Register.  

* * * * *  

8. Section 412.87 is amended by revising paragraph (b)(2) to read as follows:

§ 412.87 Additional payment for new medical services and technologies: General provisions.  

* * * * *  
(b) * * *  
(2) A medical service or technology may be considered new within 2 or 3 years after the point at which data begin to become available reflecting the inpatient hospital code (as defined in section 1886(d)(5)(K)(iii) of the Social Security Act) assigned to the new service or technology (depending on when a new code is assigned and data on the new service or technology become available for DRG recalibration). After CMS has recalibrated the DRGs, based on available data, to reflect the costs of an otherwise new medical service or technology, the medical service or technology will no longer be
considered "new" under the criterion of this section.

9. Section 412.90 is amended by revising paragraph (j) to read as follows:

§ 412.90 General rules.

(j) Medicare-dependent, small rural hospitals. For cost reporting periods beginning on or after April 1, 1990, and before October 1, 1994, and for discharges occurring on or after October 1, 1997 and before October 1, 2017, CMS adjusts the prospective payment rates for inpatient operating costs determined under subparts D and E of this part if a hospital is classified as a Medicare-dependent, small rural hospital.

10. Section 412.92 is amended by revising paragraph (e)(3) introductory text to read as follows:

§ 412.92 Special treatment: Sole community hospitals.

(e) * * *

(3) Effective for cost reporting periods beginning on or after October 1, 2017, the intermediary determines a lump sum adjustment amount not to exceed the difference between the hospital’s Medicare inpatient operating costs and the hospital’s total DRG revenue for inpatient operating costs based on DRG-adjusted prospective payment rates for inpatient operating costs (including outlier payments for inpatient operating costs determined under subpart F of this part and additional payments made for inpatient operating costs for hospitals that serve a disproportionate share of low-income patients as determined under § 412.106 and for indirect medical education costs as determined under § 412.105). Effective for cost reporting periods beginning on or after October 1, 2017, the MAC determines a lump sum adjustment amount equal to the difference between the hospital’s fixed Medicare inpatient operating costs and the hospital’s total MS–DRG revenue based on MS–DRG-adjusted prospective payment rates for inpatient operating costs (including outlier payments for inpatient operating costs determined under subpart F of this part and additional payments made for inpatient operating costs for hospitals that serve a disproportionate share of low-income patients as determined under § 412.106 and for indirect medical education costs as determined under § 412.105) multiplied by the ratio of the hospital’s fixed Medicare inpatient operating costs to its total Medicare inpatient operating costs.

11. Section 412.101 is amended by revising paragraph (b)(2) introductory text and adding paragraph (e) to read as follows:


* * *

(b) * * *

(2) In order to qualify for this adjustment, a hospital must meet the following criteria, subject to the provisions of paragraph (e) of this section:

(e) Special treatment regarding hospitals operated by the Indian Health Service (IHS) or a Tribe. For discharges occurring in FY 2018 and subsequent fiscal years—

(1) A hospital operated by the IHS or a Tribe will be considered to meet the applicable mileage criterion specified under paragraph (b)(2) of this section if it is located more than the specified number of road miles from the nearest subsection (d) hospital operated by the IHS or a Tribe.

(2) A hospital, other than a hospital operated by the IHS or a Tribe, will be considered to meet the applicable mileage criterion specified under paragraph (b)(2) of this section if it is located more than the specified number of road miles from the nearest subsection (d) hospital other than a subsection (d) hospital operated by the IHS or a Tribe.

12. Section 412.106 is amended by adding paragraph (g)(1)(iii)(C) to read as follows:

§ 412.106 Special treatment: Hospitals that serve a disproportionate share of low-income patients.

* * *

(g) * * *

(1) * * *

(iii) * *

(C) * *

(4) For fiscal year 2018, CMS will base its estimates of the amount of hospital uncompensated care on utilization data for Medicaid and Medicare SSI patients, as determined by CMS in accordance with paragraphs (b)(2)(i) and (4) of this section, using data on Medicaid utilization from 2012 and 2013 cost reports from the most recent HCRIS database extract and 2012 cost report data submitted to CMS by IHS or Tribal hospitals and the most recent available 2 years of data on Medicare SSI hospital utilization (or, for Puerto Rico hospitals, a proxy for Medicare SSI utilization data), and data on uncompensated care costs, defined as charity care costs plus non-Medicare bad debt costs from 2014 cost reports from the most recent HCRIS database extract.

13. Section 412.140 is amended by revising paragraphs (c)(2) and (d)(2) to read as follows:

§ 412.140 Participation, data submission, and validation requirements under the Hospital Inpatient Quality Reporting (IQR) Program.

* * *

(c) * * *

(2) Extraordinary circumstances exceptions. CMS may grant an exception with respect to quality data reporting requirements in the event of extraordinary circumstances beyond the control of the hospital. CMS may grant an exception as follows:

(i) For circumstances not relating to the reporting of electronic clinical quality measure data, a hospital participating in the Hospital IQR Program that wishes to request an exception with respect to quality data reporting requirements must submit its request to CMS within 90 days of the date that the extraordinary circumstances occurred. For circumstances relating to the reporting of electronic clinical quality measures, a hospital participating in the Hospital IQR Program that wishes to request an exception must submit its request to CMS by April 1 following the end of the reporting calendar year in which the extraordinary circumstances occurred. Specific requirements for submission of a request for an exception are available on QualityNet.org.

(ii) CMS may grant an exception to one or more hospitals that have not requested an exception if: CMS determines that a systemic problem with CMS data collection systems directly affected the ability of the hospital to submit data; or if CMS determines that an extraordinary circumstance has affected an entire region or locale.

(d) * * *

(2)(i) A hospital meets the chart-abstracted validation requirement with respect to a fiscal year if it achieves a 75-percent score, as determined by CMS.

(ii) A hospital meets the eCQM validation requirement with respect to a fiscal year if it submits at least 75 percent of sampled eCQM measure medical records in a timely and complete manner, as determined by CMS.
§ 412.211 Puerto Rico rates for Federal fiscal year 2004 and subsequent fiscal years.

(a) * * *

(f) * * *

(3) * * *

(iii) Any wage index adjustment made under this paragraph (f) is effective for a period of 3 fiscal years, except that hospitals in a qualifying county may elect to waive the application of the wage index adjustment. A hospital may waive the application of the wage index adjustment by notifying CMS in writing within 45 days of the date of public display of the annual notice of proposed rulemaking for the hospital inpatient prospective payment system at the Office of the Federal Register.

* * *

§ 412.230 Criteria for an individual hospital seeking redesignation to another rural area or an urban area.

(a) * * *

(3) Special rules for sole community hospitals and rural referral centers. To be redesignated under the special rules in this paragraph, the hospital must submit documentation of the approval of sole community hospital or rural referral center status to the MGCRB no later than the first business day after January 1.

(i) A hospital that is approved as a rural referral center or a sole community hospital, or both, does not have to demonstrate a close proximity to the area to which it seeks redesignation.

(ii) If a hospital that is approved as a rural referral center or a sole community hospital, or both, qualifies for urban redesignation, it is redesignated to the urban area that is closest to the hospital or to the hospital’s geographic home area. If the hospital is closer to another rural area than to any urban area, it may seek redesignation to either the closest rural area or the closest urban area.

* * *

(d) * * *

(3) Rural referral center exceptions.

For the exceptions in this paragraph to apply, the hospital must submit documentation of the approval of rural referral center status to the MGCRB no later than the first business day after January 1.

(i) If a hospital was ever approved as a rural referral center, it does not have to demonstrate that it meets the average hourly wage criterion set forth in paragraph (d)(1)(iii) of this section.

(ii) If a hospital was ever approved as a rural referral center, it is required to meet only the criteria that applies to rural hospitals under paragraph (d)(1)(iv) of this section, regardless of its actual location in an urban or rural area.

* * *

§ 412.273 Withdrawing an application, terminating an approved 3-year classification, or cancelling a previous withdrawal or termination.

(a) * * *

(c) * * *

(1) * * *

(ii) After the MGCRB issues a decision, provided that the request for withdrawal is received by the MGCRB within 45 days of the date of public display of the Office of the Federal Register of CMS’ annual notice of proposed rulemaking concerning changes to the inpatient hospital prospective payment system and proposed payment rates for the fiscal year for which the application has been filed.

* * *

§ 412.500 Basis and scope of subpart.

(a) * * *

(7) Section 411 of Public Law 114–10 which revises the annual update to the LTCH PPS standard Federal payment rate in FY 2018.

(8) Public Law 114–255 which at—

(i) Section 15004 amended the moratorium on increasing beds in existing LTCHs and LTCH satellite facilities and amended high cost outlier payment requirements;

(ii) Section 15006 amended moratoria on certain payment policies;

(iii) Section 15007 amended the average length of stay requirements;

(iv) Section 15009 temporarily excepted certain spinal cord specialty hospitals from the site neutral payment rate; and

(v) Section 15010 temporarily excepted certain wound care discharge from certain LTCHs from the site neutral payment rate.

* * *

§ 412.522 Application of site neutral payment rate.

(a) * * *

(b) * * *

(3) Temporary exception for certain severe wound discharges.—(i) Definitions. For purposes of this paragraph (b)(3) the following definitions are applicable:

Severe wound means a wound which is a stage 3 wound, stage 4 wound, unstageable wound, non-healing surgical wound, fistula, as identified by the applicable code on the claim from the long-term care hospital.

Wound means an injury, usually involving division of tissue or rupture of the integument or mucous membrane with exposure to the external environment.

(ii) Discharges for severe wounds. A discharge that occurs in a cost reporting period beginning during fiscal year 2018 for a patient who was treated for a severe wound that meets all of the following criteria is excluded from the site neutral payment rate specified under this section:

(A) The severe wound meets the definition specified in paragraph (b)(3)(i) of this section.

(B) The discharge is from a long-term care hospital that is described in § 412.230(e)(2)(i) and meets the criteria of § 412.22(f); and

(C) The discharge is classified under MS–LTC–DRG 539, 540, 602, or 603.

(4) Temporary exception for certain spinal cord specialty hospitals. For discharges in cost reporting periods beginning in fiscal years 2018 and 2019, the site neutral payment rate specified under this section does not apply if such discharge is from a long-term care hospital that meets each of the following requirements:

(i) The hospital was a not-for-profit long-term care hospital on June 1, 2014, as determined by cost report data;

(ii) Of the discharges in calendar year 2013 from the long-term care hospital for which payment was made under subpart Q, at least 50 percent were classified under MS–LTC–DRGs 28, 29, 52, 57, 551, 573, and 963; and

(iii) The long-term care hospital discharged inpatients (including both individuals entitled to, or enrolled for, benefits under Medicare Part A and individuals not so entitled or enrolled) during fiscal year 2014 who had been admitted from at least 20 of the 50

* * *
States determined by the States of residency of such inpatients.

§ 412.523 Methodology for calculating the Federal prospective payment rates.

(c) * * * * * * * * *
(d) * * * * * * * * *

(1) Outlier payments. CMS adjusts the LTCH PPS standard Federal payment rate by a reduction factor of 8 percent, the estimated proportion of outlier payments under § 412.525(a) payable for discharges described in § 412.522(a)(2) (notwithstanding the provisions of § 412.525(a)(2)) for FY 2018 and subsequent years.

(5) Adjustment for changes to the short-stay outlier policy. The standard Federal rate determined under paragraph (c)(3) of this section is permanently adjusted by a one-time factor so that estimated aggregate payments to LTCH PPS standard Federal rate cases in FY 2018 are projected to equal estimated aggregate payments that would have been paid for such cases without regard to the change in the short-stay outlier policy for FY 2018 under § 412.529(c)(5).

§ 412.525 Adjustments to the Federal prospective payment.

(a) * * *
(b) * * *
(c) * * *

(2)(i) The fixed loss-amount for discharges from a long-term care hospital described under § 412.522(a)(2) is determined for the long-term care hospital prospective payment system payment year, using the LTC–DRG relative weights that are in effect at the start of the applicable long-term care hospital prospective payment system payment year.

(ii) For FY 2018 and subsequent years, the fixed-loss amount for long-term care hospital discharges described under § 412.522(a)(2) is determined such that the estimated proportion of outlier payments under paragraph (a) of this section payable for such discharges is projected to be equal to 99.6875 of 8 percent.

§ 412.525(a)(2)(ii) for FY 2018 and subsequent years.

§ 412.529 Special payment provision for short-stay outliers.

(c) * * * * * * *

(3) Discharges occurring on or after July 1, 2007 and before December 29, 2007 and discharges occurring on or after December 29, 2012 and on or before September 30, 2017. For discharges from long-term care hospitals described under § 412.23(e)(2)(i) occurring on or after July 1, 2007, and on or before December 29, 2007 and discharges occurring on or after December 29, 2012, and on or before September 30, 2017, the LTCH prospective payment system adjusted payment amount for a short-stay outlier case is adjusted by either of the following:

(4) Discharges occurring on or after October 1, 2017. For discharges occurring on or after October 1, 2017, short-stay outlier payments are determined according to paragraph (c)(2)(iv) of this section.

§ 412.23(e)(2)(i) for FY 2018 and subsequent years.

§ 412.23(e), except as specified in paragraph (a)(2) of this section, effective for discharges occurring on or after October 1, 2018.

§ 412.560 Requirements under the Long-Term Care Hospital Quality Reporting Program (LTCH QRP).

(a) Participation in the LTCH QRP. A long-term-care hospital must begin submitting data on measures specified under sections 1886(m)(5)(D), 1899B(c)(1) and 1899B(d)(1) of the Act, and standardized patient assessment data required under section 1899B(b)(1) of the Act, under the LTCH QRP by no later than the first day of the calendar quarter subsequent to 30 days after the date on its CMS Certification Number (CCN) notification letter.

(b) Data submission requirements and payment impact. (1) Except as provided in paragraph (c) of this section, a long-term care hospital must submit to CMS data on measures specified under sections 1886(m)(5)(D), 1899B(c)(1) and 1899B(d)(1) of the Act, and standardized patient assessment data required under section 1899B(b)(1) of the Act. Such data must be submitted in a form and manner, and at a time, specified by CMS.

(c) Exception and extension request requirements. Upon request by a long-term care hospital, CMS may grant an exception or extension with respect to the measures data and standardized patient assessment data reporting requirements, for one or more quarters, in the event of certain extraordinary circumstances beyond the control of the long-term care hospital, subject to the following:

(1) A long-term care hospital that wishes to request an exception or extension with respect to the measures data and standardized patient assessment data reporting requirements must submit its request to CMS within 90 days of the date that the extraordinary circumstances occurred.

(3) * * * * *

(vii) The date on which the long-term care hospital will be able to again submit measures data and standardized patient assessment data under the LTCH QRP and a justification for the proposed date.
(ii) A systemic problem with one of CMS’ data collection systems directly affected the ability of the long-term care hospital to submit measures data and standardized patient assessment data.

* * * * *

(d) * * *

(1) Written notification of noncompliance decision. CMS will send a long-term care hospital written notification of a decision of noncompliance with the measures data and standardized patient assessment data reporting requirements for a particular fiscal year. CMS also will use the Quality Improvement and Evaluation system (QIES) Assessment Submission and Processing (ASAP) System to provide notification of noncompliance to the long-term care hospital.

(2) * * *

(vii) Accompanying documentation that demonstrates compliance of the long-term care hospital with the LTCH QRP requirements. This documentation must be submitted electronically at the same time as the reconsideration request as an attachment to the email.

* * * * *

(f) Data completion thresholds. (1) Long-term care hospitals must meet or exceed two separate data completeness thresholds: One threshold set at 80 percent for completion of measures data and standardized patient assessment data collected using the LTCH CARE Data Set submitted through the QIES ASAP System; and a second threshold set at 100 percent for completion of measures data collected using the CDC NHSN.

(2) The thresholds in paragraph (f)(1) of this section apply to all data that must be submitted under paragraph (b) of this section.

(3) A long-term care hospital must meet or exceed both thresholds in paragraph (f)(1) of this section to avoid receiving a 2 percentage point reduction to its annual payment update for a given fiscal year, beginning with the FY 2019 LTCH QRP.

PART 413—PRINCIPLES OF REASONABLE COST REIMBURSEMENT; PAYMENT FOR END-STAGE RENAL DISEASE SERVICES; OPTIONAL PROSPECTIVELY DETERMINED PAYMENT RATES FOR SKILLED NURSING FACILITIES; PAYMENT FOR ACUTE KIDNEY INJURY DIALYSIS

24. The authority citation for part 413 is revised to read as follows:

Authority: Secs. 1102, 1812(d), 1814(b), 1815, 1823(a), (f), and (n), 1861(v), 1871, 1881, 1883 and 1886 of the Social Security Act (42 U.S.C. 1302, 1395(d), 1395(b), 1395g, 1395l(a), (i), and (n), 1395x(v), 1395hh, 1395rr, 1395tt, and 1395ww); and sec. 124 of Public Law 106–113, 113 Stat. 1501A–332; sec. 3201 of Public Law 112–96, 126 Stat. 156; sec. 632 of Public Law 112–240, 126 Stat. 3254; sec. 217 of Public Law 113–93, 120 Stat. 1040; and sec. 204 of Public Law 113–295, 128 Stat. 4010; and sec. 808 of Public Law 114–27, 120 Stat. 362.

25. Section 413.24 is amended by revising paragraph (f)(4)(iv) to read as follows:

§ 413.24 Adequate cost data and cost finding.

* * * * *

(f) * * *

(iv)(A) Effective as specified in paragraphs (f)(4)(iv)(A)(1) through (4) and except as provided in paragraph (f)(4)(iv)(C) of this section, a provider must submit a hard copy of a settlement summary, if applicable, which is a statement of certain worksheet totals found within the electronic file, and the certification statement described in paragraph (f)(4)(iv)(B) of this section signed by its administrator or chief financial officer certifying the accuracy of the electronic file or the manually prepared cost report.

(1) For hospitals, effective for cost reporting periods ending on or after September 30, 1994;

(2) For skilled nursing facilities and home health agencies, effective for cost reporting periods ending on or after February 1, 1997;

(3) For hospices and end-stage renal disease facilities, effective for cost reporting periods ending on or after December 31, 2004; and

(4) For organ procurement organizations, histocompatibility laboratories, rural health clinics, Federally qualified health centers, and community mental health centers, effective for cost reporting periods ending on or after March 31, 2005.

(B) The following certification statement must immediately precede the dated original signature, or electronic signature as set forth in paragraph (f)(4)(iv)(C)(1) of this section, of the provider’s administrator or chief financial officer:

MISREPRESENTATION OR FALSEIFICATION OF ANY INFORMATION CONTAINED IN THIS COST REPORT MAY BE PUNISHABLE BY CRIMINAL, CIVIL AND ADMINISTRATIVE ACTION, FINE AND/OR IMPRISONMENT UNDER FEDERAL LAW. FURTHERMORE, IF SERVICES IDENTIFIED IN THIS REPORT WERE PROVIDED OR PROCURED THROUGH THE PAYMENT DIRECTLY OR INDIRECTLY OF A KICKBACK OR WERE OTHERWISE ILLEGAL, CRIMINAL, CIVIL AND ADMINISTRATIVE ACTION, FINES AND/OR IMPRISONMENT MAY RESULT.

I hereby certify that I have read the above certification statement and that I have examined the accompanying electronically filed or manually submitted cost report and the Balance Sheet and Statement of Revenue and Expenses prepared by [Provider Name(s) and Number(s)] for the cost reporting period beginning [begin and ending ] and that to the best of my knowledge and belief, this report and statement are true, correct, complete and prepared from the books and records of the provider in accordance with applicable instructions, except as noted. I further certify that I am familiar with the laws and regulations regarding the provision of health care services, and that the services identified in this cost report were provided in compliance with such laws and regulations.

(C) Effective for cost reporting periods beginning on or after October 1, 2017—

(1) A provider that is required to file an electronic cost report may elect to electronically submit the settlement summary, if applicable, and the certification statement with an electronic signature of the provider’s administrator or chief financial officer. The following checkbox for electronic signature and submission will immediately follow the certification statement as set forth in paragraph (f)(4)(iv)(B) of this section and must be checked if electronic signature and submission is elected.

☐ I have read and agree with the above certification statement. I certify that I intend my electronic signature on this certification statement to be the legally binding equivalent of my original signature.

(2) A provider that is required to file an electronic cost report but does not elect to electronically submit the certification statement with an electronic signature, must submit a hard copy of the settlement summary, if applicable, and a certification statement with an original signature of the provider’s administrator or chief financial officer as set forth in paragraphs (f)(4)(iv)(A) and (B) of this section.

* * * * *

26. Section 413.65 is amended by revising paragraph (m) introductory text to read as follows:
§ 413.65 Requirements for a determination that a facility or an organization has provider-based status.

(a) * * * *

(m) Status of Indian Health Service and Tribal facilities and organizations. Facilities and organizations operated by the Indian Health Services and Tribes will be considered to be departments of hospitals operated by the Indian Health Service or Tribes if they furnish only services that are billed, using the CCN of the main provider and with the consent of the main provider, as if they had been furnished by a department of a hospital operated by the Indian Health Service or a Tribe and they are:

* * * *

27. Section 413.70 is amended by—

a. Redesignating paragraph (a)(6)(iii) as paragraph (a)(6)(iv); and

b. Adding a new paragraph (a)(6)(iii); and

c. Revising newly redesignated paragraph (a)(6)(iv).

The addition and revision read as follows:

§ 413.70 Payment for services of a CAH.

(a) * * * *

(iii) Exception for decertified EHR technology. Beginning with the fiscal year 2018 payment adjustment year, the Secretary shall exempt a CAH that is not a qualifying CAH from the application of the payment adjustment under paragraph (a)(6)(i) of this section if the Secretary determines that compliance with the requirement for being a meaningful EHR user is not possible because the certified EHR technology used by the CAH has been decertified under ONC’s Health IT Certification Program. In order to be considered for an exception, a CAH must submit an application, in the manner specified by CMS, demonstrating that the certified EHR technology was decertified during the 12-month period preceding the applicable EHR reporting period for the payment adjustment year, or during the applicable EHR reporting period for the payment adjustment year, and that the CAH made a good faith effort to obtain another certified EHR technology for that EHR reporting period. Applications requesting this exception must be submitted by November 30 after the end of the applicable payment adjustment year, or a later date specified by CMS.

(iv) Exceptions granted under paragraphs (a)(6)(ii) and (iii) of this section are subject to annual renewal, but in no case may a CAH be granted such an exception for more than 5 years.

* * * *

§ 414.134 Depreciation: Allowance for depreciation based on asset costs.

(f) * * * *

1. General. Depreciable assets may be disposed of through sale, scrapping, trade-in, exchange, demolition, abandonment, condemnation, fire, theft, or other casualty.

(ii) Disposal of an asset before December 1, 1997. If disposal of a depreciable asset, including the sale or scrapping of an asset before December 1, 1997, results in a gain or loss, an adjustment is necessary in the provider’s allowable cost.

(A) The amount of a gain included in the determination of allowable cost is limited to the amount of depreciation previously included in Medicare allowable costs.

(B) The amount of a loss to be included is limited to the undepreciated basis of the asset permitted under the program.

(C) The treatment of the gain or loss depends upon the manner of disposition of the asset, as specified in paragraphs (f)(2) through (6) of this section.

(ii) Disposal of an asset on or after December 1, 1997. No gain or loss is recognized on either the sale or scrapping of an asset that occurs on or after December 1, 1997, regardless of whether the asset is sold incident to a provider’s change of ownership, or otherwise sold or scrapped as an asset of a Medicare participating provider. Gains or losses on dispositions other than sales or scrapping are recognized to the same extent as prior to December 1, 1997.

* * * *

PART 414—PAYMENT FOR PART B MEDICAL AND OTHER HEALTH SERVICES

29. The authority citation for part 414 continues to read as follows:

Authority: Secs. 1102, 1871, and 1881(b)(1) of the Social Security Act (42 U.S.C. 1302, 1320b–8, and 1395hh) and section 371 of the Public Health Service Act (42 U.S.C. 273).

30. Section 414.68 is amended by redesignating paragraphs (c)(7) through (14) as paragraphs (c)(8) through (15), respectively, and adding new paragraph (c)(7) to read as follows:

§ 414.68 Imaging accreditation.

* * * *

(c) * * * *

7. A statement acknowledging that the organization agrees to make all Medicare final accreditation survey reports (including statements of deficiencies) and acceptable plans of correction publicly available on the organization’s Web site within 90 days after such information is made available to those facilities for the most recent 3 years, on an ongoing basis. This acknowledgement includes all full, follow-up, focused, and complaint surveys, regardless of whether they are performed onsite or offsite.

PART 416—AMBULATORY SURGICAL SERVICES

31. The authority citation for part 416 is revised to read as follows:

Authority: Secs. 1102, 1138, and 1871 of the Social Security Act (42 U.S.C. 1302, 1320b–8, and 1395hh) and section 371 of the Public Health Service Act (42 U.S.C. 273).

32. Section 416.35 is amended by revising paragraph (d) introductory text to read as follows:

§ 416.35 Termination of agreement.

* * * *

(d) Notice to the public. Prompt notice of the date and effect of termination is given to the public by—

* * * *

PART 486—CONDITIONS FOR COVERAGE OF SPECIALIZED SERVICES FURNISHED BY SUPPLIERS

33. The authority citation for part 486 continues to read as follows:

Authority: Secs. 1102, 1138, and 1871 of the Social Security Act (42 U.S.C. 1302, 1320b–8, and 1395hh) and section 371 of the Public Health Service Act (42 U.S.C. 273).

34. Section 486.312 is amended by revising paragraph (e) to read as follows:

§ 486.312 De-certification.

* * * *

(e) Public notice. Once CMS approves the date for a voluntary termination, the OPO must provide prompt public notice in the service area of the date of de-certification and such other information as CMS may require. In the case of involuntary termination or nonrenewal of an agreement, CMS also provides notice to the public in the service area of the date of de-certification. No payment under titles XVIII or XIX of the Act will be made with respect to organ procurement costs attributable to the OPO on or after the effective date of de-certification.
PART 488—SURVEY, CERTIFICATION, AND ENFORCEMENT PROCEDURES

35. The authority citation for part 488 is revised to read as follows:

Authority: Secs. 1102, 1128l, 1864, 1865, 1871, and 1875 of the Social Security Act, unless otherwise noted (42 U.S.C. 1302, 1320a–7, 1395aa, 1395bb, 1395hh, and 1395ll).

36. Section 488.5 is amended by adding paragraph (a)(21) to read as follows:

§ 488.5 Application and re-application procedures for national accrediting organizations.

(a) * * *

(21) A statement acknowledging that the organization agrees to make all Medicare final accreditation survey reports (including statements of deficiencies) and acceptable plans of correction publicly available on the organization’s Web site within 90 days after such information is made available to those facilities for the most recent 3 years, on an ongoing basis. This acknowledgement includes all triennial, full, follow-up, focused, and complaint surveys, regardless of whether they are performed onsite or offsite.

* * * * *

PART 489—PROVIDER AGREEMENTS AND SUPPLIER APPROVAL

37. The authority citation for part 489 continues to read as follows:

Authority: Secs. 1102, 1128l, 1820(E), 1861, 1864(M), 1866, 1869, and 1871 of the Social Security Act (42 U.S.C. 1302, 1395i–3, 1395x, 1395aa(m), 1395cc, 1395ff, and 1395hh).

38. Section 489.52 is amended by revising paragraph (c)(2) introductory text to read as follows:

§ 489.52 Termination by the provider.

* * * * *

(c) * * *

(2) The notice must—

* * * * *

PART 495—STANDARDS FOR THE ELECTRONIC HEALTH RECORD TECHNOLOGY INCENTIVE PROGRAM

39. The authority citation for part 495 continues to read as follows:

Authority: Secs. 1102 and 1871 of the Social Security Act (42 U.S.C. 1302 and 1395hh).

40. Section 495.4 is amended by—

a. Adding in alphabetical order a definition of “Ambulatory surgical center-based EP”;

b. In the definition of “EHR reporting period,” revising paragraph (1)(ii) introductory text, adding paragraph (1)(i)(D), revising paragraph (1)(iii) introductory text, revising paragraph (2)(i) introductory text, adding paragraph (2)(i)(D) and revising paragraph (2)(iii) introductory text.

c. In the definition of “EHR reporting period for a payment adjustment year,” revising paragraph (2)(ii) introductory text, adding paragraph (2)(ii)(D), revising paragraph (2)(ii) introductory text, revising paragraph (3)(ii) introductory text, adding paragraph (3)(ii)(D), and revising paragraph (3)(iii) introductory text.

The additions and revisions read as follows:

§ 495.4 Definitions.

* * * * *

Ambulatory surgical center-based EP means an EP who furnishes 75 percent or more of his or her covered professional services in sites of service identified by the codes used in the HIPAA standard transaction as an ASC setting in the calendar year that is 2 years before the payment adjustment year.

* * * * *

EHR reporting period. * * *

(1) * * *

(ii) The following are applicable for 2015, 2016, 2017, and 2018:

* * * * *

(D) For the CY 2018 payment year under the Medicaid EHR Incentive Program:

(1) For the EP first demonstrating he or she is a meaningful EHR user, any continuous 90-day period within CY 2018.

(2) For the EP who has successfully demonstrated he or she is a meaningful EHR user in any prior year, any continuous 90-day period within CY 2018.

(iii) The following are applicable beginning within CY 2018:

* * * * *

(2) * * *

(ii) The following are applicable for 2015, 2016, 2017, and 2018:

* * * * *

(D) For the CY 2018 payment year under the Medicaid EHR Incentive Program:

(1) For the eligible hospital or CAH first demonstrating it is a meaningful EHR user, any continuous 90-day period within CY 2018.

(2) For the eligible hospital or CAH that has successfully demonstrated it is a meaningful EHR user in any prior year, any continuous 90-day period within CY 2018.

(iii) The following are applicable beginning with the FY 2019 payment year under the Medicaid EHR Incentive Program:

* * * * *

EHR reporting period for a payment adjustment year. * * *

(2) * * *

(ii) The following are applicable for 2015, 2016, 2017, and 2018:

* * * * *

(D) In 2018 as follows:

(1) If an eligible hospital has not successfully demonstrated it is a meaningful EHR user in a prior year, the EHR reporting period is any continuous 90-day period within CY 2018 and applies for the FY 2019 and 2020 payment adjustment years. For the FY 2019 payment adjustment year, the EHR reporting period must end before and the eligible hospital must successfully register for and attest to meaningful use no later than October 1, 2018.

(2) If in a prior year an eligible hospital has successfully demonstrated it is a meaningful EHR user, the EHR reporting period is any continuous 90-day period within CY 2018 and applies for the FY 2020 payment adjustment year.

(iii) The following are applicable beginning in 2019:

* * * * *

(3) * * *

(ii) The following are applicable for 2015, 2016, 2017, and 2018:

* * * * *

(D) In 2018 as follows:

(1) If a CAH has not successfully demonstrated it is a meaningful EHR user in a prior year, the EHR reporting period is any continuous 90-day period within CY 2018 and applies for the FY 2018 payment adjustment year.

(2) If in a prior year a CAH has successfully demonstrated it is a meaningful EHR user, the EHR reporting period is any continuous 90-day period within CY 2018 and applies for the FY 2018 payment adjustment year.

(iii) The following are applicable beginning in 2019:

* * * * *

41. Section 495.102 is amended by redesignating paragraph (d)(5) as paragraph (d)(6) and adding new paragraphs (d)(5) and (7) to read as follows:

§ 495.102 Incentive payments to EPs.

* * * * *

(d) * * *

(5) Exception for decertified EHR technology. The Secretary shall exempt an EP from the application of the payment adjustment for CY 2018 under paragraph (d)(1) of this section if the Secretary determines that compliance with the requirement for being a
meaningful EHR user is not possible because the certified EHR technology used by the EP has been decertified under ONC’s Health IT Certification Program. To be considered for an exception, an EP must submit, in the manner specified by CMS, an application demonstrating that the certified EHR technology was decertified during the 12-month period preceding the applicable EHR reporting period for the CY 2018 payment adjustment year, or during the applicable EHR reporting period for the CY 2018 payment adjustment year, and that the EP made a good faith effort to obtain another certified EHR technology for that EHR reporting period.

Applications requesting this exception must be submitted no later than October 1, 2017, or a later date specified by CMS.

* * * * *

(7) Payment adjustments not applicable to ambulatory surgical center-based EPs. For the CY 2017 and CY 2018 payment adjustment years, no payment adjustment under paragraphs (d)(1) through (3) of this section may be made in the case of an ambulatory surgical center-based eligible professional, as defined in §495.4.


Seema Verma,
Administrator, Centers for Medicare & Medicaid Services.

Dated: April 11, 2017

Thomas E. Price,
Secretary, Department of Health and Human Services.

Note: The following Addendum and Appendixes will not appear in the Code of Federal Regulations.

Addendum—Proposed Schedule of Standardized Amounts, Update Factors, Rate-of-Increase Percentages Effective With Cost Reporting Periods Beginning on or After October 1, 2017, and Payment Rates for LTCHs Effective for Discharges Occurring on or After October 1, 2017

I. Summary and Background

In this Addendum, we are setting forth a description of the methods and data we used to determine the proposed prospective payment rates for Medicare hospital inpatient operating costs and Medicare hospital inpatient capital-related costs for FY 2018 for acute care hospitals. We also are setting forth the rate-of-increase percentage for updating the target amounts for certain hospitals excluded from the IPPS for FY 2018. We note that, because certain hospitals excluded from the IPPS are paid on a reasonable cost basis subject to a rate-of-increase ceiling (and not by the IPPS), these hospitals are not affected by the proposed updates for the standardized amounts, offsets, and budget neutrality factors. Therefore, in this proposed rule, we are setting forth the rate-of-increase percentage for updating the target amounts for certain hospitals excluded from the IPPS that would be effective for cost reporting periods beginning on or after October 1, 2017.

In addition, we are setting forth a description of the methods and data we used to determine the proposed standard Federal payment rate that would be applicable to Medicare LTCHs for FY 2018.

In general, except for SCHs, for FY 2018, each hospital’s payment per discharge under the IPPS is based on 100 percent of the Federal national rate, also known as the national adjusted standardized amount. This amount reflects the national average hospital cost per case from a base year, updated for inflation. We note that, under section 205 of the Medicare Access and CHIP Reauthorization Act of 2015 (MACRA) (Pub. L. 114–10), enacted on April 16, 2015, the MDH program is set to expire at the end of FY 2017.

SCHs are paid based on whichever of the following rates yields the greatest aggregate payment: The Federal national rate (including, as discussed in section V.G. of the preamble of this proposed rule, uncompensated care payments under section 1886(i)(2) of the Act); the updated hospital-specific rate based on FY 1982 costs per discharge; the updated hospital-specific rate based on FY 1987 costs per discharge; the updated hospital-specific rate based on FY 1996 costs per discharge; or the updated hospital-specific rate based on FY 2006 costs per discharge. As noted, under current law, the MDH program is set to expire at the end of FY 2017.

As discussed in section V.B. of the preamble of this proposed rule, in accordance with section 1886(d)(9)(E) of the Act as amended by section 601 of the Consolidated Appropriations Act, 2016 (Pub. L. 114–113), for FY 2018, subsection (d) of the IPPS is set to expire at the end of FY 2017.

As discussed in section V.B. of the preamble of this proposed rule, in accordance with section 1886(d)(9)(E) of the Act as amended by section 601 of the Consolidated Appropriations Act, 2016 (Pub. L. 114–113), for FY 2018, subsection (d) of the IPPS is set to expire at the end of FY 2017. Because Puerto Rico hospitals are paid 100 percent of the national standardized amount and are subject to the same national standardized amount as subsection (d) of the IPPS that receive the full update, our discussion below does not include references to the Puerto Rico standardized amount or the Puerto Rico-specific wage index.

As discussed in section II. of this Addendum, we are proposing to make changes in the determination of the prospective payment rates for Medicare inpatient operating costs for acute care hospitals for FY 2018. In section III. of this Addendum, we discuss our proposed policy changes for determining the prospective payment rates for Medicare inpatient capital-related costs for FY 2018. In section IV. of this Addendum, we are setting forth the rate-of-increase percentage for determining the rate-of-increase limits for certain hospitals excluded from the IPPS for FY 2018. In section V. of this Addendum, we discuss proposed policy changes for determining the standard Federal rate for LTCHs paid under the LTCH PPS for FY 2018. The tables to which we refer in the preamble of this proposed rule are listed in section VI. of this Addendum and are available via the Internet on the CMS Web site.

II. Proposed Changes to Prospective Payment Rates for Hospital Inpatient Operating Costs for Acute Care Hospitals for FY 2018

The basic methodology for determining prospective payment rates for hospital inpatient operating costs for acute care hospitals for FY 2005 and subsequent fiscal years is set forth under §412.64. The basic methodology for determining the prospective payment rates for hospital inpatient operating costs for hospitals located in Puerto Rico for FY 2005 and subsequent fiscal years is set forth under §§412.211 and 412.212. Below we discuss the factors we are proposing to use for determining the proposed prospective payment rates for FY 2018.

In summary, the proposed standardized amounts set forth in Tables 1A, 1B, and 1C that are listed and published in section VI. of this Addendum (and available via the Internet on the CMS Web site) reflect—

• Equalization of the standardized amounts for urban and other areas at the level computed for large urban hospitals during FY 2004 and onward, as provided for under section 1886(d)(3)[A][iv][II] of the Act.
• The labor-related share that is applied to the standardized amounts to give the hospital the highest payment, as provided for under sections 1886(d)(3)[E] and 1886(d)(9)[C][iv] of the Act. For FY 2018, depending on whether a hospital submits quality data under the rules established in accordance with section 1886(b)[3][B][viii] of the Act (hereafter referred to as a hospital that submits quality data) and is a meaningful EHR user under section 1886(b)[3][B][ix] of the Act (hereafter referred to as a hospital that is a meaningful EHR user), there are four possible applicable percentage increases that can be applied to the national standardized amount. We refer readers to section V.B. of the preamble of this proposed rule for a complete discussion on the proposed FY 2018 inpatient hospital update. Below is a table with these four options:
We note that section 1886(b)(3)(B)(viii) of the Act, which specifies the adjustment to the applicable percentage increase for “subsection (d)” hospitals that do not submit quality data under the rules established by the Secretary, is not applicable to hospitals located in Puerto Rico.

In addition, section 602 of Public Law 114–113 amended section 1886(n)(6)(B) of the Act to specify that Puerto Rico hospitals are eligible for incentive payments for the meaningful use of certified EHR technology, effective beginning FY 2016, and also to apply the adjustments to the applicable percentage increase under section 1886(b)(3)(B)(ix) of the Act to Puerto Rico hospitals that are not meaningful EHR users, effective FY 2022. Accordingly, because the provisions of section 1886(b)(3)(B)(ix) of the Act are not applicable to hospitals located in Puerto Rico until FY 2022, the adjustments under this provision are not applicable for FY 2018.

- An adjustment to the standardized amount to ensure budget neutrality for DRG recalibration and reclassification, as provided for under section 1886(d)(4)(C)(iii) of the Act.
- An adjustment to ensure the wage index and labor-related share changes are budget neutral, as provided for under section 1886(d)(3)(B)(iii) of the Act (as discussed in the FY 2006 IPPS final rule (70 FR 47395) and the FY 2010 IPPS final rule (74 FR 44005). We note that section 1886(d)(3)(B)(i) of the Act requires that when we compute such budget neutrality, we assume that the provisions of section 1886(d)(3)(B)(i) of the Act (requiring a 62 percent labor-related share in certain circumstances) had not been enacted.
- An adjustment to ensure the effects of geographic reclassification are budget neutral, as provided for under section 1886(d)(3)(B)(ii) of the Act, by removing the FY 2017 budget neutrality factor and applying a revised factor.
- Removal of the adjustment in FY 2017 to offset the cost of the 3-year hold harmless transitional wage index provisions provided by CMS as a result of the implementation of the new OMB labor market area delineations (beginning with FY 2015).
- A single positive adjustment of 0.4588 in FY 2018 as required under section 15003 of the 21st Century Cures Act (Pub. L. 114–255), which amended section 7(b)(1)(B) of the TMA, as amended by section 631 of the ATRA and section 414 of the MACRA, to reduce the adjustment for FY 2018 from 0.5 percentage point to 0.4588 percentage point.
- An adjustment to remove the FY 2017 outlier offset and apply an offset for FY 2018, as provided for in section 1886(d)(3)(B) of the Act.
- As discussed in section V.M. of the preamble of this proposed rule, a factor of (1/1.006) in the calculation of the FY 2018 standardized amount. Specifically, in the FY 2017 IPPS/LTCPPS final rule (81 FR 57058 through 57060), using our authority under section 1886(d)(3)(B)(i) of the Act, we finalized a policy to include a permanent factor of (1/0.998) and a temporary one-time factor of (1.006) in the calculation of the FY 2017 standardized amount and to include a factor of (1/1.006) in the calculation of the FY 2018 standardized amount to remove the temporary one-time factor of 1.006 applied in FY 2017 to address the effects of the 0.2 percent reduction to the rate for the 2-midnight policy in effect for FY 2014, FY 2015, and FY 2016. Therefore, in this proposed rule, for FY 2018, we are removing the temporary one-time prospective increase to the FY 2017 standardized amount of 0.6 percent or a factor of 1.006.

For FY 2018, consistent with current law, we are proposing to apply the rural floor budget neutrality adjustment to hospital wage indexes. Also, consistent with section 3141 of the Affordable Care Act, instead of applying a State-level rural floor budget neutrality adjustment to the wage index, we are proposing to apply a uniform, national budget neutrality adjustment to the FY 2018 wage index for the rural floor. We note that, in section III.H.2.b. of the preamble to this proposed rule, the imputed floor is set to expire effective October 1, 2017, and we are not proposing to extend the imputed floor policy.

In prior fiscal years, CMS made an adjustment to ensure the effects of the rural community hospital demonstration program required under section 410A of Public Law 108–173, as amended by sections 3123 and 10313 of Public Law 111–148, which extended the demonstration program for an additional 5 years (FYs 2011 through 2015), were budget neutral, as provided under section 410Ac(2) of Public Law 108–173. As discussed in section V.L.3. of the preamble to this proposed rule, section 15003 of Public Law 114–255 amended section 410A of Public Law 108–173 to provide for a 10-year extension of the demonstration (in place of the 5-year extension required by the Affordable Care Act) beginning on the date immediately following the last day of the initial 5-year period under section 410A(a)(5) of Public Law 114–255. Thus, section 15003 of Public Law 114–255 requires an additional 5-year extension of the demonstration.

Regarding the costs of the demonstration specifically for FY 2018, as described in section V.L.3. of the preamble to this proposed rule, we are proposing that if the selection of additional hospitals pursuant to section 410Ag(b)(6) of Public Law 108–173 (as added by section 15003 of Public Law 114–255) is announced by June 2017, we would include in the FY 2018 IPPS/LTCPPS final rule an estimate of the costs of the demonstration for FY 2018 and the resulting budget neutrality offset amount for the newly selected hospitals (Cohort 3 hospitals) and for the previously participating hospitals (Cohorts 1 and 2 hospitals). If the final selection of the additional hospitals is not announced by June 2017, we would not be able to include an estimate of the costs of the demonstration for any participating hospitals or an estimated budget neutrality adjustment for FY 2018 in the FY 2018 IPPS/LTCPPS final rule. We refer the reader to section V.L.3. of the preamble to this proposed rule for complete details on the rural community hospital demonstration program and our proposed methodology for calculating budget neutrality for this demonstration.

A. Calculation of the Proposed Adjusted Standardized Amount

1. Standardization of Base-Year Costs or Target Amounts

In general, the national standardized amount is based on per discharge averages of adjusted hospital costs from a base period (section 1886(d)(2)(A) of the Act), updated and otherwise adjusted in accordance with the provisions of section 1886(d) of the Act. The September 1, 1983 interim final rule (48 FR 39763) contained a detailed explanation of how base-year cost data (from cost reporting periods ending during FY 1981) were established for urban and rural hospitals in the initial development of standardized amounts for the IPPS. Sections 1886(d)(2)(B) and 1886(d)(2)(C) of the Act require us to update base-year per discharge costs for FY 1984 and then standardize the cost data in order to remove the effects of certain sources of cost variations among hospitals. These effects include case-mix, differences in average wage levels, cost-of-living adjustments for Alaska.
and Hawaii, IME costs, and costs to hospitals serving a disproportionate share of low-income patients.

For FY 2018, we are proposing to rebase and revise the national labor-related and nonlabor-related shares (based on the proposed 2014-based hospital market basket discussed in section IV. of the preamble of this proposed rule). Specifically, under section 1886(d)(3)(E) of the Act, the Secretary estimates, from time to time, the proportion of payments that are labor-related and adjusts the proportion as determined by the Secretary (from time to time) of hospitals’ costs which are attributable to wages and wage-related costs of the DRG prospective payment rates. We refer to the proportion of hospitals’ costs that are attributable to wages and wage-related costs as the “labor-related share.” For FY 2018, as discussed in section IV.B.3 of the preamble of this proposed rule, we are proposing to apply a labor-related share of 68.3 percent for the national standardized amount for all IPPS hospitals (including hospitals in Puerto Rico) that have a wage index value that is greater than 1.0000. Consistent with section 1886(d)(3)(E) of the Act, we are proposing to apply the wage index to a labor-related share of 62 percent of the national standardized amount for all IPPS hospitals (including hospitals in Puerto Rico) whose wage index values are less than or equal to 1.0000.

The proposed standardized amounts for operating costs appear in Tables 1A, 1B, and 1C that are listed and published in section VI. of the Addendum to this proposed rule and are available via the Internet on the CMS Web site.

2. Computing the National Average Standardized Amount

Section 1886(d)(3)(A)(iv)(I) of the Act requires that, beginning with FY 2004 and thereafter, an equal standardized amount be computed for all hospitals at the level computed for large urban hospitals during FY 2003, updated by the applicable percentage update. Accordingly, we are proposing to calculate the FY 2016 national average wage standardized amount on the respective of whether a hospital is located in an urban or rural location.

3. Updating the National Average Standardized Amount

Section 1886(b)(3)(B) of the Act specifies the applicable percentage increase used to update the standardized amount for payment for inpatient hospital operating costs. We note that, in compliance with section 404 of the MMA, in this proposed rule, we are proposing to use the revised and rebased 2014-based IPPS operating and capital market baskets for FY 2018. As discussed in section V.B. of the preamble of this proposed rule, in accordance with section 1886(b)(3)(B) of the Act, as amended by section 3401(a) of the Affordable Care Act, we are proposing to reduce the FY 2018 applicable percentage increase (which is based on IHS Global Insight, Inc.’s fourth quarter 2016 forecast of the proposed 2014-based IPPS market basket) by the MFP adjustment (the 10-year moving average of MFP for the period ending FY 2018) of 0.4 percentage point, which is calculated based on IGI’s fourth quarter 2016 forecast.

In addition, in accordance with section 1886(b)(3)(B)(i) of the Act, as amended by sections 3401(a) and 10319(a) of the Affordable Care Act, we are proposing to further update the standardized amount for FY 2018 by the estimated market basket percentage increase less 0.75 percentage point for hospitals in all areas. Sections 1886(b)(3)(B)(xi) and (xii) of the Act, as added and amended by sections 3401(a) and 10319(a) of the Affordable Care Act, further state that these adjustments may result in the application of a percentage less than zero. The percentage increase in the market basket reflects the average change in the price of goods and services required as inputs to provide hospital inpatient services. Based on IGI’s 2016 fourth quarter forecast of the hospital market basket increase (as discussed in Appendix B of this proposed rule), the forecast of the hospital market basket increase for FY 2018 for this proposed rule is 2.9 percent. As discussed earlier, for FY 2018, depending on whether a hospital submits quality data under the rules established in accordance with section 1886(b)(3)(B)(iii) of the Act and is a meaningful EHR user under section 1886(b)(3)(B)(ix) of the Act, there are four possible applicable percentage increases that could be applied to the standardized amount. We refer readers to section V.B. of the preamble of this proposed rule for a complete discussion on the proposed FY 2018 inpatient hospital update to the standardized amount. We also refer readers to the table above for the four applicable percentage increases that would be applied to update the national standardized amount. The proposed standardized amounts shown in Tables 1A through 1C that are published in section VI. of this Addendum and that are available via the Internet on the CMS Web site reflect these differential amounts.

Although the update factors for FY 2018 are set by law, we are required by section 1886(e)(4) of the Act to recommend, taking into account MedPAC’s recommendations, appropriate update factors for FY 2018 for both IPPS hospitals and hospital units excluded from the IPPS. Section 1886(e)(5)(A) of the Act requires that we publish our proposed recommendations in the Federal Register for public comment. Our recommendation on the update factors is set forth in Appendix B of this proposed rule.

4. Methodology for Calculation of the Average Standardized Amount

The methodology we used to calculate the proposed FY 2018 standardized amount is as follows:

- To ensure we are only including hospitals paid under the IPPS in the calculation of the standardized amount, we apply the following inclusion and exclusion criteria: include hospitals whose last four digits fall between 0001 and 0879 (section 2779A1 of Chapter 2 of the State Operations Manual at https://www.cms.gov/Regulations-and-Guidance/Guidance/Manuals/Downloads/som107c02.pdf), exclude CAGHs at the time of this proposed rule; exclude hospitals in Maryland (because these hospitals are paid under an all payer model under section 1115A of the Act); and remove PPS-excluded cancer hospitals that have a “V” in the fifth position of their provider number or a “F” or “F” in the sixth position.
- As in the past, we are proposing to adjust the FY 2018 standardized amount to remove the effects of the FY 2017 geographic reclassifications and outlier payments before applying the FY 2018 updates. We then apply budget neutrality offsets for outliers and geographic reclassifications to the standardized amount based on proposed FY 2018 payment policies.
- We do not remove the prior year’s budget neutrality adjustments for reclassification and recalibration of the DRG relative weights and for updated wage data because, in accordance with sections 1886(d)(4)(C)(iii) and 1886(d)(3)(E) of the Act, estimated aggregate payments after updates in the DRG relative weights and wage index should equal estimated aggregate payments prior to the changes. If we removed the prior year’s adjustment, we would not satisfy these conditions.

Budget neutrality is determined by comparing aggregate IPPS payments before and after making changes that are required to be budget neutral (for example, changes to MS–DRG classifications, recalibration of the MS–DRG relative weights, updates to the wage index, and different geographic reclassifications). We include outlier payments in the simulations because they may be affected by changes in these parameters.

- Consistent with our methodology established in the FY 2011 IPPS/LTCH PPS final rule (75 FR 50422 through 50433), because IME Medicare Advantage payments are made to IPPS hospitals under section 1886(d) of the Act, we believe these payments must be part of these budget neutrality calculations. However, we note that it is not necessary to include Medicare Advantage IME payments in the outlier threshold calculation or the outlier offset to the standardized amount because the statute requires that outlier payments be not less than 5 percent nor more than 6 percent of total operating DRG payments; which does not include IME and DSH payments. We refer readers to the FY 2011 IPPS/LTCH PPS final rule for a complete discussion on our methodology of identifying and adding the total Medicare Advantage IME payment amount to the budget neutrality adjustments.
- Consistent with the methodology in the FY 2012 IPPS/LTCH PPS final rule, in order to ensure that we capture only fee-for-service claims, we are only including claims with a “Claim Type” of 60 (which is a field on the MedPAR file that indicates a claim is an FFS claim).
- Consistent with our methodology established in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57277), in order to further ensure that we capture only FFS claims, we are excluding claims with a “GHOPAID” indicator of 1 (which is a field on the MedPAR file that indicates a claim is not an FFS claim and is paid by a Group Health Organization).
- Consistent with our methodology established in the FY 2011 IPPS/LTCH PPS final rule (75 FR 50422 through 50433), we examine the MedPAR file and remove...
pharmacy charges for anti-hemophilic blood factor (which are paid separately under the IPPS) with an indicator of “3” for blood clotting with a revenue code of “0636” from the covered charge field for the budget neutrality adjustments. We also remove organ acquisition charges from the covered charge field for the budget neutrality adjustments because organ acquisition is a pass-through payment not paid under the IPPS.

- The Bundled Payments for Care Improvement (BPCI) initiative, developed under authority of section 3021 of the Affordable Care Act (codified at section 1115A of the Act), is comprised of four broadly defined models of care, which link payments for multiple services beneficiaries receive during an episode of care. Under the BPCI initiative, organizations enter into payment arrangements that include financial and performance accountability for episodes of care. On January 31, 2013, CMS announced the first set of health care organizations selected to participate in the BPCI initiative, and organizations were selected in 2014. For additional information on the BPCI initiative, we refer readers to the CMS Center for Medicare and Medicaid Innovation’s Web site at: http://innovation.cms.gov/initiatives/Bundled-Payments/index.html.

In the FY 2013 IPPS/LTCH PPS final rule (77 FR 53341 through 53343), for FY 2013 and subsequent fiscal years, we finalized a methodology to treat hospitals that participate in the BPCI initiative the same as prior fiscal years for the IPPS payment modeling and ratesetting process (which includes recalibration of the MS–DRG weight factors for discharges occurring in FY 2018. (For additional information on our general policy for the reporting of hospital-specific readmission rates, consistent with section 1886(q)(6) of the Act. We discuss our proposed policy regarding the reporting of hospital-specific readmission rates for FY 2018 in section V.13.f. of the preamble of this proposed rule. (For additional information on our general policy for the reporting of hospital-specific readmission rates, consistent with section 1886(q)(6) of the Act. We refer readers to the FY 2013 IPPS/LTCH PPS final rule (77 FR 53390 through 53400),

In addition, for FY 2018, in this proposed rule, for the purpose of modeling aggregate payments when determining all budget neutrality factors, we are proposing to use proxy hospital VBP payment adjustment factors for FY 2018 based on data from a historical period because hospitals have not yet had an opportunity to review and submit corrections for their data from the FY 2018 performance period. (For additional information on our policy regarding the review and correction of hospital-specific measure rates under the Hospital VBP Program, consistent with section 1886(o)(10)(A)(ii) of the Act, we refer readers to the FY 2013 IPPS/LTCH PPS final rule (77 FR 53578 through 53581), the CY 2012 OPPS/ASC final rule with comment period (76 FR 74534 through 74547), and the Hospital Inpatient VBP final rule (76 FR 26353 through 26361).

- The Affordable Care Act also established section 1886(r) of the Act, which modifies the methodology for computing the Medicare DSH payment adjustments for Medicare DSH payments on both sides of the comparison for budget neutrality, prior to FY 2014, we included estimated Medicare DSH payments on both sides of our comparison of aggregate payments when determining all budget neutrality factors described in section II.A.4. of this Addendum.

To do this for FY 2018 (as we did for the last 4 fiscal years), we are proposing to include estimated empirically justified Medicare DSH payments that will be paid in accordance with section 1886(r)(1) of the Act and estimates of the additional uncompensated care payments made to hospitals receiving Medicare DSH payments adjustments as described by section 1886(r)(2) of the Act. That is, we are proposing to consider estimated empirically justified Medicare DSH payments at 25 percent of what would otherwise have been paid, and also the estimated additional uncompensated care payments for hospitals receiving Medicare DSH payment adjustments on both sides of our comparison of aggregate payments when determining all budget neutrality factors described in section II.A.4. of this Addendum.

- When calculating total payments for budget neutrality, to determine total payments for SCHs, we model total hospital-specific rate payments and total Federal rate payments and then include whichever one of the total payments is greater. As discussed in section V.G, of the preamble to this proposed rule and below, we are proposing to continue the FY 2014 finalized methodology under which we would take into consideration uncompensated care payments in the comparison of payments under the Federal rate and the hospital-specific rate for SCHs. Therefore, we are proposing to include estimated uncompensated care payments in this comparison.

- We are proposing to include an adjustment to the DSH payments made to hospitals that are not meaningful EHR users in our modeling of aggregate payments for budget neutrality for FY 2018. Similar to FY 2017, we are including this adjustment based on data on the prior year’s performance. Payments for hospitals would be estimated based on the proposed applicable standardized amount in Tables 1A and 1B for discharges occurring in FY 2018.

a. Proposed Recalibration of MS–DRG Relative Weights

Section 1886(d)(4)(C)(iii) of the Act specifies that, beginning in FY 1991, the annual DRG recategorization and recalibration of the relative weights must be made in a manner that ensures that aggregate payments
to hospitals are not affected. As discussed in section II.G. of the preamble of this proposed rule, we normalized the recalculated MS–DRG relative weights by an adjustment factor so that the average case relative weight after recalibration is equal to the average case relative weight before recalibration. However, equating the average case relative weight after recalibration to the average case relative weight before recalibration does not necessarily achieve budget neutrality with respect to aggregate payments to hospitals because the wages of hospitals are affected by factors other than average case relative weight. Therefore, as we have done in past years, we are proposing to make a budget neutrality adjustment to ensure that the requirement of section 1886(d)(4)(C)(ii) of the Act is met.

For FY 2018, to comply with the requirement that MS–DRG reclassification and recalibration of the relative weights be budget neutral for the standardized amount and the hospital-specific rates, we used FY 2016 discharge data to simulate payments and compared the following:

- Aggregate payments using the FY 2017 labor-related share percentages, the FY 2017 relative weights, and the FY 2017 pre-recalculated wage data, and applied the proposed FY 2018 hospital readmissions payment adjustments and estimated FY 2018 hospital VBP payment adjustments; and
- Aggregate payments using the FY 2017 labor-related share percentages, the FY 2017 relative weights, and the FY 2017 pre-recalculated wage data, and applied the same proposed FY 2018 hospital readmissions payment adjustments and estimated FY 2018 hospital VBP payment adjustments applied above.

Based on this comparison, we computed a proposed budget neutrality adjustment factor equal to 0.997573 and applied this factor to the standardized amount. As discussed in section IV. of this Addendum, we also are proposing to apply the MS–DRG reclassification and recalibration budget neutrality factor of 0.997755 to the hospital-specific rates that are effective for cost reporting periods beginning on or after October 1, 2017.

b. Updated Wage Index—Budget Neutrality Adjustment

Section 1886(d)(3)(E)(i) of the Act requires us to update the hospital wage index on an annual basis beginning October 1, 1993. This provision also requires us to make any updates or adjustments to the wage index in a manner that ensures that aggregate payments to hospitals are not affected by the change in the wage index. Section 1886(d)(3)(E)(ii) of the Act requires that we implement the wage index adjustment in a budget neutral manner. However, section 1886(d)(3)(E)(ii) of the Act sets the labor-related share at 62 percent for hospitals with a wage index less than or equal to 1.0000, and section 1886(d)(3)(E)(ii) of the Act provides that the Secretary shall calculate the budget neutrality adjustment for the adjustments or updates made under that provision as if section 1886(d)(3)(E)(ii) of the Act had not been enacted. In other words, this section of the statute requires that we implement the updates to the wage index in a budget neutral manner, but that our budget neutrality adjustment should not take into account the requirement that we set the labor-related share for hospitals with wage indexes less than or equal to 1.0000 at the more advantageous level of 62 percent. Therefore, in making the budget neutrality adjustment, section 1886(d)(3)(E)(i) of the Act prohibits us from taking into account the fact that hospitals with a wage index less than or equal to 1.0000 are paid using a labor-related share of 62 percent.

Consistent with section 1886(d)(3)(E)(i) of the Act, we are proposing to adjust 100 percent of the wage index factor for occupational mix. We describe the occupational mix adjustment in section III.E. of the preamble of this proposed rule.

To compute a proposed budget neutrality adjustment factor for wage index and labor-related share percentage changes, we used FY 2016 discharge data to simulate payments and compared the following:

- Aggregate payments using the proposed FY 2018 wage index and labor-related share percentage changes, proposed FY 2018 labor-related share percentages, proposed FY 2018 relative weights, and applied the proposed FY 2018 wage data after any reclassifications under sections 1886(d)(6)(B) and (C) and 1886(d)(10) of the Act, and
- Aggregate payments using the FY 2017 wage index and labor-related share percentage changes, FY 2017 labor-related share percentages, the FY 2017 relative weights, and applied the FY 2017 wage data after any reclassifications under sections 1886(d)(6)(B) and (C) and 1886(d)(10) of the Act.

We note that the reclassifications applied under the second simulation and comparison are those listed in Table 2 associated with this proposed rule, which is available via the Internet on the CMS Web site. This table reflects reclassification crosswalks proposed for FY 2018, and apply the proposed policies explained in section III. of the preamble to this proposed rule. Based on these simulations, we calculated a proposed budget neutrality adjustment factor of 0.988522 to ensure that the effects of these provisions are budget neutral, consistent with the statute.

The proposed FY 2018 budget neutrality adjustment factor was applied to the standardized amount after removing the effects of the FY 2017 budget neutrality adjustment factor. We note that the proposed FY 2018 budget neutrality adjustment reflects FY 2018 wage data after any reclassifications approved by the MGCRB or the Administrator at the time of development of this proposed rule.

c. Reclassified Hospitals—Proposed Budget Neutrality Adjustment

Section 1886(d)(9)(B) of the Act provides that certain rural hospitals are deemed urban. In addition, section 1886(d)(10) of the Act provides for the reclassification of hospitals based on determinations by the MGCRB. Under section 1886(d)(10) of the Act, a hospital may be reclassified for purposes of the wage index. Under section 1886(d)(6)(D) of the Act, the Secretary is required to adjust the standardized amount to ensure that aggregate payments under the IPPS after implementation of the provisions of sections 1886(d)(8)(B) and (C) and 1886(d)(10) of the Act are equal to the aggregate prospective payments that would have been made absent these provisions. We note that the wage index adjustments provided for under section 1886(d)(13) of the Act are not budget neutral.

Section 1886(d)(13)(H) of the Act provides that any increase in a wage index under section 1886(d)(13) shall not be taken into account in applying any budget neutrality adjustment with respect to such index under section 1886(d)(9)(D) of the Act. To calculate the proposed budget neutrality adjustment for FY 2018, we used FY 2016 discharge data to simulate payments and compared the following:

- Aggregate payments using the proposed FY 2018 labor-related share percentages, proposed FY 2018 relative weights, and applied the proposed FY 2018 wage data after any reclassifications under sections 1886(d)(6)(B) and (C) and 1886(d)(10) of the Act, and
- Aggregate payments using the proposed FY 2018 labor-related share percentages, proposed FY 2018 relative weights, and applied the FY 2017 wage data after any reclassifications under sections 1886(d)(6)(B) and (C) and 1886(d)(10) of the Act.

We note that the reclassifications applied under the second simulation and comparison are those listed in Table 2 associated with this proposed rule, which is available via the Internet on the CMS Web site. This table reflects reclassification crosswalks proposed for FY 2018, and apply the proposed policies explained in section III. of the preamble to this proposed rule. Based on these simulations, we calculated a proposed budget neutrality adjustment factor of 0.988522 to ensure that the effects of these provisions are budget neutral, consistent with the statute.

d. Proposed Rural Floor Budget Neutrality Adjustment

Under §412.64(e)(4), we make an adjustment to the wage index to ensure that aggregate payments after implementation of the rural floor under section 4410 of the BBA (Pub. L. 105–33) is equal to the aggregate prospective payments that would have been made in the absence of this provision. Consistent with section 3141 of the Affordable Care Act and as discussed in section III.H. of the preamble of this proposed rule and codified at §412.64(e)(4)(ii), the budget neutrality adjustment for the rural floor is a national adjustment to the wage index.

As noted above and as discussed in section III.H. of the preamble of this proposed rule, the imputed floor is set to expire effective October 1, 2017, and we are not proposing to extend the imputed floor policy.

Similar to our calculation in the FY 2015 IPPS/LTCIPPS final rule (79 FR 50369 through 50379), for FY 2018, we are proposing to calculate a national rural Puerto
 Ricardo wage index. Because there are no rural Puerto Rico hospitals with established wage data, our calculation of the proposed FY 2018 rural Puerto Rico wage index is based on the policy adopted in the FY 2008 IPPS final rule with comment period (72 FR 47523). That is, we will continue to average the wage indexes from all CBSAs (urban areas) that are contiguous (share a border with) to the rural counties to compute the rural floor (72 FR 47323; 76 FR 51594). Under the OMB labor market area delineations, except for Aricibo, Puerto Rico (CBSA 11640), all other Puerto Rico urban areas are contiguous to a rural area. Therefore, based on our existing policy, the proposed FY 2018 rural Puerto Rico wage index is calculated based on the average of the proposed FY 2018 wage indexes for the following urban areas: Aguadilla-Isabela, PR (CBSA 10380); Guayanilla, PR (CBSA 25020); Mayaguez, PR (CBSA 32420); Ponce, PR (CBSA 38660); San German, PR (CBSA 41900); and San Juan-Carolina-Cayey, PR (CBSA 41980).

To calculate the national rural floor budget neutrality adjustment factor, we are proposing to use FY 2016 discharge data to simulate payments and the proposed post-reclassified national wage indexes and compared the following:

- National simulated payments without the proposed national rural floor; and
- National simulated payments with the proposed national rural floor.

Based on this comparison, we determined a proposed national rural floor budget neutrality adjustment factor of 0.993672. The national adjustment was applied to the national wage indexes to produce a proposed national rural floor budget neutral wage index.

e. Proposed Adjustment for FY 2018

Required Under Section 414 of Public Law 114–10 (MACRA) and Section 15005 of Public Law 114–255

As stated in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56785), once the recoupment required under section 631 of the ATRA was complete, we had anticipated making a single positive adjustment in FY 2018 to offset the reductions required to recoup the $11 billion under section 631 of the ATRA. However, section 414 of the MACRA (which was enacted on April 16, 2015) replaced the single positive adjustment we intended to make in FY 2018 with a 0.5 percent positive adjustment for each of FYs 2018 through 2023. In the FY 2017 rulemaking, we indicated that we would address the adjustments for FY 2018 and later fiscal years in future rulemaking. As noted previously, section 15005 of the 21st Century Cures Act (Public Law 114–255), which was enacted December 13, 2016, amended section 7(b)[1][B] of the OMB, as amended by section 631 of the ATRA and section 414 of the MACRA, to reduce the adjustment for FY 2018 from 0.5 percentage points to 0.4588 percentage points. Therefore, for FY 2018, we are proposing to implement the required +0.4588 percent adjustment to the standardized amount. This is a permanent adjustment to payment rates. While we are not proposing future adjustments required under section 414 of the MACRA and section 15005 of Public Law

As we have done in the past, to calculate the proposed FY 2018 outlier threshold, we simulated payments by applying proposed FY 2018 payment rates and policies using cases from the FY 2016 MedPAR file. Therefore, in order to determine the proposed FY 2018 outlier threshold, we inflated the charges on the MedPAR claims by 2 years, from FY 2016 to FY 2018. As discussed in the FY 2015 IPPS/LTCH PPS final rule, we believe a methodology that is based on 1-year of charge data will provide a more stable measure to project the average charge per case because our prior methodology used a 6-month measure, which inherently uses fewer claims than a 1-year measure and makes it more susceptible to fluctuations in the average charge per case as a result of any significant charge increases or decreases by hospitals. As finalized in the FY 2017 IPPS/LTCH final rule (81 FR 57282), we are using the following methodology to calculate the charge inflation factor for FY 2018:

- To produce the most stable measure of charge inflation, we applied the following inclusion and exclusion criteria of hospitals claims in our measure of charge inflation:
  - Include hospitals whose last four digits fall between 0001 and 0099 (section 2779A1 of Chapter 2 of the State Operations Manual on the CMS Web site at https://www.cms.gov/Regulations-and-Guidance/Guidance/Manuals/Downloads/som107c02.pdf); include CAHs that were IPPS hospitals for the time period of the MedPAR data being used to calculate the charge inflation factor; include hospitals in Maryland; and remove PPS-excluded cancer hospitals who have a “V” in the fifth position of their provider number or a “E” or “F” in the sixth position.
  - We excluded Medicare Advantage IME claims for the reasons described in section I.A. of this Addendum. We refer readers to the FY 2011 IPPS/LTCH PPS final rule for a complete discussion on our methodology of identifying and adding the total Medicare Advantage IME payment amount to the budget neutrality adjustments.

In order to ensure that we capture only FFS claims, we included claims with a “Claim Type” of 60 (which is a field on the MedPAR file that indicates a claim is a FFS claim).

- In order to further ensure that we capture only FFS claims, we excluded claims with a “GHOPAI” indicator of 1 (which is a field on the MedPAR file that indicates a claim is not an FFS claim and is paid by a Group Health Organization).

- We examined the MedPAR file and removed pharmacy charges for anti-hemophilic blood factor (which are paid separately under the IPPS) with an indicator of “0636” from the covered charge field. We also removed organ acquisition charges from the covered charge field because organ acquisition is a pass-through payment not paid under the IPPS.

In the FY 2016 IPPS/LTCH final rule (80 FR 49779–49780), we stated that commenters were concerned that they were unable to replicate the calculation of the charge inflation factor that CMS used in the proposed rule. In response to those
Under this methodology, to compute the 1-year average annualized rate-of-change in charges per case for FY 2018, we compared the average covered charge per case of $53,287 ($517,993,138,897/9,720,768) from the second quarter of FY 2015 through the first quarter of FY 2016 (January 1, 2015, through December 31, 2015) to the average covered charge per case of $55,991 ($508,057,757,077/9,073,897) from the second quarter of FY 2016 through the first quarter of FY 2017 (January 1, 2016, through December 31, 2016). This rate-of-change is 5.1 percent (1.05074) or 10.4 percent (1.104055) over 2 years. The billed charges are obtained from the claim from the MedPAR file and inflated by the inflation factor specified above.

As we have done in the past, in this proposed rule, we are proposing to establish the proposed FY 2018 outlier threshold using hospital CCRs from the December 2016 update to the Provider-Specific File (PSF)—the most recent available data at the time of the development of this proposed rule. We are proposing to apply the following edits to providers’ CCRs in the PSF. We believe these edits are appropriate in order to accurately model the outlier threshold. We first search for Indian Health Service providers and those providers assigned the statewide average CCR from the current fiscal year. We then replace these CCRs with the statewide average CCR for the upcoming fiscal year per also assign the statewide average CCR (for the upcoming fiscal year) to those providers that have no value in the CCR field in the PSF or whose CCRs exceed the ceilings described later in this section (3.0 standard deviations from the mean of the log distribution of CCRs for all hospitals). We do not apply the adjustment factors described below to hospitals assigned the statewide average CCR.

For FY 2018, we also are proposing to continue to apply an adjustment factor to the CCRs for cost and charge inflation (as explained below). We are proposing that, if more recent data become available, we would use that data to calculate the final FY 2018 outlier threshold.

In the FY 2014 IPPS/LTHCH PPS final rule (78 FR 50979), we adopted a new methodology to adjust the CCRs. Specifically, we finalized a policy to compare the national average case-weighted operating and capital CCR from the most recent update of the PSF to the national average case-weighted operating and capital CCR from the same period of the prior year.

Therefore, as we did for the last 4 fiscal years, we are proposing to adjust the CCRs from the December 2016 update of the PSF by comparing the percentage change in the national average case-weighted operating CCR and capital CCR from the December 2016 update of the PSF to the national average case-weighted operating CCR and capital CCR from the December 2015 update of the PSF. We note that we used total transfer-adjusted cases from FY 2016 to determine the national average case-weighted CCRs for both sides of the comparison. As stated in the FY 2014 IPPS/LTHCH PPS final rule (78 FR 50979), we believe that it is appropriate to use the same case count on both sides of the comparison because this will produce the true percentage change in the average case-weighted operating and capital CCR from one year to the next without any effect from a change in case count on different sides of the comparison.

Using the proposed methodology above, we calculated a proposed December 2015 operating national average case-weighted CCR of 0.274139 and a proposed December 2016 operating national average case-weighted CCR of 0.26579. We then calculated the percentage change between the two national operating case-weighted CCRs by subtracting the December 2015 operating national average case-weighted CCR from the December 2016 operating national average case-weighted CCR and then dividing the result by the December 2015 national operating average case-weighted CCR. This resulted in a proposed national operating CCR adjustment factor of 0.979187.

We used the same methodology proposed above to make the same adjustments for the national capital CCR. Specifically, we calculated a December 2015 capital national average case-weighted CCR of 0.024074 and a December 2016 capital national average case-weighted CCR of 0.022967. We then calculated the percentage change between the two national capital case-weighted CCRs by subtracting the December 2015 capital national average case-weighted CCR from the December 2016 capital national average case-weighted CCR and then dividing the result by the December 2015 capital national average case-weighted CCR. This resulted in a proposed national capital CCR adjustment factor of 0.950568.

As discussed in section III.B.3. of the preamble to the FY 2011 IPPS/LTHCH PPS final rule (75 FR 50160 and 50161) and in section III.H.3. of the preamble of this proposed rule, in accordance with section 10324(a) of the Affordable Care Act, we created a wage index floor of 1.0000 for all hospitals located in States determined to be frontier States. We note that the frontier State floor adjustments would be calculated and applied after rural floor budget neutrality adjustments are calculated for all labor market areas, in order to ensure that no hospital in a frontier State would receive a wage index less than 1.0000 due to the proposed rural floor adjustment. In accordance with section 10324(a) of the Affordable Care Act, the frontier State adjustment will not be subject to budget neutrality, and will only be extended to hospitals geographically located within a frontier State. However, for purposes of estimating the proposed outlier threshold for FY 2018, it was necessary to adjust the proposed wage index of those eligible hospitals in a frontier State when calculating the proposed outlier threshold that results in outlier payments being 5.1 percent of total payments for FY 2018. If we did not take the above into account, our estimate of total FY 2018 payments would be too low, and, as a result, our proposed outlier threshold would be too high, such that estimated outlier payments would be less than our projected 5.1 percent of total payments.

As we did in establishing the FY 2009 outlier threshold (73 FR 57891), in our projection of FY 2018 outlier payments, we may need to make any adjustments for the possibility that hospitals’ CCRs and outlier payments may be reconciled upon cost report settlement. We continue to believe that, due to the policy implemented in the June 9, 2003 Outlier Final Rule (68 FR 34494), CCRs will no longer fluctuate significantly and, therefore, few hospitals
will actually have these ratios reconciled upon cost report settlement. In addition, it is difficult to predict the specific hospitals that will have CCRs and outlier payments reconciled in any given year. We note that we have instructed MACs to identify for CMS any instance for which a hospital’s actual CCR for the cost reporting period fluctuates plus or minus 10 percentage points compared to the interim CCR used to calculate outlier payments when a bill is processed; and (2) the total outlier payments for the hospital exceed 5.1 percent of total operating or capital costs for that period. Our simulations assume that CCRs accurately measure hospital costs based on information available to us at the time we set the outlier threshold. For these reasons, we are proposing not to make any assumptions regarding the effects of reconciliation on the outlier threshold calculation.

As described in sections V.I. and V.J. respectively, of the preamble of this proposed rule, sections 1886(q) and 1886(o) of the Act establish the Hospital Readmissions Reduction Program and the Hospital VBP Program, respectively. We do not believe that it is appropriate to include the hospital VBP payment adjustments and the hospital readmissions payment adjustments in the proposed outlier threshold calculation or the proposed outlier threshold calculation for the hospitals for which the MAC computes operating CCRs greater than 1.17 or capital CCRs greater than 0.161, or hospitals for which the MAC is unable to calculate a CCR (as described in § 412.84(i)(3)(ii) of the Act). Statewide average CCRs are used to determine whether a hospital qualifies for outlier payments. Table 8A listed in section VI. of this Addendum (and available only via the Internet on the CMS Web site) contains the proposed statewide average operating CCRs for urban hospitals and for rural hospitals for which the MAC is unable to compute a hospital-specific CCR within the above range. These statewide average ratios would be effective for discharges occurring on or after October 1, 2017, and would supersede our previously published statewide average ratios from the prior fiscal year. Table 8B listed in section VI. of this Addendum (and available via the Internet on the CMS Web site) contains the comparable proposed statewide average capital CCRs. As previously stated, the proposed CCRs in Tables 8A and 8B would be used during FY 2018 when hospital-specific CCRs based on the latest settled cost report either are not available or are outside the range noted above. Table 8C listed in section VI. of this Addendum (and available via the Internet on the CMS Web site) contains the proposed statewide average total CCRs under the LTCH PPS as discussed in section V. of this Addendum.

We finally note that we published a manual update (Change Request 3966) to our outlier policy on October 12, 2005, which updated Chapter 3, Section 20.1.2 of the Medicare Claims Processing Manual. The manual update covered an array of topics, including CCRs, reconciliation, and the time value of money. We encourage hospitals that are assigned the statewide average operating and/or capital CCRs to work with their MAC on a possible alternative operating and/or capital CCR as explained in Change Request 3966. Use of an alternative CCR developed by the hospital in conjunction with the MAC can avoid possible overpayments or underpayments at cost report settlement, thereby ensuring better accuracy when making outlier payments and negating the need for outlier reconciliation. We also note that a hospital may request an alternative operating or capital CCR ratio at any time as long as the guidelines of Change Request
3966 are followed. In addition, as mentioned above, we published an additional manual update (Change Request 7192) to our outlier policy on December 3, 2010, which also updated Chapter 3, Section 20.1.2 of the Medicare Claims Processing Manual. The manual update outlines the outlier reconciliation process for hospitals and Medicare contractors. To download and view the manual instructions on outlier reconciliation, we refer readers to the CMS Web site: http://www.cms.hhs.gov/manuals/downloads/clm104c03.pdf.

(3) FY 2016 Outlier Payments

Our current estimate, using available FY 2016 claims data, is that actual outlier payments for FY 2016 were approximately 5.37 percent of actual total MS–DRG payments. Therefore, the data indicate that, for FY 2016, the percentage of actual outlier payments relative to actual total payments is higher than we projected for FY 2016. Consistent with the policy and statutory interpretation we have maintained since the inception of the IPPS, we do not make retroactive adjustments to outlier payments to ensure that total outlier payments for FY 2016 are equal to 5.1 percent of total MS–DRG payments. As explained in the FY 2003 Outlier Final Rule (68 FR 34502), if we were to make retroactive adjustments to outlier payments to ensure actual total payments are 5.1 percent of MS–DRG payments (by retroactively adjusting outlier payments), we would be removing the important aspect of the prospective nature of the IPPS. Because such an across-the-board adjustment would either lead to more or less outlier payments for all hospitals, hospitals would no longer be able to reliably approximate their payment for a patient while the patient is still hospitalized. We believe it would be neither necessary nor appropriate to make such an adjustment to outlier payments.

Furthermore, we believe it is consistent with the statutory language at section 1886(d)(5)(A)(iv) of the Act not to make retroactive adjustments to outlier payments. This section calls for the Secretary to ensure that outlier payments are equal to or greater than 5 percent and less than or equal to 6 percent of projected or estimated (not actual) MS–DRG payments. We believe that an important goal of a PPS is predictability. Therefore, we believe that the fixed-loss outlier threshold should be projected based on the best available historical data and should not be adjusted retroactively. A retroactive change to the fixed-loss outlier threshold would affect all hospitals subject to the IPPS, thereby undercutting the predictability of the system as a whole.

We note that because the MedPAR claims data for the entire FY 2017 will not be available until after September 30, 2017, we are unable to provide an estimate of actual outlier payments for FY 2017 based on FY 2017 claims data in this proposed rule. We will provide an estimate of actual FY 2017 outlier payments in the FY 2019 IPPS/LTCH PPS proposed rule.

5. Proposed FY 2018 Standardized Amount

The adjusted standardized amount is divided into labor-related and nonlabor-related portions. Tables 1A and 1B listed and published in section VI. of this Addendum (and available via the Internet on the CMS Web site) contain the national standardized amounts that we are proposing to apply to all hospitals, except hospitals located in Puerto Rico, for FY 2018. The proposed standardized amount for hospitals in Puerto Rico is shown in Table 1C listed and published in section VI. of this Addendum (and available via the Internet on the CMS Web site). The proposed amounts shown in Tables 1A and 1B differ only in that the labor-related share applied to the standardized amounts in Table 1B is 62 percent. In accordance with sections 1886(d)(3)(E) and 1886(d)(9)(C)(iv) of the Act, we are proposing to apply a labor-related share of 62 percent, unless application of that percentage would result in lower payments to a hospital than would otherwise be made. In effect, the statutory provision means that we will apply a labor-related share of 62 percent for all hospitals whose wage indexes are less than or equal to 1.0000.

In addition, Tables 1A and 1B include the proposed standardized amounts reflecting the proposed applicable percentage increases for FY 2018. The proposed labor-related and nonlabor-related portions of the national average standardized amounts for Puerto Rico hospitals for FY 2018 are set forth in Table 1C listed and published in section VI. of this Addendum (and available via the Internet on the CMS Web site). Similar to above, section 1886(d)(9)(C)(iv) of the Act, as amended by section 403(b) of Public Law 108–173, provides that the labor-related share for hospitals located in Puerto Rico be 62 percent, unless the application of that percentage would result in lower payments to the hospital.

The following table illustrates the changes from the FY 2017 national standardized amount to the proposed FY 2018 national standardized amount. The second through fifth columns display the proposed changes from the FY 2017 standardized amounts for each applicable FY 2018 standardized amount. The first row of the table shows the updated (through FY 2017) average standardized amount after restoring the FY 2017 offsets for outlier payments, geographic reclassification budget neutrality, new labor market delineation wage index transition budget neutrality and removing the FY 2017 2-midnight rule one-time prospective increase. The MS–DRG recategorization and recalculation and wage index budget neutrality adjustment factors are cumulative. Therefore, those FY 2017 adjustment factors are not removed from this table.

### Changes From FY 2017 Standardized Amounts To The Proposed FY 2018 Standardized Amounts

<table>
<thead>
<tr>
<th>Hospital submitted quality data and is a meaningful EHR user</th>
<th>Hospital submitted quality data and is NOT a meaningful EHR user</th>
<th>Hospital did NOT submit quality data and is a meaningful EHR user</th>
<th>Hospital did NOT submit quality data and is NOT a meaningful EHR user</th>
</tr>
</thead>
<tbody>
<tr>
<td>FY 2018 Base Rate after removing:</td>
<td>FY 2017 Geographic Reclassification Budget Neutrality (0.988136).</td>
<td>FY 2017 Operating Outlier Offset (0.949898).</td>
<td>FY 2017 2-Midnight Rule One-Time Prospective Increase (1.006).</td>
</tr>
<tr>
<td>If Wage Index is Greater Than 1.0000:</td>
<td>If Wage Index is Greater Than 1.0000:</td>
<td>If Wage Index is Greater Than 1.0000:</td>
<td>If Wage Index is Greater Than 1.0000:</td>
</tr>
<tr>
<td>Labor (68.3%):</td>
<td>Labor (68.3%):</td>
<td>Labor (68.3%):</td>
<td>Labor (68.3%):</td>
</tr>
<tr>
<td>$3,993.72.</td>
<td>$3,993.72.</td>
<td>$3,993.72.</td>
<td>$3,993.72.</td>
</tr>
<tr>
<td>Nonlabor (30.4%):</td>
<td>Nonlabor (30.4%):</td>
<td>Nonlabor (30.4%):</td>
<td>Nonlabor (30.4%):</td>
</tr>
<tr>
<td>$1,853.60.</td>
<td>$1,853.60.</td>
<td>$1,853.60.</td>
<td>$1,853.60.</td>
</tr>
<tr>
<td>If Wage Index is Less Than or Equal to 1.0000:</td>
<td>If Wage Index is Less Than or Equal to 1.0000:</td>
<td>If Wage Index is Less Than or Equal to 1.0000:</td>
<td>If Wage Index is Less Than or Equal to 1.0000:</td>
</tr>
<tr>
<td>Labor (62%):</td>
<td>Labor (62%):</td>
<td>Labor (62%):</td>
<td>Labor (62%):</td>
</tr>
<tr>
<td>$3,625.34.</td>
<td>$3,625.34.</td>
<td>$3,625.34.</td>
<td>$3,625.34.</td>
</tr>
<tr>
<td>Nonlabor (38%):</td>
<td>Nonlabor (38%):</td>
<td>Nonlabor (38%):</td>
<td>Nonlabor (38%):</td>
</tr>
<tr>
<td>$2,221.98.</td>
<td>$2,221.98.</td>
<td>$2,221.98.</td>
<td>$2,221.98.</td>
</tr>
<tr>
<td>Proposed FY 2018 Update Factor:</td>
<td>1.0175</td>
<td>0.99575</td>
<td>1.01025</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0.9885</td>
</tr>
</tbody>
</table>
We note that, in recent years, we have estimated the MS–DRG recalibration budget neutrality factor, wage index budget neutrality factor, recategorization budget neutrality factor and operating outlier factor to six decimal places. While we are not proposing any changes at this time, we are interested in receiving comments from the public as to the continued necessity of six decimal places for these four estimates or if fewer decimal places would be sufficient.

B. Proposed Adjustments for Area Wage Levels and Cost-of-Living

Tables 1A through 1C, as published in section VI. of this Addendum (and available via the Internet on the CMS Web site), contain the proposed labor-related and nonlabor-related shares that we are proposing to use to calculate the prospective payment rates for hospitals located in the 50 States, the District of Columbia, and Puerto Rico for FY 2018. This section addresses two types of adjustments to the standardized amounts that are made in determining the proposed prospective payment rates as described in this Addendum.

1. Proposed Adjustment for Area Wage Levels

Sections 1886(d)(3)(E) and 1886(d)(9)(C)(iv) of the Act require that we make an adjustment to the labor-related portion of the national prospective payment rate to account for area differences in hospital wage levels. This adjustment is made by multiplying the labor-related portion of the adjusted standardized amounts by the appropriate wage index for the area in which the hospital is located. For FY 2018, as discussed in section IV. B.3. of the preamble of this proposed rule, we are proposing to apply a labor-related share of 68.3 percent for the national standardized amounts for all IPPS hospitals (including hospitals in Puerto Rico) that have a wage index value that is greater than 1.0000. Consistent with section 1886(d)(3)(E) of the Act, we are proposing to apply the wage index to a labor-related share of 62 percent of the national standardized amount for all IPPS hospitals (including hospitals in Puerto Rico) whose wage index values are less than or equal to 1.0000. In section III. of the preamble of this proposed rule, we discuss the data and methodology for the proposed FY 2018 wage index.

2. Proposed Adjustment for Cost-of-Living in Alaska and Hawaii

Section 1886(d)(5)(H) of the Act provides discretionary authority to the Secretary to make adjustments as the Secretary deems appropriate to take into account the unique circumstances of hospitals located in Alaska and Hawaii. Higher labor-related costs for these two States are taken into account in the adjustment for area wages described above. To account for higher nonlabor-related costs for these two States, we multiply the nonlabor-related portion of the standardized amount for hospitals in Alaska and Hawaii by an adjustment factor. For FY 2011 and in prior fiscal years, we used the most recent cost-of-living adjustment (COLA) factors obtained from the U.S. Office of Personnel Management (OPM) Web site at http://www.opm.gov/oca/colas.asp to update this nonlabor portion.

In the FY 2012 IPPS/LTCH PPS final rule (76 FR 51797), we explained that sections 1911 through 1919 of the Nonforeign Area Retirement Equity Assurance Act, as contained in subtitle B of title XIX of the National Defense Authorization Act (NDAA) for Fiscal Year 2010 (Pub. L. 111–84, October 28, 2009), transitions the Alaska and Hawaii COLAs to locality pay. We finalized that, for FY 2012, as OPM transitioned away from COLAs, we would continue to use the same “frozen” COLA factors (published by OPM) that we used to adjust payments in FY 2011 (which were based on OPM’s 2009 COLA factors) to adjust the nonlabor-related portion of the standardized amount for hospitals located in Alaska and Hawaii. We refer readers to the FY 2012 IPPS/LTCH PPS final rule for a more detailed discussion of our rationale for continuing to use the frozen COLAs in FY 2012.

In the FY 2013 IPPS/LTCH PPS final rule (77 FR 53700 and 53701), for FY 2013, we continued to use the same COLA factors that were used to adjust payments in FY 2012 (as originally used to adjust payments in FY 2011, which were based on OPM’s 2009 COLA factors). We also established a methodology to update the COLA factors published by OPM every 4 years (at the same time as the update of the labor-related share of the IPPS market basket), beginning in FY 2014. We refer readers to the FY 2013 IPPS/LTCH PPS proposed rule (77 FR 28145 and 28146) for a detailed description of this methodology. For FY 2014, we updated the COLA factors for Alaska and Hawaii published by OPM for 2009 using the methodology finalized in the FY 2013 IPPS/LTCH PPS final rule (77 FR 53700 and 53701).

For FY 2018, we are proposing to continue to update the COLA factors published by OPM for 2009 (as these are the last COLA factors OPM published prior to transitioning from COLAs to locality pay) using the methodology that we finalized in the FY 2013 IPPS/LTCH PPS final rule and implemented for the FY 2014 IPPS update. Specifically, we are proposing to update the 2009 OPM COLA factors by a comparison of the growth in the Consumer Price Indices (CPIs) for Anchorage, AK, and Honolulu, HI,
We note that the reweighted CPI for Honolulu, HI grew faster than the reweighted CPI for the average U.S. city over the 2009 to 2016 time period, at 13.7 percent and 10.5 percent, respectively. As a result, for FY 2018, we calculated proposed COLA factors for the City and County of Honolulu, County of Kauai, County of Maui, and County of Kalawao to be 1.29 compared to the FY 2013 COLA factor of 1.25 (which was based on OPM’s published COLA factors for 2009, as described above). However, as stated above, we are applying our methodology as finalized in the FY 2013 IPPS/LTCH PPS final rule to incorporate a cap of 1.25 for these areas. In addition, the proposed COLA factor we calculated for the County of Hawaii for FY 2018 is 1.21 compared to the FY 2013 COLA factor of 1.18. The COLA factors adopted in FY 2014 using this same methodology can be found in the table above.

Similarly, the reweighted CPI for Anchorage, AK grew faster than the reweighted CPI for the average U.S. city over the 2009 to 2016 time period, at 12.4 percent and 10.5 percent, respectively. As a result, for FY 2018, we calculated proposed COLA factors for the City of Anchorage, City of Fairbanks, and City of Juneau to be 1.25 compared to the FY 2013 COLA factor of 1.23. For FY 2018, we calculated a proposed COLA factor of 1.27 for the rest of Alaska compared to the FY 2013 COLA factor of 1.25. However, as stated above, we are applying our methodology as finalized in the FY 2013 IPPS/LTCH PPS final rule to incorporate a cap of 1.25 for the Rest of Alaska.

As stated above, the COLA factors adopted in the FY 2014 IPPS/LTCH PPS final rule were based on the same methodology used to determine the proposed FY 2018 COLA factors but utilizing BLS data from 2009 through 2012 (the most recent data available at the time of this rulemaking) rather than through 2016 (the most recent data available at the time of this rulemaking). Compared to the FY 2014 COLA factors, the proposed FY 2018 COLA factors are higher—with all areas either reaching or exceeding the cap of 1.25 except the County of Hawaii.

### COST-OF-LIVING ADJUSTMENT FACTORS: ALASKA AND HAWAI'I HOSPITALS

<table>
<thead>
<tr>
<th>Area</th>
<th>FY 2013</th>
<th>FY 2014 through FY 2017</th>
<th>Proposed FY 2018</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alaska:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>City of Anchorage and 80-kilometer (50-mile) radius by road</td>
<td>1.23</td>
<td>1.23</td>
<td>1.25</td>
</tr>
<tr>
<td>City of Fairbanks and 80-kilometer (50-mile) radius by road</td>
<td>1.23</td>
<td>1.23</td>
<td>1.25</td>
</tr>
<tr>
<td>City of Juneau and 80-kilometer (50-mile) radius by road</td>
<td>1.23</td>
<td>1.23</td>
<td>1.25</td>
</tr>
<tr>
<td>Rest of Alaska</td>
<td>1.25</td>
<td>1.25</td>
<td>1.25</td>
</tr>
<tr>
<td>Hawaii:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>City and County of Honolulu</td>
<td>1.25</td>
<td>1.25</td>
<td>1.25</td>
</tr>
<tr>
<td>County of Kauai</td>
<td>1.18</td>
<td>1.19</td>
<td>1.21</td>
</tr>
<tr>
<td>County of Maui and County of Kalawao</td>
<td>1.25</td>
<td>1.25</td>
<td>1.25</td>
</tr>
</tbody>
</table>

**C. Calculation of the Proposed Prospective Payment Rates**

**General Formula for Calculation of the Prospective Payment Rates for FY 2018**

In general, the operating prospective payment rate for all hospitals (including hospitals in Puerto Rico) paid under the IPPS, except SCHs, for FY 2018 equals the Federal rate (which includes uncompensated care payments).

SCHs are paid based on whichever of the following rates yields the greatest aggregate payment: The Federal national rate (which, as discussed in section V.G. of the preamble of this proposed rule, includes uncompensated care payments); the updated hospital-specific rate based on FY 1982 costs per discharge; the updated hospital-specific rate based on FY 1987 costs per discharge; the updated hospital-specific rate based on FY 1996 costs per discharge; or the updated hospital-specific rate based on FY 2006 costs per discharge to determine the rate that yields the greatest aggregate payment.

The prospective payment rate for SCHs for FY 2018 equals the higher of the applicable rate.
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Federal rate, or the hospital-specific rate as described below.

1. Operating and Capital Federal Payment Rate and Outlier Payment Calculation

   Note: The formula below is used for actual claim payment and is also used by CMS to project the outlier threshold for the upcoming fiscal year. The difference is the source of some of the variables in the formula. For example, operating and capital CCRs for actual claim payment are from the PSF while CMS uses an adjusted CCR as described above to project the threshold for the upcoming fiscal year. In addition, charges for a claim payment are from the bill while charges to project the threshold are from the MedPAR data with an inflation factor applied to the charges as described earlier.

   Step 1—Determine the MS–DRG and MS–DRG relative weight for each claim based on the ICD–10–CM procedure and diagnosis codes on the claim.

   Step 2—Select the applicable average standardized amount depending on whether the hospital submitted qualifying quality data and is a meaningful EHR user, as described above.

   Step 3—Compute the operating and capital Federal payment rate:

   —Federal Payment Rate for Operating Costs = MS–DRG Relative Weight × ([Labor–Related Applicable Standardized Amount × Applicable CBSA Wage Index) + (Nonlabor–Related Applicable Standardized Amount × Cost–of–Living Adjustment) × (1 + IME + (DSH × 0.25))]
   —Operating Payment for Capital Costs = MS–DRG Relative Weight × Geographic Adjustment Factor × (1 + IME + DSH)

   Step 4—Determine operating and capital costs:

   —Operating Costs = (Billed Charges × Operating CCR)
   —Capital Costs = (Billed Charges × Capital CCR)

   Step 5—Compute operating and capital outlier threshold (CMS applies a geographic adjustment to the operating and capital outlier threshold to account for local cost variation):

   —Operating CCR to Total CCR = (Operating CCR)/(Operating CCR + Capital CCR)
   —Operating Outlier Threshold = [(Fixed Loss Threshold × ([Labor–Related Portion × CBSA Wage Index) + Nonlabor–Related Portion]) × Operating CCR to Total CCR + Federal Payment with IME, DSH + Uncompensated Care Payment + New Technology Add–On Payment Amount]

   Step 6—Compute operating and capital outlier payments:

   —Marginal Cost Factor = 0.80 or 0.90 (depending on the MS–DRG)
   —Operating Outlier Payment = (Operating Costs – Operating Outlier Threshold) × Marginal Cost Factor
   —Capital Outlier Payment = (Capital Costs – Capital Outlier Threshold) × Marginal Cost Factor

   The payment rate may then be further adjusted for hospitals that qualify for a low-volume payment adjustment under section 1886(d)(12) of the Act and 42 CFR 412.101(b). The base-operating DRG payment amount may be further adjusted by the hospital readmissions payment adjustment and the hospital VBP payment adjustment as described under sections 1886(g) and 1886(o) of the Act, respectively. Payments also may be reduced by the 1-percent adjustment under the HAC Reduction Program as described in section 1886(p) of the Act. We also make new technology add–on payments in accordance with section 1886(d)(5)(K) and (L) of the Act. Finally, we add the uncompensated care payment to the total claim payment amount. As noted in the formula above, we take uncompensated care payments and new technology add–on payments into consideration when calculating outlier payments.

<table>
<thead>
<tr>
<th>FY 2018</th>
<th>Hospital submitted quality data and is a meaningful EHR user</th>
<th>Hospital submitted quality data and is NOT a meaningful EHR user</th>
<th>Hospital did NOT submit quality data and is a meaningful EHR user</th>
<th>Hospital did NOT submit quality data and is NOT a meaningful EHR user</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proposed Market Basket Rate-of-Increase</td>
<td>2.9</td>
<td>2.9</td>
<td>2.9</td>
<td>2.9</td>
</tr>
<tr>
<td>Proposed Adjustment for Failure to Submit Quality Data under Section 1886(b)(3)(B)(viii) of the Act</td>
<td>0.0</td>
<td>0.0</td>
<td>−0.725</td>
<td>−0.725</td>
</tr>
<tr>
<td>Proposed Adjustment for Failure to be a Meaningful EHR User under Section 1886(b)(3)(B)(ix) of the Act</td>
<td>0.0</td>
<td>−2.175</td>
<td>0.0</td>
<td>−2.175</td>
</tr>
<tr>
<td>Proposed MFP Adjustment under Section 1886(b)(3)(B)(ix) of the Act</td>
<td>−0.4</td>
<td>−0.4</td>
<td>−0.4</td>
<td>−0.4</td>
</tr>
<tr>
<td>Statutory Adjustment under Section 1886(b)(3)(B)(xii) of the Act</td>
<td>−0.75</td>
<td>−0.75</td>
<td>−0.75</td>
<td>−0.75</td>
</tr>
<tr>
<td>Proposed Applicable Percentage Increase Applied to Hospital-Specific Rate</td>
<td>1.75</td>
<td>−0.425</td>
<td>1.025</td>
<td>−1.15</td>
</tr>
</tbody>
</table>

For a complete discussion of the applicable percentage increase applied to the hospital-specific rates for SCHs, we refer readers to section V.B. of the preamble of this proposed rule.

In addition, because SCHs use the same MS–DRGs as other hospitals when they are paid based in whole or in part on the hospital-specific rate, the hospital-specific rate is adjusted by a budget neutrality factor to ensure that changes to the MS–DRG classifications and the recalibration of the MS–DRG relative weights are made in a manner so that aggregate IPPS payments are unaffected. Therefore, the hospital-specific rate for an SCH is adjusted by the proposed MS–DRG reclassification and recalibration.
proposed update factor for FY 2018 under § 412.308(c)(1), to account for capital input price increases and other factors. The regulations at § 412.308(c)(2) also provide that the capital Federal rate be adjusted annually by a factor equal to the estimated proportion of outlier payments under the capital Federal rate to total capital payments under the capital Federal rate. In addition, § 412.308(c)(3) requires that the capital Federal rate be reduced by an adjustment factor equal to the estimated proportion of payments for exceptions under § 412.348. (We note that, as discussed in the FY 2013 IPPS/LTCH PPS final rule (77 FR 53705), there is generally no longer a need for an exceptions payment adjustment factor.) However, in limited circumstances, an additional payment exception for extraordinary circumstances is provided for under § 412.348(f) for qualifying hospitals. Therefore, in accordance with § 412.308(c)(3), an exceptions payment adjustment factor may need to be applied if such payments are made. Section 412.308(c)(3) requires that the capital Federal rate be adjusted so that the effects of the annual DRG reclassification and the recalibration of DRG weights and changes in the geographic adjustment factor (GAF) are budget neutral.

Section 412.374 provides for payments to hospitals located in Puerto Rico under the IPPS for acute care hospital inpatient capital-related costs. In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57066), we revised § 412.374 to add paragraph (e) to provide that, effective with discharges on or after October 1, 2016, capital IPPS payments to hospitals located in Puerto Rico are based on 100 percent of the Federal rate.

A. Determination of the Proposed Federal Hospital Inpatient Capital-Related Prospective Payment Rate Update for FY 2018

In the discussion that follows, we explain the factors that we are proposing to use to determine the capital Federal rate for FY 2018. In particular, we show why the proposed FY 2018 capital Federal rate would increase approximately 1.03 percent, compared to the FY 2017 capital Federal rate. As discussed in the impact analysis in Appendix A to this proposed rule, we estimate that capital payments per discharge would increase approximately 2.4 percent during that same period. Because capital payments constitute approximately 10 percent of hospital payments, a percent change in the capital Federal rate yields only approximately a 0.1 percent change in actual payments to hospitals.

1. Proposed Projected Capital Standard Federal Rate Update

a. Description of the Update Framework

Under § 412.308(c)(1), the capital standard Federal rate is updated on the basis of an analytical framework that takes into account changes in a capital input price index (CPI) and several other policy adjustment factors. Specifically, we adjust the projected CPI rate of change as appropriate each year for case-mix index-related changes, for intensity, and for errors in previous CPI forecasts. The proposed update factor for FY 2018 under that framework is 1.2 percent based on a projected 1.2 percent increase in the proposed 2014-based CPI, a 0.0 percentage point adjustment for intensity, a 0.0 percentage point adjustment for case-mix, a 0.0 percentage point adjustment for the DRG reclassification and recalibration, and a 0.0 percentage point adjustment for a forecast error correction of 0.0 percentage point. As discussed in section III.C of this Addendum, we continue to believe that the CPI is the most appropriate input price index for capital costs to measure capital price changes in a given year. We also explain the basis for the proposed FY 2018 CPI projection in that same section of this Addendum. Below we describe the policy adjustments that we are proposing to apply in the update framework for FY 2018.

The case-mix index is the measure of the average DRG weight for cases paid under the IPPS. Because the DRG weight determines the prospective payment for each case, any percentage increase in the case-mix index corresponds to an equal percentage increase in capital payments.

The case-mix index can change for any of several reasons:

• The average resource use of Medicare patient changes (“real” case-mix change);

• Changes in hospital documentation and coding of patient records result in higher-weighted DRG assignments (“coding effects”); and

• The annual DRG reclassification and recalibration changes may not be budget neutral (“reclassification effect”).

We define real case-mix change as actual changes in the mix of Medicare patients and the mix of resource requirements of Medicare patients as opposed to changes in documentation and coding behavior that result in assignment of cases to higher-weighted DRGs, but do not reflect higher resource requirements. The capital update framework includes the same case-mix index adjustment used in the former operating IPPS update framework (as discussed in the May 18, 2004 IPPS proposed rule for FY 2005 [69 FR 28816]). (We no longer use an update framework to make a reclassification adjustment for FY 2005 (70 FR 47707).) For FY 2018, we are projecting a 0.5 percent total increase in the case-mix index. We estimated that the 0.5 percent increase will equal 0.5 percent for FY 2018. The net adjustment for change in case-mix is the difference between the projected real increase in case-mix and the projected total increase in case-mix. Therefore, the proposed net adjustment for case-mix change in FY 2018 is 0.0 percentage point.

The capital update framework also contains an adjustment for the effects of DRG reclassification and recalibration. This adjustment is intended to remove the effect on total payments of prior year’s changes to the DRG classification and recalibration relative weights, in order to retain budget neutrality for all case-mix index-related changes other than those due to patient severity of illness. Due to the lag time in the availability of data, there is a 2-year lag in data used to determine the adjustment for the effects of DRG reclassification and recalibration. For
example, we have data available to evaluate the effects of the FY 2016 DRG reclassification and recalibration as part of our proposed update for FY 2018. We estimate that FY 2016 DRG reclassification and recalibration resulted in no change in the case-mix when compared with the case-mix index that would have resulted if we had not made the reclassification and recalibration changes to the DRGs. Therefore, we are proposing to make a 0.0 percentage point adjustment for reclassification and recalibration in the update framework for FY 2018.

The capital update framework also contains an adjustment for forecast error. The input price index forecast is based on historical trends and relationships ascertainable at the time the update factor is established for the upcoming year. In any given year, there may be unanticipated price fluctuations that may result in differences between the actual increase in prices and the forecast used in calculating the update factor. The prospective payment rate under the framework, we make an adjustment for forecast error only if our estimate of the change in the capital input price index for any year is off by 0.25 percentage point or more. There is a 2-year lag between the forecast and the availability of data to develop a measurement of the forecast error. Historically, when a forecast error of the CPI is greater than 0.25 percentage point in absolute terms, it is reflected in the update recommended under this framework. A forecast error of 0.2 percentage point was calculated for the FY 2016 update, for which there are historical data. That is, current historical data indicate that the forecasted FY 2016 CPI (1.3 percent) used in calculating the FY 2016 update factor was 0.2 percentage points higher than actual realized price increases (1.1 percent). However, as this does not exceed the 0.25 percentage point threshold, we are proposing not to make an adjustment for forecast error in the update for FY 2018.

Under the IPPS capital update framework, we also make an adjustment for changes in intensity. Historically, we calculated this adjustment using the same methodology and data that were used in the past under the framework for operating IPPS. The intensity factor for the operating update framework reflected how hospital services are utilized to produce the final product, that is, the discharge. This component accounts for changes in the use of quality-enhancing services, for changes within DRG severity, and for expected modification of practice patterns to remove non-cost-effective services. Our intensity measure is based on a 5-year average.

We calculate case-mix constant intensity as the change in total cost per discharge, adjusted for price level changes (the CPI for hospital and related services) and changes in real capital costs. Without reliable estimates of the proportions of the overall annual intensity changes that are due, respectively, to ineffective practice patterns and the combination of quality-enhancing new technologies and complexity within the DRG system, we assume that one-half of the annual change is due to each of these factors.

The capital update framework thus provides an add-on to the input price index rate of increase of one-half of the estimated annual increase in intensity, to allow for increases within DRG severity and the adoption of quality-enhancing technology.

In this proposed rule, we are proposing to continue to use a Medicare-specific intensity measure that is based on a 5-year adjusted average of cost per discharge for FY 2018 (we refer readers to the FY 2011 IPPS/LTCF PPS final rule (75 FR 50436) for a full description of our Medicare-specific intensity measure). Specifically, for FY 2018, we are proposing to use an intensity measure that is based on an average of cost per discharge data from the 5-year period beginning with FY 2011 and extending through FY 2015. Based on these data, we estimated that case-mix constant intensity declined during FYs 2011 through 2015. In the past, when we found intensity to be declining, we believed a zero (rather than a negative) intensity adjustment was appropriate. Consistent with this approach, because we estimate that intensity will decline during that 5-year period, we believe it is appropriate to continue to apply a zero intensity adjustment for FY 2018. Therefore, we are proposing to make a 0.0 percentage point adjustment for intensity in the update for FY 2018.

Above, we described the basis of the components we are proposing to use to develop the proposed 1.2 percent capital update factor under the capital update framework for FY 2018 as shown in the following table.

### CMS Proposed FY 2018 Update Factor to the Capital Federal Rate

<table>
<thead>
<tr>
<th>Factor</th>
<th>FY 2018 Proposed Update Factor to the Capital Federal Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intensity</td>
<td>1.2</td>
</tr>
<tr>
<td>Case-Mix Adjustment</td>
<td>0.0</td>
</tr>
<tr>
<td>Real Across DRG Change</td>
<td>0.5</td>
</tr>
<tr>
<td>Projected Case-Mix Change</td>
<td>0.5</td>
</tr>
<tr>
<td>Total Proposed Update</td>
<td>1.2</td>
</tr>
</tbody>
</table>

* The capital input price index represents the proposed 2014-based CPI.

b. Comparison of CMS and MedPAC Update Recommendation


2. Proposed Outlier Payment Adjustment Factor

Section 412.312(c) establishes a unified outlier payment methodology for inpatient operating and inpatient capital-related costs. A single set of thresholds is used to identify outlier cases for both inpatient operating and inpatient capital-related payments. Section 412.308(c)(2) provides that the standard Federal rate for inpatient capital-related costs be reduced by an adjustment factor equal to the estimated proportion of capital-related outlier payments to total inpatient capital-related PPS payments. The outlier thresholds are set so that operating outlier payments are projected to be 5.1 percent of total operating IPPS DRG payments.

For FY 2017, we estimated that outlier payments for capital would equal 6.14 percent of inpatient capital-related payments based on the capital Federal rate in FY 2017. Based on the thresholds as set forth in section I.A. of this Addendum, we estimate that outlier payments for capital-related costs would equal 5.66 percent for inpatient capital-related payments based on the capital Federal rate in FY 2018. Therefore, we are proposing to apply an outlier adjustment factor of 0.9434 in determining the capital Federal rate for FY 2018. Thus, we estimate that the percentage of capital outlier payments to total capital Federal rate payments for FY 2018 will be lower than the percentage for FY 2017.

3. Proposed Budget Neutrality Adjustment Factor for Changes in DRG Classifications and Weights and the GAF

Section 412.308(c)(4)(ii) requires that the capital Federal rate be adjusted so that aggregate payments for the fiscal year based on the capital Federal rate after any changes resulting from the annual DRG reclassification and recalibration and changes in the GAF are projected to equal aggregate payments that would have been made on the basis of the capital Federal rate without such changes. The budget neutrality factor for DRG reclassifications and recalibrations and changes in the GAF is applied in determining the capital IPPS Federal rate, and is applicable for all hospitals, including those hospitals located in Puerto Rico.

To determine the proposed national capital rate factors for FY 2018, we compared estimated aggregate capital Federal rate payments based on the FY 2017 MS–DRG classifications and relative weights and the FY 2017 GAF to estimated aggregate capital Federal rate payments based on the FY 2017 MS–DRG classifications and relative weights and the proposed FY 2018 GAFs. To achieve budget neutrality for the changes in the national GAFs, based on calculations using updated data, we are proposing to apply an incremental budget neutrality adjustment factor of 0.9997 for FY 2018 to the previous cumulative FY 2017 adjustment factor of 0.9850, yielding an adjustment factor of 0.9847 through FY 2018.

We then compared estimated aggregate capital Federal rate payments based on the FY 2017 MS–DRG relative weights and the
proposed FY 2018 GAFs to estimated aggregate capital Federal rate payments based on the cumulative effects of the proposed FY 2018 MS–DRG classifications and relative weights and the proposed FY 2018 GAFs. The proposed incremental adjustment factor for DRG classifications and changes in relative weights is 0.9994. The proposed cumulative adjustment factor for MS–DRG classifications and changes in relative weights and for changes in the GAFs through FY 2018 is 0.9842. (We note that all the values are calculated with unrounded numbers.)

The GAF/DRG budget neutrality adjustment factors are built permanently into the capital rates; that is, they are applied cumulatively in determining the capital Federal rate. This follows the requirement under § 412.308(c)(4)(iii) that estimated aggregate payments each year be no more or less than they would have been in the absence of the annual DRG reclassification and recalibration and changes in the GAFs.

The methodology used to determine the recalibration and geographic adjustment factor (GAF/DRG) budget neutrality adjustment is similar to the methodology used in establishing budget neutrality adjustments for the IPPS for operating costs. One difference is that, under the operating IPPS, the budget neutrality adjustments for the effect of geographic reclassifications are determined separately from the effects of other changes in the hospital wage index and the MS–DRG relative weights. Under the capital IPPS, there is a single GAF/DRG budget neutrality adjustment factor for changes in the GAF (including geographic reclassification) and the MS–DRG relative weights. In addition, there is no adjustment for the effects that geographic reclassification has on the other payment parameters, such as the payments for DSH or IME.

The proposed cumulative adjustment factor of 0.9992 (the product of the proposed incremental national GAF budget neutrality adjustment factor of 0.9997 and the proposed incremental DRG budget neutrality adjustment factor of 0.9994) accounts for the MS–DRG reclassifications and recalibration and for changes in the GAFs. It also incorporates the effects on the GAFs of FY 2018 geographic reclassification decisions made by the MGCRB compared to FY 2017 decisions. However, it does not account for changes in payments due to changes in the DSH and IME adjustment factors.

As discussed in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57062), we made an adjustment of (1/0.998) to the national capital Federal rate to remove the 0.2 percent reduction (an adjustment factor of 0.998) to the national capital Federal rate to offset the estimated increase in capital IPPS expenditures associated with the 2-midnight policy. This was consistent with the adjustment to the operating IPPS standardized amount and hospital-specific payment rates. In addition, consistent with the approach for the operating IPPS standardized amount and hospital-specific payment rates and for the reasons discussed in the FY 2017 IPPS/LTCH PPS final rule, we made a one-time prospective adjustment of 1.006 in FY 2017 to the national capital Federal rate to address the effect of the 0.2 percent reduction to the national capital Federal rates in effect for FY 2014, FY 2015, and FY 2016. Furthermore, as provided for in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57294) we are removing this one-time prospective adjustment through an adjustment of (1/1.006) to the national capital Federal rate in FY 2018, consistent with the approach for the operating IPPS standardized amount and hospital-specific payment rates (as discussed in section V.M. of the preamble of this proposed rule). We refer readers to sections V.M. and V.LC. of the preamble of this proposed rule for a complete discussion of these issues.

4. Proposed Capital Federal Rate for FY 2018

For FY 2017, we established a capital Federal rate of $446.79 (81 FR 68947 through 68949 (Correction Notice)). We are proposing to establish an update of 1.2 percent in determining the FY 2018 capital Federal rate for all hospitals. As a result of this proposed update, the proposed budget neutrality factor for changes in the GAFs, the national capital Federal rate in FY 2018, consistent with the approach for the operating IPPS standardized amount and hospital-specific payment rates (as discussed in section V.M. of the preamble of this proposed rule). We refer readers to sections V.M. and V.LC. of the preamble of this proposed rule for a complete discussion of these issues.

For FY 2018, we established a capital Federal rate of $451.37 (81 FR 68949). We are proposing an update of 1.02 percent in determining the FY 2018 capital Federal rate for all hospitals. As a result of this proposed update, the proposed budget neutrality factor for changes in the GAFs, the national capital Federal rate in FY 2018, consistent with the approach for the operating IPPS standardized amount and hospital-specific payment rates (as discussed in section V.M. of the preamble of this proposed rule). We refer readers to sections V.M. and V.LC. of the preamble of this proposed rule for a complete discussion of these issues.

<table>
<thead>
<tr>
<th>FY 2017</th>
<th>Proposed FY 2018</th>
<th>Proposed change</th>
<th>Proposed percent change</th>
</tr>
</thead>
<tbody>
<tr>
<td>Update Factor</td>
<td>1.0090</td>
<td>1.0120</td>
<td>0.08</td>
</tr>
<tr>
<td>GAF/DRG Adjustment Factor</td>
<td>0.9990</td>
<td>0.9992</td>
<td>0.08</td>
</tr>
<tr>
<td>Outlier Adjustment Factor</td>
<td>0.9396</td>
<td>0.9434</td>
<td>0.04</td>
</tr>
<tr>
<td>Recalibration and change adjustment Factor</td>
<td>0.9994</td>
<td>1.0060</td>
<td>0.06</td>
</tr>
<tr>
<td>Capital Federal Rate</td>
<td>$446.79</td>
<td>$451.37</td>
<td>1.00</td>
</tr>
</tbody>
</table>

1. The update factor and the GAF/DRG budget neutrality adjustment factors are built permanently into the capital Federal rates. Thus, for example, the proposed incremental change from FY 2017 to FY 2018 resulting from the application of the proposed 0.9992 GAF/DRG budget neutrality adjustment factor for FY 2018 is a proposed net change of 0.9992 (or −0.08 percent).

2. The outlier reduction factor is not built permanently into the capital Federal rate; that is, the factor is not applied cumulatively in determining the capital Federal rate. Thus, for example, the proposed net change resulting from the application of the proposed FY 2018 outlier adjustment factor is 0.9434/0.9386 or 1.0051 (or 0.51 percent).

3. Proposed percent change may not sum due to rounding.
In this proposed rule, we also are providing the following chart that shows how the proposed FY 2018 capital Federal rate differs from the final FY 2017 capital Federal rate as presented in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57291 through 57295) as corrected in the Correction Notice published October 5, 2016 (81 FR 68954).

**COMPARISON OF FACTORS AND ADJUSTMENTS: PROPOSED FY 2018 CAPITAL FEDERAL RATE AND FINAL FY 2017 CAPITAL FEDERAL RATE**

<table>
<thead>
<tr>
<th>Final FY 2017</th>
<th>Proposed FY 2018</th>
<th>Change</th>
<th>Percent change</th>
</tr>
</thead>
<tbody>
<tr>
<td>Update Factor</td>
<td>1.0090</td>
<td>1.0120</td>
<td>1.20</td>
</tr>
<tr>
<td>GAF/DRG Adjust Factor</td>
<td>0.9990</td>
<td>0.9992</td>
<td>-0.08</td>
</tr>
<tr>
<td>Outlier Adjustment Factor</td>
<td>0.9386</td>
<td>1.0051</td>
<td>0.51</td>
</tr>
<tr>
<td>Permanent 2-midnight Policy Adjustment Factor</td>
<td>1.002</td>
<td>1.00</td>
<td>0.00</td>
</tr>
<tr>
<td>One-Time 2-midnight Policy Adjustment Factor</td>
<td>1.006</td>
<td>1.0103</td>
<td>1.03</td>
</tr>
<tr>
<td>Capital Federal Rate</td>
<td>$446.79</td>
<td>$451.37</td>
<td>0.08</td>
</tr>
</tbody>
</table>

**C. Capital Input Price Index**

The CIPI capital price changes needs to reflect this vintage nature of capital. Therefore, the CIPI was developed to capture the vintage nature of capital by using a weighted-average of past capital purchase prices up to and including the current year. We periodically update the base year for the operating input price index to reflect the changing composition of inputs for operating and capital expenses. For this FY 2018 IPPS/LTCH PPS proposed rule, we are proposing to base and revise the IPPS operating and capital market baskets to reflect a 2014 base year. For a complete discussion of this proposed rebasing, we refer readers to section IV. of the preamble of this proposed rule.

**B. Calculation of the Inpatient Capital-Related Prospective Payments for FY 2018**

For purposes of calculating payments for each discharge during FY 2018, the capital Federal rate is adjusted as follows: (Standard Federal Rate) × (DRG weight) × (GAF) × (COLA for hospitals located in Alaska and Hawaii) × (1 + DSH Adjustment Factor + IME Adjustment Factor, if applicable). The result is the adjusted capital Federal rate.

Hospitals also may receive outlier payments for those cases that qualify under the thresholds established for each fiscal year. Section 412.312(c) provides for a single set of thresholds to identify outlier cases for both inpatient and outpatient capital-related payments. The proposed thresholds for FY 2018 are in section I.A. of this Addendum. For FY 2018, a case would qualify as a cost outlier if the cost for the case plus the (operating) IME and DSH payments (including both the empirically justified Medicare DSH payment and the estimated uncompensated care payment, as discussed in section I.A.4.g.(1) of this Addendum) is greater than the prospective payment rate for the MS–DRG plus the fixed-loss amount of $26,713.

Currently, as provided under § 412.304(c)(2), we pay a new hospital 85 percent of its reasonable costs during the first 2 years of operation unless it elects to receive payment based on 100 percent of the capital Federal rate. Effective with the third year of operation, we pay the hospital based on 100 percent of the capital Federal rate (that is, the same methodology used to pay all other hospitals subject to the capital PPS).

**V. Proposed Changes to the Payment Rates**

**A. Proposed LTCH PPS Standard Federal Payment Rate for FY 2018**

In this FY 2018 IPPS/LTCH PPS proposed rule, the proposed FY 2018 rate-of-increase percentage for updating the target amounts for the 11 cancer hospitals, children’s hospitals, the short-term acute care hospitals located in the U.S. Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa, and RNHICs is the estimated percentage increase in the IPPS operating market basket for FY 2018, in accordance with applicable regulations at § 413.40. Based on IHS Global Insight, Inc.’s 2016 fourth quarter forecast, we estimate that the proposed FY 2018 rate-of-increase percentage that would be applied to the FY 2017 target amounts in order to determine the FY 2018 target amounts is 2.9 percent. The IRF PPS, the IFP PPS, and the LTCH PPS are updated annually. We refer readers to section VIII. of the preamble of this proposed rule and section V. of the Addendum to this proposed rule for the proposed update changes to the Federal payment rates for LTCHs under the LTCH PPS for FY 2018. The annual updates for the IRF PPS and the IFP PPS are issued by the agency in separate Federal Register documents.

**Proposed Changes to Payment Rates for Excluded Hospitals: Proposed Rate-of-Increase Percentages for FY 2018**

Payments for services furnished in children’s hospitals, 11 cancer hospitals, and hospitals located outside the 50 States, the District of Columbia and Puerto Rico (that is, short-term acute care hospitals located in the U.S. Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa), and RNHICs, are excluded from the IPPS rate update for FY 2018. Therefore, for children’s hospitals, the 11 cancer hospitals, hospitals located outside the 50 States, the District of Columbia and Puerto Rico (that is, short-term acute care hospitals located in the U.S. Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa), and RNHICs, the proposed FY 2018 rate-of-increase percentage that would be applied to the FY 2017 target amounts in order to determine the FY 2018 target amounts is 2.9 percent.
market basket of goods and services for LTCHs. We established this policy of annually updating the standard Federal payment rate because, at that time, we believed that was the most appropriate method for updating the rate for years after the initial implementation of the LTCH PPS in FY 2003. Therefore, under § 412.523(c)(3)(ii), for RYs 2004 through 2006, the annual update to the LTCH PPS standard Federal payment rate was equal to the previous rate year’s Federal rate updated by the most recent estimate of increases in the appropriate market basket of goods and services included in covered inpatient LTCH services.

In determining the annual update to the standard Federal payment rate for FY 2007, based on our ongoing monitoring activity, we believed that, rather than solely using the most recent estimate of the LTCH PPS market basket update as the basis of the annual update factor, it was appropriate to adjust the standard Federal payment rate to account for the effect of documentation and coding in a prior period that was unrelated to patients’ severity of illness (71 FR 27818). Accordingly, we established under § 412.523(c)(3)(iii) that the annual update to the standard Federal payment rate for FY 2007 was zero percent based on the most recent estimate of the LTCH PPS market basket update at that time, offset by an adjustment to account for changes in case-mix in prior periods due to the effect of documentation and coding that were unrelated to patients’ severity of illness. For FY 2008 through FY 2011, we also made an adjustment to account for the effect of documentation and coding that was unrelated to patients’ severity of illness in establishing the annual update to the standard Federal payment rate as set forth in the regulations at § 412.523(c)(3)(iv) through (c)(3)(vii). For FYs 2012 through 2017, we updated the standard Federal payment rate by the most recent estimate of the LTCH PPS market basket at that time, including additional statutory adjustments required by sections 1886(m)(3)(A)(iii), (iv) (citing section 1886(m)(3)(A)(ii)), 1886(m)(3)(A)(iii), and 1886(m)(4) of the Act as set forth in the regulations at § 412.523(c)(3)(vii) through (c)(3)(xiii)).

Section 1886(m)(3)(A) of the Act, as added by section 3401(c) of the Affordable Care Act, specifies that, for rate year 2010 and each subsequent rate year, any annual update to the standard Federal payment rate shall be reduced:

• For rate year 2010 through 2019, by the other adjustment specified in section 1886(m)(3)(A)(ii) and (m)(4) of the Act; and
• For rate year 2012 and each subsequent rate year, by the productivity adjustment described in section 1886(b)(3)(B)(ii)(I) of the Act (which we refer to as “the multifactor productivity (MFP) adjustment”) as discussed in section VIII.E.2. of the preamble of this proposed rule.

Section 1886(m)(3)(B) of the Act provides that the application of paragraph (3) of section 1886(m) of the Act may result in the annual update being less than zero for a rate year, and may result in payment rates for a rate year being less than such payment rates for the preceding rate year. (As noted in section VIII.C.2.h. of the preamble of this proposed rule, the annual update to the LTCH PPS occurs on October 1 and we have adopted the term “fiscal year” (FY) rather than “rate year” (RY) under the LTCH PPS beginning October 1, 2010. Therefore, for purposes of this proposed rule, we are discussing the annual update for the LTCH PPS, including the provisions of the Affordable Care Act, we use the term “fiscal year” rather than “rate year” for 2011 and subsequent years.)

Notwithstanding those provisions, however, under section 114–10 (the MACRA) requires a 1.0 percent update in FY 2018.

For FY 2017, consistent with our historical practice, we established an update to the LTCH PPS standard Federal payment rate based on the full estimated LTCH PPS market basket increase of 2.8 percent and the 1.05 percent point reductions required by section 1886(m)(3)(A)(ii) and 1886(m)(3)(A)(ii) with 1886(m)(4)(F) of the Act. Accordingly, at § 412.523(c)(3)(xii) of the regulations, we established an annual update of 1.75 percent to the standard Federal payment rate for FY 2017 (81 FR 57296 through 57297). In addition, as discussed in that same final rule, the annual update for FY 2017 was further reduced by 2.0 percentage points for LTCHs that failed to submit quality reporting data in accordance with the requirements of the LTCH QRP under section 1886(m)(5) of the Act.

Section 411(e) of the MACRA amended section 1886(m)(3) of the Act by providing an additional special rule for FY 2018. Specifically, as amended, section 1886(m)(3)(C) of the Act requires that the annual update for FY 2018, after applications of the reductions for the MFP adjustment and the “other adjustment” (under section 1886(m)(3)(A)) is 1 percent. (For additional details, refer to section VIII.E.2. of the preamble of this proposed rule.) Accordingly, in this proposed rule, we are proposing an annual update for the LTCH PPS standard Federal payment rate of 1.0 percent for FY 2018 as required by section 1886(m)(3) of the Act. Accordingly, we are proposing an annual update to the LTCH PPS standard Federal payment rate of −1.0 percent for FY 2018 to reflect the required quality reporting data for FY 2017 in accordance with the LTCH QRP, the annual update is reduced by 2.0 percentage points as required by section 1886(m)(5) of the Act.

Development of the Proposed FY 2018 LTCH PPS Standard Federal Payment Rate

Consistent with our historical practice, for FY 2018, we are proposing to apply the annual update to the LTCH PPS standard Federal payment rate from the previous year. Furthermore, in determining the proposed LTCH PPS standard Federal payment rate for FY 2018, we also are proposing to make certain regulatory adjustments, consistent with past practices. Specifically, in determining the proposed FY 2018 LTCH PPS standard Federal payment rate, we are proposing to apply a budget neutrality adjustment factor for the proposed changes related to the area wage adjustment (that is, proposed changes to the wage data and proposed labor-related share) in accordance with § 412.523(d)(4) and a proposed budget neutrality adjustment factor for the proposed change to the SSO payment methodology (discussed in VIII.D. of the preamble of this proposed rule).

For FY 2017, we established an annual update to the LTCH PPS standard Federal payment rate of 1.75 percent based on the full estimated LTCH PPS market basket increase of 2.8 percent, less the MFP adjustment of 0.3 percentage point consistent with section 1886(m)(3)(A)(i) of the Act and less the 0.75 percentage point required by sections 1886(m)(3)(A)(ii) and (m)(4)(F) of the Act. Accordingly, at § 412.523(c)(3)(xii), we established an annual update to the LTCH PPS standard Federal payment rate for FY 2017 of 1.75 percent. That is, we applied an update factor of 1.0175 to the FY 2016 LTCH PPS Federal rate of $41,762.85 to determine the FY 2017 LTCH PPS standard Federal payment rate. We also applied an area wage level budget neutrality factor for FY 2017 of 0.999593 to the LTCH PPS standard Federal payment rate to ensure that any changes to the area wage level adjustment would not result in any change in estimated aggregate LTCH PPS payments. Consequently, we established an LTCH PPS standard Federal payment rate for FY 2017 of $42,476.41 (calculated as $41,762.85 × 1.0175 × 0.999593) (81 FR 57296 through 57297).

In this proposed rule, as required by statute, we are proposing an annual update to the LTCH PPS standard Federal payment rate of 1.0 percent for FY 2018 (as described above). Accordingly, under § 412.523(c)(3)(xii), we are proposing to apply a factor of 1.01 to the FY 2017 LTCH PPS standard Federal payment rate of $42,476.41 to determine the proposed FY 2018 LTCH PPS standard Federal payment rate. Also, under proposed § 412.523(c)(3)(xii), in conjunction with the provisions of § 412.523(d)(4), we are proposing to apply an annual update to the LTCH PPS standard Federal payment rate of −1.0 percent (that is, a proposed update factor of 0.99) for FY 2018 for LTCHs that fail to submit the required quality reporting data for FY 2017 in accordance with the LTCH QRP. The annual update is reduced by 2.0 percentage points for LTCHs that fail to submit quality reporting data as required by section 1886(m)(5) of the Act.

Consistent with § 412.523(d)(4), we also are proposing to apply an area wage level budget neutrality factor to the proposed FY 2018 LTCH PPS standard Federal payment rate of 1.000077, based on the best available wage index data at this time, to ensure that any proposed changes to the area wage level adjustment (that is, the proposed annual update of the wage index values and labor-related share) would not result in any change (increase or decrease) in estimated aggregate LTCH PPS payments. Specifically, in determining the proposed FY 2018 LTCH PPS standard Federal payment rate, we are proposing to apply a budget neutrality adjustment factor for the proposed changes related to the area wage adjustment (that is, proposed changes to the wage data and proposed labor-related share) in accordance with § 412.523(d)(4) and a proposed budget neutrality adjustment factor for the proposed change to the SSO payment methodology (discussed in VIII.D. of the preamble of this proposed rule). Accordingly, we are proposing an LTCH PPS standard Federal payment rate of $41,497.20 (calculated as $42,476.41 × 1.01 × 1.000077...
Under the authority of section 123 of the BBRA, an area delineation as defined in section 307(b) of the BIPA, we established an adjustment to the LTCH PPS standard Federal payment rate to account for differences in LTCH area wage levels used under § 412.525(c). The labor-related share of the LTCH PPS standard Federal payment rate is adjusted to account for geographic differences in area wage levels by applying the applicable LTCH PPS wage index. The applicable LTCH PPS wage index is computed using wage data from inpatient acute care hospitals without regard to reclassification under section 1886(d)(8) or section 1886(d)(10) of the Act.

When we implemented the LTCH PPS, we established a 5-year transition to the full area wage level adjustment. The area wage level adjustment was completely phased-in for cost reporting periods beginning in FY 2003. Therefore, for cost reporting periods beginning on or after October 1, 2006, the applicable LTCH area wage index values are the full LTCH PPS area wage index values calculated based on acute care hospital inpatient wage index data without taking into account geographic reclassification under section 1886(d)(8) and section 1886(d)(10) of the Act. For additional information on the phase-in of the area wage level adjustment under the LTCH PPS, we refer readers to the August 30, 2002 LTCH PPS final rule (67 FR 56015 through 56019) and the FY 2006 LTCH PPS Final Rule (71 FR 51808).

2. Proposed Geographic Classifications (Labor Market Areas) for the LTCH PPS Standard Federal Payment Rate

In adjusting for the differences in area wage levels under the LTCH PPS, the labor-related portion of an LTCH’s Federal prospective payment adjustment is determined using the geographic classification (labor market area) in which the LTCH is located. Specifically, the application of the LTCH PPS wage level adjustment under existing § 412.525(c) is made based on the location of the LTCH—either in an “urban area” or a “rural area,” as defined in § 412.503, and the LTCH wage level adjustment most appropriately accounts for and reflects the relative hospital wage levels in the geographic area of the hospital as compared to the national average hospital wage level.

We noted that this policy was consistent with the IPPS policy adopted in FY 2015 under § 412.64(b)(1)(ii)(D) of the regulations (79 FR 49951 through 499963). For additional information on the CBSA-based labor market area geographic classification delineations currently used under the LTCH PPS and the history of the labor market area definitions used under the LTCH PPS, we refer readers to the FY 2015 IPPS/LTCH PPS final rule (79 FR 50180 through 50185).

In general, it is our historical practice to update the CBSA-based labor market area delineations annually based on the most recent updates issued by OMB. Generally, OMB issues major revisions to statistical areas every 10 years, based on the results of the decennial census. However, OMB occasionally issues updates and revisions to statistical areas in the years between the decennial censuses. As discussed in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56913 through 56914), OMB issued OMB Bulletin No. 15–01 on July 15, 2015 to update and supersede Bulletin No. 13–10. Bulletin No. 15–01 and its attachment provide detailed information on the update to statistical areas since the February 28, 2013 release of Bulletin No. 13–10 and are based on the application of the 2010 Standards for Metropolitan and Micropolitan Statistical Areas to Census Bureau population estimates for July 1, 2012, and July 1, 2013. A copy of this bulletin may be obtained on the CMS Web site at: https://www.obamawhitehouse.archives.gov/sites/default/files/omb/bulletins/2015/15-01.pdf.

We believe that these revisions to the CBSA-based labor market area delineations will ensure that the LTCH PPS area wage level adjustment most appropriately accounts for and reflects the relative hospital wage levels in the geographic area of the hospital as compared to the national average hospital wage level based on the best available data that reflect the local economies and area wage levels of the hospitals that are currently located in these geographic areas (81 FR 57298).

Therefore, we are proposing to continue to use the LTCH PPS wage levels under the LTCH PPS area wage level adjustments adopted under the LTCH PPS, effective October 1, 2017 (as adopted in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57298)). Moreover, the proposed FY 2018 LTCH PPS wage index values in Tables 12A and 12B listed in section VI. of the Addendum of this proposed rule (which are available via the Internet on the CMS Web site) reflect the revisions to the CBSA-based labor market area delineations described above. We note that, as discussed in section III.A.2. of the preamble of this proposed rule, the revisions to the CBSA-based delineations adopted under the IPPS, effective beginning October 1, 2016.

3. Proposed Labor-Related Share for the LTCH PPS Standard Federal Payment Rate

Under the payment adjustment for the differences in area wage levels under § 412.525(c), the labor-related share of an LTCH’s standard Federal payment rate adjustment is calculated based on the applicable LTCH market basket. The labor-related share currently represents the sum of the labor-related portion of operating costs (Wages and Salaries; Employee Benefits; Professional Fees Labor-Related; Administrative and Business Support Services; and All-Other: Labor-Related Services) and a labor-related portion of capital costs using the applicable LTCH PPS market basket. Additional background information on the historical development of the labor-related share under the LTCH PPS can be found in the FY 2007 LTCH PPS final rule (71 FR 27810 through 27817 and 27829 through 27830) and the FY 2012 IPPS/LTCH PPS final rule (76 FR 51766 through 51769 and 51808).

For FY 2013, we revised and rebased the market basket used under the LTCH PPS by adopting the newly created FY 2009-based LTCH-specific market basket. In addition, beginning in FY 2013, we determined the labor-related share annually as the sum of the relative importance of each labor-related cost category of the 2009-based LTCH-specific market basket for the respective fiscal year based on the best available data. For more details, we refer readers to the FY 2013 IPPS/ LTCH PPS final rule (77 FR 53477 through 53479). As noted previously, we rebased and revised the 2009-based LTCH-specific market basket to reflect a 2013 base year.

In conjunction with that policy, as discussed in section VIII.C. of the preamble of this proposed rule, we are proposing to establish that the LTCH PPS labor-related share for FY 2018 is the sum of the FY 2018 relative importance of each labor-related cost category in the 2013-based LTCH market basket using the most recent available data. Specifically, we are proposing to establish that the labor-related share for FY 2018 would include the sum of the labor-related portion of operating costs from the 2013-based LTCH market basket (that is, the sum of the FY 2018 relative importance share of Wages and Salaries; Employee Benefits; Professional Fees; Labor-Related; Administrative and Facilities Support Services; Installation, Maintenance, and Repair Services; All Other: Labor-related Services) and a portion of the Capital-related costs from the 2013-based LTCH market basket. Based on IGI’s fourth quarter 2016 forecast of the 2013-based LTCH market basket, we are proposing to establish a labor-related share under the LTCH PPS for FY 2018 of 66.3 percent. This labor-related share is determined using the same methodology as employed in calculating all previous LTCH.
PPS labor-related shares. Consistent with our historical practice, we also are proposing that if more recent data become available, we would use that data, if appropriate, to determine the final FY 2018 labor-related share in the final rule.

The proposed labor-related share for FY 2018 is the sum of the FY 2018 relative importance of each labor-related cost category, and would reflect the different rates of price change for these cost categories between the base year (2013) and FY 2018. The sum of the relative importance for FY 2018 is concentrated on the labor market (Wages and Salaries; Employee Benefits; Professional Fees; Labor-Related; Administrative and Facilities Support Services; Installation, Maintenance, and Repair Services; All Other: Labor-Related Services) is 62.1 percent. The portion of capital-related costs that is influenced by the local labor market is estimated to be 46 percent (the same percentage applied to the 2009-based LTCH-specific market basket).

Because the relative importance for capital-related costs under our policies is 9.2 percent of the 2013-based LTCH market basket in FY 2018, we are proposing to take 46 percent of 9.2 percent to determine the labor-related share of capital-related costs for FY 2018 (0.46 x 9.2). The result is 4.2 percent, which we added to 62.1 percent for the operating cost amount to determine the total proposed labor-related share for FY 2018. Therefore, we are proposing that the labor-related share under the LTCH PPS for FY 2018 is 66.3 percent.

4. Proposed Wage Index for FY 2018 for the LTCH PPS Standard Federal Payment Rate

Historically, we have established LTCH PPS area wage index values calculated from acute care IPPS hospital wage data without taking into account geographic reclassification under sections 1886(d)(8) and 1886(d)(10) of the Act (67 FR 56019).

The area wage level adjustment established under the LTCH PPS is based on an LTCH’s actual location without regard to the “urban” or “rural” designation of any related or affiliated hospital.

In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57299 through 57301), we calculated the FY 2017 LTCH PPS area wage index values using the same data used for the FY 2017 acute care hospital IPPS (that is, data from cost reporting periods beginning during FY 2013), without taking into account geographic reclassification under sections 1886(d)(8) and 1886(d)(10) of the Act, as these were the most relevant complete data available at that time. In that same final rule, we indicated that we computed the FY 2017 LTCH PPS area wage index values, consistent with the urban and rural geographic classifications (labor market areas) that were in place at that time and consistent with the pre-reclassified IPPS wage index policy (that is, our historical policy of not taking into account geographic reclassifications in determining payments under the LTCH PPS).

As with the IPPS wage index, wage data for multicampus hospitals with campuses located in different labor market areas (CBSAs) are apportioned to each CBSA where the campus (or campuses) are located. We also continued to use our existing policy for determining area wage index values for areas where there are no IPPS wage data.

Consistent with our historical methodology, as discussed in this FY 2018 IPPS/LTCH PPS proposed rule, to determine the applicable area wage index values for the FY 2018 LTCH PPS beginning Federal payment rate, under the broad authority of section 123 of the BBRA, as amended by section 307(b) of the BIPA, we are proposing to use wage data collected from cost reports submitted by IPPS hospitals for cost reporting periods beginning during FY 2014, without taking into account geographic reclassification under sections 1886(d)(8) and 1886(d)(10) of the Act, because these data are the most recent complete data available. We also note that these are the same data we are using to compute the FY 2018 acute care hospital inpatient wage index, as discussed in section III. of the preamble of this proposed rule. We are proposing to compute the proposed FY 2018 LTCH PPS standard Federal payment rate area wage index values consistent with the “urban” and “rural” geographic classifications (that is, labor market area delineations, including the proposed updates, as previously discussed in section V.B. of this Addendum) and our historical policy of not taking into account IPPS geographic reclassifications under sections 1886(d)(8) and 1886(d)(10) of the Act in determining payments under the LTCH PPS. We also are proposing to continue to apportion wage data for multicampus hospitals with campuses located in different labor market areas to each CBSA where the campuses are located, consistent with the IPPS policy.

Lastly, consistent with our existing methodology for determining the LTCH PPS wage index values, for FY 2018, we are proposing to continue to use our existing policy for determining area wage index values for areas where there are no IPPS wage data. Under our existing methodology, the LTCH PPS wage index value for urban CBSAs with no IPPS wage data would be determined by using an average of all of the urban areas in the State where the LTCH PPS wage index value for rural areas with no IPPS wage data would be determined by using the unweighted average of the wage indices from all of the CBSAs that are contiguous to the rural counties of the State. Based on the FY 2014 IPPS wage data that we are proposing to use to determine the proposed FY 2018 LTCH PPS standard Federal payment rate area wage index values in this proposed rule, there are no IPPS wage data for the urban area of Hinesville, GA (CBSA 25980). Consistent with the methodology discussed above, we calculated the proposed FY 2018 wage index value for CBSA 25980 as the average of the wage index values for all of the other urban areas within the state of Georgia (that is, CBSAs 10500, 12020, 12060, 12260, 15260, 16860, 17980, 19240, 24750, 42340, 42580, 43540, 46160, 46460, 46750, and 47580), as shown in Table 12A, which is listed in section VI. of the Addendum to this proposed rule and available via the Internet on the CMS Web site. We note that, as IPPS wage data are dynamic, it is possible that urban areas without IPPS wage data will vary in the future.

Based on the FY 2014 IPPS wage data that we are proposing to use to determine the proposed FY 2018 LTCH PPS standard Federal payment rate area wage index values in this proposed rule, there are no rural areas without IPPS hospital wage data. Therefore, it is not necessary to use our established methodology to calculate a proposed LTCH PPS standard Federal payment rate wage index value for proposed rural areas with no IPPS wage data for FY 2018. We note that, as IPPS wage data are dynamic, it is possible that the number of rural areas without IPPS wage data will vary in the future.

The proposed FY 2018 LTCH PPS standard Federal payment rate wage index values that would be applicable for LTCH PPS standard Federal payment rate changes occurring on or after October 1, 2017, through September 30, 2018, are presented in Table 12A (for urban areas) and Table 12B (for rural areas), which are listed in section VI. of the Addendum of this proposed rule and available via the Internet on the CMS Web site.

5. Proposed Budget Neutrality Adjustment for Proposed Changes to the LTCH PPS Standard Federal Payment Rate Area Wage Level Adjustment

Historically, the LTCH PPS wage index and labor-related share are updated annually based on the latest available data. Under § 412.525(c)(4), any changes to the area wage index values or labor-related shares are to be made in a budget neutral manner such that estimated aggregate LTCH PPS payments are unaffected; that is, will be neither greater than nor less than estimated aggregate LTCH PPS payments without such changes to the area wage level adjustment. Under this policy, we determine an area wage-level adjustment budget neutrality factor that will be applied to the standard Federal payment rate to ensure that any changes to the area wage level adjustments are budget neutral such that any changes to the area wage index values or labor-related share would not result in any change (increase or decrease) in estimated aggregate LTCH PPS payments.

Accordingly, under § 412.525(c)(4), we apply an area wage level adjustment budget neutrality factor in determining the standard Federal payment rate, and also established a methodology for calculating an area wage level adjustment budget neutrality factor. (For additional information on the establishment of our budget neutrality policy for changes to the area wage level adjustment, we refer readers to the FY 2012 IPPS/LTCH PPS final rule (76 FR 51771 through 51773 and 51809).)

In this proposed rule, for FY 2018 LTCH PPS standard Federal payment rate cases, in accordance with § 412.525(c)(4), we are proposing to apply an area wage level adjustment budget neutrality factor to adjust the LTCH PPS standard Federal payment rate to account for the estimated effect of the proposed changes to the area wage level adjustment under § 412.525(c)(1) on estimated aggregate LTCH PPS payments using a methodology that is consistent with the methodology we established in the FY 2012 IPPS/LTCH PPS final rule (76 FR 51771). Specifically, we are proposing to determine an area wage level adjustment...
budget neutrality factor that would be applied to the LTCH PPS standard Federal payment rate under § 412.523(d)(4) for FY 2018 using the following methodology:

Step 1—We estimated aggregate LTCH PPS standard Federal payment rate payments using the FY 2017 wage index values and the FY 2017 labor-related share of 66.5 percent (as established in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57099 and 57100)).

Step 2—We estimated aggregate LTCH PPS standard Federal payment rate payments using the proposed FY 2018 wage index values (as shown in Tables 12A and 12B listed in the Addendum to this proposed rule and available via the Internet on the CMS Web site) and the proposed FY 2018 labor-related share of 66.3 percent (based on the latest available data as previously discussed in this Addendum).

Step 3—We calculated the ratio of these estimated total LTCH PPS standard Federal payment rate payments by dividing the estimated total LTCH PPS standard Federal payment rate payments using the proposed FY 2018 area wage level adjustments (calculated in Step 1) by the estimated total LTCH PPS standard Federal payment rate payments using the proposed FY 2018 area wage level adjustments (calculated in Step 2) to determine the proposed area wage level adjustment budget neutrality factor for FY 2018 LTCH PPS standard Federal payment rate payments.

Step 4—We then applied the proposed FY 2018 area wage level adjustment budget neutrality factor from Step 3 to determine the proposed FY 2018 LTCH PPS standard Federal payment rate after the application of the proposed FY 2018 annual update (discussed previously in section V.A. of this Addendum).

We note that, with the exception of cases subject to the transitional blend payment rate provisions in the first 2 years and certain temporary exemptions for certain spinal cord specialty hospitals and certain severe wound cases, all the FY 2017 LTCH PPS standard Federal payment rate, only LTCH PPS cases that meet the statutory criteria to be excluded from the site neutral payment rate (that is, LTCH PPS standard Federal payment rate cases) are paid based on the LTCH PPS standard Federal payment rate. Because the area wage level adjustment under § 412.525(c) is an adjustment to the LTCH PPS standard Federal payment rate, we only used data from claims that would have qualified for payment at the LTCH PPS standard Federal payment rate if such rate had been in effect at the time of discharge to calculate the FY 2017 LTCH PPS standard Federal payment rate area wage level adjustment budget neutrality factor described above.

For this proposed rule, using the steps in the methodology previously described, we determined a proposed FY 2018 LTCH PPS standard Federal payment rate area wage level adjustment budget neutrality factor of 1.000077. Accordingly, in section V.A. of the Addendum to this proposed rule, to determine the proposed FY 2018 LTCH PPS standard Federal payment rate, we are proposing to apply an area wage level adjustment budget neutrality factor of 1.000077, in accordance with § 412.523(d)(4). The proposed FY 2018 LTCH PPS standard Federal payment rate shown in Table 1E of the Addendum to this proposed rule reflects this adjustment factor.

C. Proposed Cost-of-Living Adjustment (COLA) for LTCHs Located in Alaska and Hawaii

Under § 412.525(b), a cost-of-living adjustment (COLA) is provided for LTCHs located in Alaska and Hawaii to account for the higher costs incurred in those States. Specifically, we apply a COLA to payments to LTCHs located in Alaska and Hawaii by multiplying the nonlabor-related portion of the standard Federal payment rate by the applicable COLA factors established annually by CMS. Higher labor-related costs for LTCHs located in Alaska and Hawaii are taken into account in the area wage level area wage levels previously described.

Under our current methodology, we update the COLA factors for Alaska and Hawaii every 4 years (at the same time as the update to the labor-related share of the IPPS market basket) (77 FR 53712 through 53713). This methodology is based on a comparison of the growth in the Consumer Price Indexes (CPIs) for Anchorage, Alaska, and Honolulu, Hawaii, relative to the growth in the CPI for the average U.S. city as published by the Bureau of Labor Statistics (BLS). It also includes a 25-percent cap on the CPI-updated COLA factors. Under our current policy, we update the COLA factors using the methodology described above every 4 years; the first year began in FY 2014. For FY 2014, we updated the COLA factors for Alaska and Hawaii published by OPM for 2009 using the methodology finalized in FY 2013. (For additional details on our current methodology for updating the COLA factors for Alaska and Hawaii, we refer readers to the FY 2013 IPPS/LTCH PPS final rule (77 FR 50985 through 50987), when developing the FY 2014 IPPS/LTCH PPS final rule, we continue to believe that determining updated COLA factors using this methodology would appropriately adjust the nonlabor-related portion of the LTCH PPS standard Federal payment rate for LTCHs located in Alaska and Hawaii.

For FY 2018, we are proposing to continue to update the COLA factors published by OPM for 2009 (as these are the last COLA factors OPM published prior to transitioning from COLAs to locality pay) using the methodology that we finalized in the FY 2013 IPPS/LTCH PPS final rule and implemented for the FY 2014 IPPS update. Specifically, we are proposing to update the 2009 OPM COLA factors by a comparison of the growth in the Consumer Price Indices (CPIs) for Anchorage, Alaska, and Honolulu, Hawaii, relative to the growth in the CPI for the average U.S. city as published by the Bureau of Labor Statistics (BLS). Because BLS publishes CPI data for only Anchorage and Honolulu, using the methodology we finalized in the FY 2013 IPPS/LTCH PPS final rule, we use the comparison of the growth in the overall CPI relative to the growth in the CPI for those cities to update the COLA factors for all areas in Alaska and Hawaii, respectively. We believe that the relative price differences between these cities and the U.S. (as measured by the CPIs mentioned above) are appropriate proxies for the relative price differences between the “other areas” of Alaska and Hawaii and the United States.

BLS publishes the CPI for All Items for Anchorage, Honolulu, and for the average U.S. city. However, consistent with our methodology finalized in the FY 2014 IPPS/LTCH PPS final rule, we are creating reweighted CPIs for each of the respective areas to reflect the underlying composition of the IPPS market basket nonlabor-related share. The current composition of the CPI for All Items for the respective areas is approximately 40 percent commodities and 60 percent services. However, the IPPS nonlabor-related share is comprised of a different mix of commodities and services. Therefore, we create reweighted indexes for Anchorage, Honolulu, and the average U.S. city using the respective CPI commodities index and CPI services index (at the national level) and then reweight these indexes to approximate 55 percent commodities/45 percent services shares shares obtained from the proposed 2014-based IPPS market basket. We create reweighted indexes using BLS data for 2009 through 2016—the most recent data available at the time of this proposed rulemaking. In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50985 through 50987), we created reweighted indexes based on the FY 2010-based IPPS market basket (which was adopted for the FY 2014 update) and BLS data for 2009 through 2012 (the most recent BLS data at the time of the FY 2014 IPPS/LTCH PPS rulemaking).

We continue to believe this methodology is appropriate because we continue to make a COLA for LTCHs located in Alaska and Hawaii by multiplying the nonlabor-related portion of the LTCH PPS standard Federal rate by a COLA factor. We note that OPM’s COLA factors were calculated with a statutorily mandated cap of 25 percent. As stated in the FY 2014 IPPS/LTCH PPS final rule (78 FR 50987), when developing the COLA update methodology we finalized in the FY 2013 IPPS/LTCH final rule, we exercised our discretionary authority to adjust payments to LTCHs in Alaska and Hawaii by incorporating this cap. In applying this finalized methodology for updating the COLA factors, our proposal for FY 2018 continues to use a 25-percent cap, as our proposal is based on OPM’s COLA factors (updated by the methodology described earlier).

Applying this methodology, the COLA factors that we are proposing to establish for FY 2018 to adjust the nonlabor related portion of the LTCH PPS standard Federal rate for LTCHs located in Alaska and Hawaii are shown in the table below. For comparison purposes, we also are showing the FY 2013 COLA factors (which were based on OPM’s published COLA factors for 2009) and the COLA factors for FYs 2014 through 2017.
We note that the reweighted CPI for Honolulu, HI grew faster than the reweighted CPI for the average U.S. city over the 2009 to 2016 time period at 13.7 percent and 10.5 percent, respectively. As a result, for FY 2018, we calculated COLA factors for the City and County of Honolulu, County of Kauai, and County of Maui and County of Kalawao to be 1.29 compared to the FY 2013 COLA factor of 1.25 (which were based on OPM’s published COLA factors for 2009, as described above). However, as stated above, we are applying our methodology as finalized in the FY 2013 IPPS/LTCPPS final rule to incorporate a cap of 1.25 for these areas. In addition, the proposed COLA factor we calculated for the County of Hawaii for FY 2018 is 1.21 compared to the FY 2013 COLA factor of 1.18. The COLA factors adopted in FY 2014 using this same methodology can be found in the table above.

Similarly, the reweighted CPI for Anchorage, AK grew faster than the reweighted CPI for the average U.S. city over the 2009 to 2016 time period, at 12.4 percent and 10.5 percent, respectively. As a result, for FY 2018, we calculated proposed COLA factors for the City of Anchorage, City of Fairbanks, and City of Juneau to be 1.25 compared to the FY 2013 COLA factor of 1.23. For FY 2018, we calculated a proposed COLA factor of 1.27 for the Rest of Alaska compared to the FY 2013 COLA factor of 1.25. However, as stated above, we are applying our methodology as finalized in the FY 2013 IPPS/LTCPPS final rule to incorporate a cap of 1.25 for the rest of Alaska.

As stated above, the COLA factors adopted in the FY 2014 IPPS/LTCPPS final rule were based on the same methodology used to determine the proposed FY 2018 COLA factors but utilizing BLS data from 2009 through 2012 (the most recent data available at the time of the FY 2014 rulemaking) rather than through 2016 (the most recent data available at the time of this rulemaking). Compared to the FY 2014 COLA factors, the proposed FY 2018 COLA factors are higher—with all areas either reaching or exceeding the cap of 1.25 except the County of Hawaii.

D. Proposed Adjustment for LTCH PPS High-Cost Outlier (HCO) Cases

1. HCO Background

From the beginning of the LTCH PPS, we have included an adjustment to account for cases in which there are extraordinarily high costs relative to the costs of most discharges. Under this policy, additional payments are made based on the degree to which the estimated cost of a case (which is calculated by multiplying the Medicare allowable covered charge by the hospital’s overall hospital CCR) exceeds a fixed-loss amount. This policy results in greater payment accuracy under the LTCH PPS and the Medicare program, and the LTCH sharing the financial risk for the treatment of extraordinarily high-cost cases.

We retained the basic tenets of our HCO policy in FY 2018 when we implemented the dual rate LTCH PPS payment structure under section 1206 of Public Law 113–67. LTCH discharges that meet the criteria for exclusion from the site neutral payment rate (that is, LTCH PPS standard Federal payment rate cases) are paid at the LTCH PPS standard Federal payment rate, which includes, as applicable, HCO payments under § 412.523(e). LTCH discharges that do not meet the criteria for exclusion are paid at the site neutral payment rate, which includes, as applicable, HCO payments under § 412.522(c)(2)(i). In this rule, we established separate fixed-loss amounts and targets for the different LTCH PPS payment rates. Under this bifurcated policy, the historic 8 percent HCO target was retained for LTCH PPS standard Federal payment rate cases, with the fixed-loss amount calculated using only data from LTCH cases that would have been paid at the LTCH PPS standard Federal payment rate if that rate had been in effect at the time of those discharges. For site neutral payment rate cases, we adopted the operating IPPS HCO target (currently 5.1 percent) and set the fixed-loss amount for site neutral payment rate cases at the value of the IPPS fixed-loss amount. Under the HCO policy for both payment rates, an LTCH receives 80 percent of the difference between the estimated cost of the case and the applicable HCO threshold, which is the sum of the LTCH PPS payment rate for the case and the applicable fixed-loss amount for such case.

In order to maintain budget neutrality, consistent with the budget neutrality requirement for HCO payments to LTCH PPS standard Federal payment rate cases, we also adopted a budget neutrality requirement for HCO payments to site neutral payment rate cases by applying a budget neutrality factor to the LTCH PPS payment for those site neutral payment rate cases. (We refer readers to § 412.522(c)(2)(i) of the regulations for further details.) We note that, during the 2-year transitional period, the site neutral payment rate in HCO budget neutrality factor did not apply to the LTCH PPS standard Federal payment rate portion of the blended rate at § 412.522(c)(3) payable to site neutral payment rate cases. (For additional details on the HCO policy adopted for site neutral payment rate cases under the dual rate LTCH PPS payment structure, including the budget neutrality adjustment for HCO payments to site neutral payment rate cases, we refer readers to the FY 2016 IPPS/LTCPPS final rule (80 FR 49617 through 49623.).)

2. Determining LTCH CCRs Under the LTCH PPS

a. Background

As noted above, CCRs are used to determine payments for HCO adjustments for both payment rates under the LTCH PPS, and also are currently used to determine payments for SSO cases under § 412.529 as well as payments for site neutral payment rate cases. (We note that the provisions of § 412.529 are only applicable to LTCH PPS standard Federal payment rate cases). However, if our proposed SSO payment method is finalized, CCRs would no longer be used to determine the payment adjustment for SSO cases. Therefore, if our proposed SSO policies are finalized, this discussion would no longer be relevant to all HCO and site neutral payment rate calculations.

As noted earlier, currently in determining HCO, SSO, and the site neutral payment rate (regardless of whether the case is also an HCO) payments, we generally calculate the estimated cost of the case by multiplying the LTCH’s overall CCR by the Medicare allowable charges for the case. An overall CCR is used because the LTCH PPS uses a single prospective payment per discharge that covers both inpatient operating and capital-related costs. The LTCH’s overall CCR is generally computed based on the sum of LTCH operating and capital costs (as described in Section 150.24, Chapter 3, of the Medicare Claims Processing Manual (Pub. 100–4)) as compared to total Medicare charges (that is, the sum of its operating and capital inpatient routine and ancillary charges), with those values determined from either the most recently settled cost report or the most recent tentatively settled cost report.

## Table 1: Proposed Cost-of-Living Adjustment Factors for Alaska and Hawaii Under the LTCH PPS for FY 2018

<table>
<thead>
<tr>
<th>Area</th>
<th>FY 2013</th>
<th>FY 2014 through FY 2017</th>
<th>Proposed FY 2018</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alaska:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>City of Anchorage and 80-kilometer (50-mile) radius by road</td>
<td>1.23</td>
<td>1.23</td>
<td>1.25</td>
</tr>
<tr>
<td>City of Fairbanks and 80-kilometer (50-mile) radius by road</td>
<td>1.23</td>
<td>1.23</td>
<td>1.25</td>
</tr>
<tr>
<td>City of Juneau and 80-kilometer (50-mile) radius by road</td>
<td>1.23</td>
<td>1.23</td>
<td>1.25</td>
</tr>
<tr>
<td>Rest of Alaska</td>
<td>1.25</td>
<td>1.25</td>
<td>1.25</td>
</tr>
<tr>
<td>Hawaii:</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>City and County of Honolulu</td>
<td>1.25</td>
<td>1.25</td>
<td>1.25</td>
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<tr>
<td>County of Hawaii</td>
<td>1.18</td>
<td>1.19</td>
<td>1.21</td>
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<tr>
<td>County of Kauai</td>
<td>1.25</td>
<td>1.25</td>
<td>1.25</td>
</tr>
<tr>
<td>County of Maui and County of Kalawao</td>
<td>1.25</td>
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<td>1.25</td>
</tr>
</tbody>
</table>
whichever is from the latest cost reporting period. However, in certain instances, we use an alternative CCR, such as the statewide average CCR, a CCR that is specified by CMS, or one that is requested by the hospital. (We refer readers to § 412.525(a)(4)(iv) of the regulations for further details regarding HCO adjustments for either LTCH PPS payment rate, § 412.529(f)(4) for SSO adjustments under the current policy, and § 412.522(c)(1)(ii) for the site neutral payment rate, respectively.)

The LTCH statewide average CCR is then compared to the LTCH total CCR ceiling. Under our established policy, an LTCH with a calculated CCR in excess of the applicable maximum CCR threshold (that is, the LTCH total CCR ceiling, which is calculated as 3 standard deviations from the national geometric average CCR) is generally assigned the applicable statewide CCR. This policy is premised on a belief that calculated CCRs above the LTCH total CCR ceiling are most likely due to faulty data reporting or entry, and that using an erroneous data should not be used to identify and make payments for outlier cases.

b. LTCH Total CCR Ceiling

Consistent with our historical practice, we are proposing to use the most recent data to determine the LTCH total CCR ceiling for FY 2018 in this proposed rule. Specifically, in this proposed rule, using our established methodology for determining the LTCH total CCR ceiling based on IPPS total CCR data from the December 2016 update of the Provider Specific File (PSF), which is the most recent data available, we are proposing to establish an LTCH total CCR ceiling of 1.276 under the LTCH PPS for FY 2018 in accordance with § 412.525(a)(4)(iv)(C)(2) for HCO cases under either payment rate and § 412.522(c)(1)(ii) for the site neutral payment rate. Also, consistent with our historical practice, we are proposing that if more recent data are available, we would use it to establish the LTCH total CCR ceiling for FY 2018 in the final rule. (For additional information on our methodology for determining the LTCH total CCR ceiling, we refer readers to the FY 2007 IPPS final rule (71 FR 48118 through 48119).)

c. LTCH Statewide Average CCRs

Our general methodology for determining the statewide average CCRs used under the LTCH PPS is similar to the established methodology for determining the LTCH total CCR ceiling because it is based on “total” IPPS CCR data. (For additional information on our methodology for determining statewide average CCRs under the LTCH PPS, we refer readers to the FY 2007 IPPS final rule (71 FR 48119 through 48120).) Under the LTCH PPS HCO policy for cases paid under either payment rate at § 412.525(a)(4)(iv)(C)(2), the current SSO policy at § 412.529(f)(4)(iii)(B), and the site neutral policy at § 412.522(c)(1)(ii), the MAC may use a statewide average CCR, which is established annually by CMS, if it is unable to determine an accurate CCR for an LTCH in one of the following circumstances: (1) New LTCHs that have not yet submitted their first Medicare cost report (a new LTCH is defined as an entity that has not accepted assignment of an existing hospital’s provider agreement in accordance with § 489.18); (2) LTCHs whose calculated CCR is in excess of the LTCH total CCR ceiling; and (3) other LTCHs for whom data with which to calculate a CCR are not available (for example, missing or faulty data). (Other sources of data that the MAC may consider in determining an LTCH’s CCR include data from a different cost reporting period for the LTCH, data from the cost reporting period preceding the period in which the LTCH is unable to determine an accurate CCR for rural hospitals that would be effective for discharges occurring on or after October 1, 2017, through September 30, 2018, in Table 8C in section VI of the Addendum to this proposed rule (and available via the Internet on the CMS Web site). Consistent with our historical practice, we also are proposing that if more recent data become available, we would use that data to determine the LTCH PPS statewide average total CCRs for FY 2018 in the final rule.

Under the current LTCH PPS labor market areas, all areas in Delaware, the District of Columbia, New Jersey, and Rhode Island are classified as rural. Therefore, there are no rural statewide average total CCRs listed for those jurisdictions in Table 8C. This policy is consistent with the policy that we established when we revised our methodology for determining the applicable LTCH statewide average CCRs in the FY 2007 IPPS final rule (71 FR 48119 through 48121) and is the same policy applied under the IPPS. In addition, although Connecticut has areas that are designated as rural, in our calculation of the LTCH statewide average CCRs, there was no data available from short-term, acute care IPPS hospitals or LTCHs located in that area as of December 2016. Therefore, consistent with our existing methodology, we are proposing to use the national average total CCR for rural IPPS hospitals for rural Connecticut in Table 8C. While Massachusetts also has rural areas, the statewide average CCR for rural areas in Massachusetts is based on one provider whose CCR is an atypical 1.222. Because this is much higher than the statewide urban average of 0.646 and furthermore implies costs outside a defined geographic area, we are proposing to use the national average total CCR for rural hospitals for hospitals located in rural Massachusetts. Furthermore, consistent with our existing methodology, in determining the urban and rural statewide average total CCRs for Maryland LTCHs paid under the LTCH PPS, we are proposing to continue to use, as a proxy, the national average total CCR for urban IPPS hospitals and the national average total CCR for rural IPPS hospitals, respectively. We are using this proxy because we believe that the CCR data in the PSF for Maryland hospitals may not be entirely accurate (as discussed in greater detail in the FY 2007 IPPS final rule (71 FR 48120)).

d. Reconciliation of HCO and SSO Payments

Under the HCO policy for cases paid under either payment rate at § 412.525(a)(4)(iv)(D) and the current SSO policy at § 412.529(f)(4)(iv), the payments for HCO and SSO cases are subject to reconciliation. Specifically, any such payments are reconciled at settlement based on the CCR that is calculated based on the cost report coinciding with the discharge. However, under our proposed changes to the SSO payment methodology discussed in section VIII.D. of the preamble of this proposed rule, we are proposing to replace cost as a consideration for payment to SSO cases. As such, consistent with our proposed changes to the SSO payment methodology, we are proposing that SSO payments would no longer be subject to reconciliation.

Specifically, we are proposing to revise paragraph (l) of § 412.529 to specify that SSO payments would be reconciled only for discharges occurring before October 1, 2017. We note that this proposal is dependent upon adoption of our proposed SSO payment methodology, and if those changes are not finalized, we would not finalize this proposal either.

For additional information on the reconciliation policy, we refer readers to Sections 150.26 through 150.28 of the Medicare Claims Processing Manual (Pub. 100–4), as added by Change Request 7192 (Transmittal 2111; December 3, 2010), and the FY 2009 LTCH PPS final rule (73 FR 26820 through 26821).

3. High-Cost Outlier Payments for LTCH PPS Standard Federal Payment Rate Cases

a. Proposed Changes to High-Cost Outlier Payments for LTCH PPS Standard Federal Payment Rate Cases

When we implemented the LTCH PPS, we established a fixed-loss amount so that total estimated outlier payments are projected to equal 8 percent of total estimated payments under the LTCH PPS (67 FR 56022 through 56026). Furthermore, § 412.523(d)(1) requires the LTCH PPS standard Federal payment rate be adjusted by a reduction factor of 8 percent, the estimated proportion of outlier payments under § 412.525(a) payable to LTCH PPS standard Federal payment rate cases. Section 15004(b) of the 21st Century Cures Act (Pub. L. 114–255) amended section 1886(m) of the Act by adding new paragraph (7), which specifies certain treatment of HCO payments for fiscal years beginning on or after October 1, 2017. Specifically, section 1886(m)(7)(A) of the Act requires, beginning in FY 2018, that the LTCH PPS standard Federal payment rate be reduced as if estimated HCO payments for standard Federal payment rate cases would be equal to 8 percent of estimated aggregate payments for standard Federal payment rate cases for
a given year. In other words, section 1866(m)(7)(A) of the Act makes our existing regulatory budget neutrality requirement at § 412.523(d)(1) for the 8 percent HCO target for standard Federal payment rate cases a statutory requirement beginning in FY 2018. In addition, section 1866(m)(7)(B) of the Act requires, beginning in FY 2018, that the fixed-loss amount for HCO payments for LTCH PPS standard Federal payment rate cases be determined so that the estimated aggregate amount of HCO payments for such cases (that is, 99.6875 percent of the 8 percent estimated aggregate payments for standard Federal payment rate cases (that is, 7.975 percent). In other words, sections 1866(m)(7)(A) and (7)(B) require that we adjust the standard Federal payment rate each year to ensure budget neutrality for HCO payments as if estimated aggregate HCO payments made for standard Federal payment rate discharges remain at 8 percent, while the fixed-loss amount for the HCO payments is set each year so that the estimated aggregate HCO payments for standard Federal payment rate cases are 7.975 percent of estimated aggregate payments for standard Federal payment rate cases.

More specifically, section 1866(m)(7)(A) of the Act stipulates that, for fiscal years beginning on or after October 1, 2017, the Secretary shall reduce the standard Federal payment rate as if the estimated aggregate amount of HCO payments for standard Federal payment rate discharges for each such fiscal year would be equal to 8 percent of estimated aggregate payments for standard Federal payment rate discharges for each such fiscal year; while section 1866(m)(7)(B) of the Act states that the Secretary shall set the fixed loss amount for HCO payments such that the estimated aggregate amount of HCO payments made for standard Federal payment rate discharges for fiscal years beginning on or after October 1, 2017, shall be equal to 99.6875 percent of 8 percent of estimated aggregate payments for standard Federal payment rate discharges for each such fiscal year; while section 1866(m)(7)(C) of the Act requires that any reduction in payments resulting from the application of paragraph (B) shall not be taken into account in applying any budget neutrality provision. Finally, section 1866(m)(7)(D) of the Act provides there will be no effect on HCO payments to site neutral payment rate cases by this certain treatment of HCO payments by requiring that this paragraph shall not apply with respect to the computation of the applicable site neutral payment rate under section 1866(m)(6) of the Act.

To codify the treatment of HCO payments provided by section 15004(b) of the 21st Century Cures Act (discussed earlier), we are proposing to revise § 412.523(a) by redesignating paragraph (2) as paragraph (2)(i) and adding paragraph (2)(ii) which would specify that, for FY 2018 and subsequent years, the fixed-loss amount for LTCH discharges described under § 412.522(a)(2) is determined such that the estimated proportion of outlier payments under § 412.522(a) that are payable for such discharges is projected to be equal to 99.6875 percent of 8 percent. We also are proposing to make conforming changes to § 412.523(d)(1) to specify that the provisions under proposed § 412.525(a)(2) would not affect the reduction factor of 8 percent that is applied to the LTCH PPS standard Federal payment rate cases.

h. Establishment of the Proposed Fixed-Loss Amount for LTCH PPS Standard Federal Payment Rate Cases for FY 2018

When we implemented the dual rate LTCH PPS payment structure beginning in FY 2016, we established that, in general, the historical LTCH PPS HCO policy will continue to apply to LTCH PPS standard Federal payment rate cases.

That is, the fixed-loss amount and target for LTCH PPS standard Federal payment rate cases is determined using the LTCH PPS HCO policy adopted when the LTCH PPS was first implemented, but we limited the data used under that policy to LTCH cases that would have been LTCH PPS standard Federal payment rate cases if the statutory changes had been in effect at the time of those discharges.

To determine the applicable fixed-loss amount for LTCH PPS standard Federal payment rate cases in FY 2018, we estimate outlier payments and total LTCH PPS payments for each LTCH PPS standard Federal payment rate case (or for each case that would have been a LTCH PPS standard Federal payment rate case if the statutory changes had been in effect at the time of the discharge) using claims data from the MedPAR files.

Historically, the applicable fixed-loss amount for LTCH PPS standard Federal payment rate cases results in estimated total outlier payments being projected to be equal to 8 percent of projected total LTCH PPS payments for LTCH PPS standard Federal payment rate cases. We use MedPAR claims data and CCRs based on data from the most recent PSF (or from the applicable statewide average CCR if an LTCH’s CCR data are faulty or unavailable). Under this proposal, we would increase from FY 2016 ($16,432) to our estimated total LTCH PPS payments in FY 2017, which exceeds the 8 percent target by 0.6 percentage points. We continue to believe, as discussed in detail in the FY 2017 IPPS/LTCH PPS proposed rule (81 FR 25287), this increase is largely attributable to rate-of-change (that is, increases in the Medicare allowable charges on the claims data in the MedPAR file. In particular, using the historic 8-percent target for projected aggregate outlier payments (absent the required changes under the 21st Century Cures Act for comparison purposes), the proposed HCO threshold would be $29,934, and thus represents a 36-percent increase from the final FY 2017 HCO threshold of $21,943. However, this increase is in line with previous proposed increases of the HCO threshold, such as the 36-percent increase from FY 2016 ($16,432) to our proposed FY 2017 HCO threshold ($22,728). We further note that the proposed FY 2017 HCO threshold was established based on the most recent data available at that time (specifically, the December 2015 update of the FY 2015 MedPAR file and the December 2015 update of the PSF), and in the FY 2017 final rule, based on the March 2016 update of the FY 2015 MedPAR file and the March 2016 update to the PSF, we finalized a somewhat lower HCO threshold of $21,943. Consistent with our historical practice of using the best data available, we are proposing that, when determining the fixed-loss amount for LTCH PPS standard Federal payment rate cases for FY 2018 in the final rule, we would use the most recent available LTCH claims data and CCR data at that time. The amount of the fixed-loss amount occurred in the first few years after the implementation of the LTCH PPS, due, in part, to the changes in LTCH behavior (such as Medicare beneficiary treatment patterns) in response to the new payment system and the lack of data and information available to predict how those changes would...
Furthermore, in accordance with 1886(m)(7)(B) of the Act as discussed above. We are proposing to continue to apply a budget neutrality factor to LTCH PPS standard Federal payment rate cases to offset our historic 8 percent HCO target for LTCH PPS standard Federal payment rate cases that would be necessary to achieve budget neutrality if the estimated aggregate HCO payments were set to be equal to 8 percent. As described in detail above, our calculation of the proposed fixed-loss amount for LTCH PPS standard Federal payment rate cases for FY 2018 of $30,081 is generally consistent with the methodology used to establish the FY 2017 LTCH PPS fixed-loss amount (absent the modification from an HCO target of 8 percent to the now statutorily required 7.975 percent HCO target).

c. Application of the High-Cost Outlier Policy to Short Stay Outlier (SSO) Cases

Under our implementation of the dual rate LTCH PPS payment structure required by statute, we are proposing that LTCH PPS standard Federal payment rate cases (that is, LTCH discharges that meet the criteria for exclusion from the site neutral payment rate) would be continued on the LTCH PPS standard Federal payment rate, and would include all of the existing payment adjustments under § 412.525(d), such as the adjustments for SSO cases under § 412.529. Under some rare circumstances, an LTCH discharge can qualify as an SSO case (as defined in the regulations at § 412.529 in conjunction with § 412.530) and also as an HCO case, as discussed in the August 30, 2002 final rule (67 FR 56026). In this scenario, a patient could be hospitalized for less than five-sixths of the geometric average length of stay for the specific MS–LTAC–DRG, and yet incur extraordinarily high treatment costs. If the estimated costs exceeded the HCO threshold (that is, the SSO payment plus the applicable fixed-loss amount), the discharge is eligible for payment at an HCO rate (We note that, under our proposed change to the SSO policy discussed in section VIII.D. of the preamble of this proposed rule, SSO cases would still be eligible to qualify for an HCO payment.) Therefore, for an SSO case in FY 2018, we are proposing to establish that the HCO payment would be 80 percent of the difference between the estimated cost of the case and the outlier threshold (the sum of the proposed fixed-loss amount of $30,081 and the amount paid under the proposed SSO policy as specified in § 412.529).

4. Proposed High-Cost Outlier Payments for Site Neutral Payment Rate Cases

Under § 412.525(a), site neutral payment rate cases will be paid under the site neutral payment rate for costs that exceed the HCO threshold that is equal to 80 percent of the difference between the estimated cost of the case and the applicable HCO threshold (80 FR 49618 through 49629). In the following discussion, we will refer to the statutory transitional payment method for cases that are paid the site neutral payment rate for LTCH discharges occurring in cost reporting periods beginning during FY 2016 or FY 2017 uses a blended payment rate, which is determined as 50 percent of the standard Federal prospective payment rate amount for the discharge ($412.522(c)(3)). The transitional blended payment rate uses the same blend percentages (that is, 50 percent) for both years of the 2-year transition period. For FY 2016, the site neutral payment rate cases discharged in FY 2016 and in a cost reporting period that ends before October 1, 2017 continue to be paid under the blended payment rate. However, site neutral payment rate cases discharged in FY 2018 during the LTCH’s cost reporting period beginning on or after October 1, 2017 will no longer be paid under the blended payment rate and instead will be paid the site neutral payment rate amount as determined under § 412.522(c)(1). As such, for FY 2018 discharges paid under the transitional payment method, the discussion that follows applies only to cases that are discharged in cost reporting periods that begin before October 1, 2017 and in a cost reporting period that ends after October 1, 2017 continue to be paid under the blended payment rate. Under § 412.522(c)(3)(i) of the blended payment rate (as well as to FY 2018 discharges paid the site neutral payment rate amount determined under § 412.522(c)(1)).

When we implemented the application of the site neutral payment rate in FY 2016, in examining the appropriate fixed-loss amount for site neutral payment rate cases, we considered how LTCH discharges based on historical claims data would have been classified under the dual rate LTCH PPS payment structure and the CMS’ Office of the Actuary projections regarding LTCHs for FY 2018. We again relied on these considerations and actuarial projections in FY 2017 because the historical claims data available in FY 2017 predated the LTCH PPS dual rate payment system. Similarly, for FY 2018, we continue to rely on these considerations and actuarial projections because, due to the rolling effective date of the site neutral payment policy, not all claims in FY 2016 were paid under the site neutral payment system.

For both FY 2016 and FY 2017, at that time our actuaries projected that the proportion of cases that would qualify as LTCH PPS standard Federal payment rate cases versus site neutral payment rate cases under the statutory provisions would remain consistent with what is reflected in the historical LTCH PPS claims data. Although our actuaries did not project an immediate change in the proportions found in the historical data, they did project cost and resource changes to account for the lower payment rates. Our actuaries also projected that the costs and resource use for cases paid at the site neutral payment rate would likely be lower, on average, than the costs and resource use for cases paid at the LTCH PPS standard Federal payment rate and would include all of the existing payment adjustments under § 412.525(d), such as the adjustments for SSO cases under § 412.529. Under some rare circumstances, an LTCH discharge can qualify as an SSO case (as defined in the regulations at § 412.529 in conjunction with § 412.530) and also as an HCO case, as discussed in the August 30, 2002 final rule (67 FR 56026). In this scenario, a patient could be hospitalized for less than five-sixths of the geometric average length of stay for the specific MS–LTAC–DRG, and yet incur extraordinarily high treatment costs. If the estimated costs exceeded the HCO threshold (that is, the SSO payment plus the applicable fixed-loss amount), the discharge is eligible for payment at an HCO rate (We note that, under our proposed change to the SSO policy discussed in section VIII.D. of the preamble of this proposed rule, SSO cases would still be eligible to qualify for an HCO payment.) Therefore, for an SSO case in FY 2018, we are proposing to establish that the HCO payment would be 80 percent of the difference between the estimated cost of the case and the outlier threshold (the sum of the proposed fixed-loss amount of $30,081 and the amount paid under the proposed SSO policy as specified in § 412.529).
For these reasons, we continue to believe that the most appropriate fixed-loss amount for site neutral payment rate cases would generally be paid based on an IPPS comparable per diem amount under the statutory LTCH PPS payment changes that began in FY 2016, which, in the majority of cases, would be lower than the payments that would have been paid if those statutory changes were not enacted. In light of these projections and expectations, we discussed that we believed that the use of a single fixed-loss amount and HCO target for all LTCH cases would be problematic. In addition, we discussed that we did not believe that it would be appropriate for comparable LTCH PPS site neutral payment rate cases to receive dramatically different HCO payments from those cases that would be paid under the IPPS (80 FR 49617 through 49619 and 81 FR 57305 through 57307). For those reasons, we stated that we believed that the most appropriate fixed-loss amount for site neutral payment rate cases for both FY 2016 and FY 2017 would be equal to the IPPS fixed-loss amount that year. Therefore, we established the fixed-loss amount for site neutral payment rate cases as the FY 2016 and FY 2017 IPPS fixed-loss amounts, in FY 2016 and FY 2017 respectively. In particular, in FY 2017, we established that the fixed-loss amount for site neutral payment rate cases is the FY 2017 IPPS fixed-loss amount of $23,570.

As noted earlier, because not all claims in the data used for this proposed rule were subject to the site neutral payment rate system, we continue to rely on the same consideration and actuarial projections used in FY 2016 and FY 2017 when developing a fixed-loss amount for site neutral payment rate cases for FY 2018. Because our actuaries continue to project that site neutral payment rate cases in FY 2018 will continue to mirror an IPPS case paid under the same MS–DRG, we continue to believe that it would be inappropriate for comparable LTCH PPS site neutral payment rate cases to receive dramatically different HCO payments from those cases that would be paid under the IPPS. As with FY 2016 and FY 2017, our actuaries project that the costs and resource use for FY 2018 cases paid at the site neutral payment rate would likely be lower, on average, than the costs and resource use for cases paid at the LTCH PPS standard Federal payment rate and will likely mirror the costs and resource use for IPPS cases assigned to the same MS–DRG, regardless of whether the proportion of site neutral payment rate cases in the future remains similar to what is found based on the historical data. (Based on the most recent FY 2016 LTCH claims data, approximately 58 percent of LTCH cases would have been paid the LTCH PPS standard Federal payment rate and approximately 42 percent of LTCH cases would have been paid the site neutral payment rate if those rates had been in effect at that time and discharges occurring in FY 2016, regardless of LTCHs’ cost reporting period beginning dates.)

For these reasons, we continue to believe that the most appropriate fixed-loss amount for site neutral payment rate cases for FY 2018 is the IPPS fixed-loss amount for FY 2018. Therefore, consistent with past practice, for FY 2018, we are proposing that the applicable HCO threshold for site neutral payment rate cases is the sum of the site neutral payment rate for the case and the IPPS fixed-loss amount. That is, we are proposing a fixed-loss amount for site neutral payment rate cases equal to 7.1 percent of the IPPS fixed-loss amount discussed in section II.A.4.g.(1) of the Addendum to this proposed rule. We continue to believe that this policy would result in HCO payments for similar cases under the LTCH PPS and promote fairness between the two systems. Accordingly, for FY 2018, we are proposing to calculate a HCO payment for site neutral payment rate cases with costs that exceed the HCO threshold amount, which is equal to 80 percent of the difference between the estimated cost of the case and the outlier threshold (the sum of proposed site neutral payment rate payment and the proposed fixed-loss amount for site neutral payment rate cases). In establishing a HCO policy for site neutral payment rate cases, we established a budget neutrality adjustment under §412.522(c)(2)(i). We established this requirement because we believed, and continue to believe, that the HCO policy for site neutral payment rate cases should be budget neutral, just as the HCO policy for LTCH PPS standard Federal payment rate cases are budget neutral, meaning that estimated site neutral payment rate HCO payments should not result in any change in estimated aggregate payments.

To ensure that estimated HCO payments payable to site neutral payment rate cases in FY 2018 would not result in any increase in estimated aggregate FY 2018 LTCH PPS payments, under the budget neutrality requirement at §412.522(c)(2)(i), it is necessary to reduce site neutral payment rate payments (or the portion of the blended payment rate payment for FY 2018 discharges occurring in LTCH cost reporting periods beginning before October 1, 2017) by 5.1 percent. As with FY 2016, we estimated additional HCO payments payable to those FY 2018 site neutral payment rate cases. In order to achieve this, for FY 2018, in general we are proposing to continue to use the policy adopted for FY 2017.

As discussed earlier, consistent with the IPPS HCO payment threshold, we estimate our proposed fixed-loss threshold of $26,713 results in HCO payments for site neutral payment rate cases equal to 5.1 percent of the site neutral payment rate payments that are based on the IPPS comparable per diem amount. As such, to ensure estimated HCO payments payable for site neutral payment rate cases in FY 2018 would not result in any increase in estimated aggregate FY 2018 LTCH PPS payments, under the budget neutrality requirement at proposed revised §412.522(c)(2)(i), it is necessary to reduce the site neutral payment rate amount paid under §412.522(c)(1)(i) by 5.1 percent to account for the estimated additional HCO payments payable for site neutral payment rate cases in FY 2018. In order to achieve this, for FY 2018, we are proposing to apply a proposed budget neutrality factor of 0.949 (that is, the decimal equivalent of a 5.1 percent reduction, determined as 1.0–5.1/100 = 0.949) to the site neutral payment rate for those site neutral payment rate cases paid under §412.522(c)(1)(i). We note that, consistent with the policy adopted for FY 2016, under this proposed fixed-loss amount and site neutral payment rate cases assigned to the same MS–DRG; that is, 5.1 percent. In other words, we estimated that HCO payments for site neutral payment rate cases would be 5.1 percent of the site neutral payment rate payments (80 FR 49805 and 81 FR 57307).

E. Proposed Update to the IPPS Comparable/ Equivalent Amounts To Reflect the Statutory Changes to the IPPS DSH Payment Adjustment Methodology

In the FY 2014 IPPS/LTCH PPS final rule (78 FR 50766), we established a policy to reflect the changes to the Medicare IPPS DSH payment adjustment methodology made by section 3133 of the Affordable Care Act in the calculation of the “IPPS comparable amount” under the SSO policy at §412.529 and the “IPPS equivalent amount” under the 25 percent threshold payment adjustment policy at §412.534 and §412.536. Historically, the determination of both the “IPPS comparable amount” and the “IPPS equivalent amount” includes an amount for inpatient operating costs “for the costs of serving a disproportionate share of low-income patients.” Under the statutory changes to the Medicare DSH payment adjustment methodology that began in FY 2014, in general, eligible IPPS hospitals receive an empirically justified Medicare DSH payment equal to 25 percent of the amount they otherwise would have received under the statutory formula for Medicare DSH payments prior to the amendments made by the Affordable Care Act. The remaining amount, equal to an estimate of 75 percent of the amount that otherwise would have been paid as Medicare DSH payments, reduced to reflect changes in the percentage of individuals who are uninsured, is made available to make additional payments to each hospital that qualifies for Medicare DSH payments and that has uncompensated care. The additional uncompensated care payments are based on the total amount of uncompensated care for a given time period relative to the total amount of uncompensated care for that same time period reported by all IPPS hospitals that receive Medicare DSH payments.

To reflect the statutory changes to the Medicare DSH payment adjustment...
methodology in the calculation of the “IPPS comparable amount” and the “IPPS equivalent amount” under the LTCH PPS, we stated that we will include a reduced Medicare DSH payment amount that reflects the projected percentage of the payment amount that would have been made prior to the amendments made by the Affordable Care Act that will be paid to eligible IPPS hospitals. As a result, for FY 2018, we propose that the reduction in the amount of Medicare DSH payments pursuant to section 1886(f)(1) of the Act, along with the payments for uncompensated care under section 1886(f)(2) of the Act, would result in overall Medicare DSH payments of 66.52 percent of the amount of Medicare DSH payments that would otherwise have been made in the absence of the amendments made by the Affordable Care Act (that is, 25 percent + 43.51 percent = 68.51 percent).

In this proposed rule, for FY 2018, we are proposing to establish that the calculation of the “IPPS comparable amount” under §412.525(b) (the “IPPS equivalent amount” under §412.538 would include an applicable operating Medicare DSH payment amount that is equal to 68.51 percent of the operating Medicare DSH payment amount that would have been paid based on the statutory Medicare DSH payment formula but for the amendments made by the Affordable Care Act. Furthermore, consistent with our historical practice, we are proposing that if more recent data become available, if appropriate, we would use that data to determine this factor in the final rule.

F. Computing the Proposed Adjusted LTCH PPS Federal Prospective Payments for FY 2018

Section 412.525 sets forth the adjustments to the LTCH PPS standard Federal payment rate. Under the dual rate LTCH PPS payment structure, only LTCH PPS cases that meet the statutory criteria to be excluded from the site neutral payment rate are paid based on the LTCH PPS standard Federal payment rate. Under §412.525(c), the proposed LTCH PPS standard Federal payment rate is adjusted to account for differences in area wages by multiplying the proposed labor-related share of the LTCH PPS standard Federal payment for a case by the applicable LTCH PPS wage index (the proposed FY 2018 values are shown in Tables 12A through 12B listed in section VI. of the Addendum of this proposed rule and available via the Internet on the CMS Web site). The proposed LTCH PPS standard Federal payment rate is then adjusted to account for the higher costs of LTCHs located in Alaska and Hawaii by the applicable COLA factors (the proposed FY 2018 factors are shown in the chart in section V.C. of this Addendum) in accordance with section 1886(m)(5) of the Act. The resulting amount is the proposed adjusted LTCH PPS standard Federal payment rate.

To calculate the LTCH’s total adjusted Federal prospective payment for this Medicare patient case in FY 2018, we compute the wage-adjusted proposed Federal prospective payment amount by multiplying the unadjusted proposed FY 2018 LTCH PPS standard Federal payment rate ($41,497.20 by the proposed labor-related share (66.3 percent) and the proposed wage index value (1.0563). This wage-adjusted amount was then added to the proposed nonlabor-related portion of the unadjusted proposed LTCH PPS standard Federal payment rate (33.7 percent; adjusted for cost of living, if applicable) to determine the adjusted proposed LTCH PPS standard Federal payment rate, which is then multiplied by the proposed MS–LTC–DRG relative weight (0.9158) to calculate the total adjusted proposed LTCH PPS standard Federal prospective payment for FY 2018 ($39,421.67). The table below illustrates the components of the calculations in this example.

<table>
<thead>
<tr>
<th>Proposed LTCH PPS Standard Federal Prospective Payment Rate</th>
<th>$41,497.20</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proposed Labor-Related Share</td>
<td>$27,512.64</td>
</tr>
<tr>
<td>Proposed Wage Index (CBSA 16974)</td>
<td>$13,984.56</td>
</tr>
<tr>
<td>Proposed Nonlabor-Related Portion of the LTCH PPS Standard Federal Payment Rate</td>
<td>$29,061.60</td>
</tr>
<tr>
<td>Proposed Adjusted LTCH PPS Standard Federal Payment Amount</td>
<td>$39,421.67</td>
</tr>
</tbody>
</table>

VI. Tables Referenced in This Proposed Rule and Available Only Through the Internet on the CMS Web Site

This section lists the tables referred to throughout the preamble of this proposed rule and in this Addendum. In the past, a majority of these tables were published in the Federal Register as part of the annual proposed and final rules. However, similar to FYs 2012 through 2017, for the FY 2018 rulemaking cycle, the IPPS and LTCH tables will not be published in the Federal Register in the annual IPPS/LTC PPS proposed and final rules and will be available only through the Internet. Specifically, all IPPS tables listed below, with the exception of IPPS Tables 1A, 1B, 1C, and 1D, and LTCH PPS Table 1E will be available only through the Internet. IPPS Tables 1A, 1B, 1C, and 1D, and LTCH PPS Table 1E are displayed at the end of this section and will continue to be published in the Federal Register as part of the annual proposed and final rules.
As discussed in the FY 2016 IPPS/LTCH PPS final rule (80 FR 49807), we streamlined and consolidated the wage index tables for FY 2016 and subsequent fiscal years.

As discussed in sections II.F.14., II.F.15.b., II.F.16., II.F.17.a., and II.F.19.a.1., a.3., and c.1. of the preamble of this proposed rule, we developed the following ICD–10–CM and ICD–10–PCS code tables for FY 2018: Table 6A—New Diagnosis Codes; Table 6B—New Procedure Codes; Table 6C—Invalid Diagnosis Codes; Table 6D—Invalid Procedure Codes; Table 6E—Revised Diagnosis Code Titles; Table 6F—Revised Procedure Code Titles; Table 6G.1—Proposed Secondary Diagnosis Order Additions to the CC Exclusion List; Table 6G.2—Proposed Principal Diagnosis Order Additions to the CC Exclusion List; Table 6H.1—Proposed Secondary Diagnosis Order Deletions to the CC Exclusion List; Table 6H.2—Proposed Principal Diagnosis Order Deletions to the CC Exclusion List; Table 6I.1—Proposed Additions to the MCC List; Table 6I.2—Proposed Deletions to the MCC List; Table 6J.1—Proposed Additions to the CC List; Table 6J.2—Proposed Deletions to the CC List; and Table 6P—Proposed ICD–10–CM and ICD–10–PCS Code Designations, MCE and MS–DRG Changes. In addition, under the Affordable Care Act, a hospital will receive under section 3133 of the Affordable Care Act.

The following IPPS tables for this FY 2018 proposed rule are available only through the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/index.html. Click on the link on the left side of the screen titled, “FY 2018 IPPS Proposed Rule Home Page” or “Acute Inpatient—Files for Download.”

Table 2—Proposed Case-Mix Index and Wage Index Table by CCN—FY 2016

Table 3—Proposed Wage Index Table by CBSA—FY 2018

Table 5—List of Proposed Medicare Severity Diagnosis-Related Groups (MS–DRGs), Relative Weighting Factors, and Geometric and Arithmetic Mean Length of Stay—FY 2018

Table 6A—New Diagnosis Codes—FY 2018

Table 6B—New Procedure Codes—FY 2018

Table 6C—Invalid Diagnosis Codes—FY 2018

Table 6D—Invalid Procedure Codes—FY 2018

Table 6E—Revised Diagnosis Code Titles—FY 2018

Table 6F—Revised Procedure Code Titles—FY 2018

Table 6G.1—Proposed Secondary Diagnosis Order Additions to the CC Exclusions List—FY 2018

Table 6G.2—Proposed Principal Diagnosis Order Additions to the CC Exclusions List—FY 2018

Table 6H.1—Proposed Secondary Diagnosis Order Deletions to the CC Exclusions List—FY 2018

Table 6H.2—Proposed Principal Diagnosis Order Deletions to the CC Exclusions List—FY 2018

Table 6I.1—Proposed Additions to the MCC List—FY 2018

Table 6I.2—Proposed Deletions to the MCC List—FY 2018

Table 6J.1—Proposed Additions to the CC List—FY 2018

Table 6J.2—Proposed Deletions to the CC List—FY 2018

Table 6P—Proposed ICD–10–CM and ICD–10–PCS Code Designations, MCE and MS–DRG Changes—FY 2018

Table 4—Proposed Wage Index Table by CBSA—FY 2018

Table 5—List of Proposed Medicare Severity Diagnosis-Related Groups (MS–DRGs), Relative Weighting Factors, and Geometric and Arithmetic Mean Length of Stay—FY 2018

Table 6A—New Diagnosis Codes—FY 2018

Table 6B—New Procedure Codes—FY 2018

Table 6C—Invalid Diagnosis Codes—FY 2018

Table 6D—Invalid Procedure Codes—FY 2018

Table 6E—Revised Diagnosis Code Titles—FY 2018

Table 6F—Revised Procedure Code Titles—FY 2018

Table 6G.1—Proposed Secondary Diagnosis Order Additions to the CC Exclusions List—FY 2018

Table 6G.2—Proposed Principal Diagnosis Order Additions to the CC Exclusions List—FY 2018

Table 6H.1—Proposed Secondary Diagnosis Order Deletions to the CC Exclusions List—FY 2018

Table 6H.2—Proposed Principal Diagnosis Order Deletions to the CC Exclusions List—FY 2018

Table 6I.1—Proposed Additions to the MCC List—FY 2018

Table 6I.2—Proposed Deletions to the MCC List—FY 2018

Table 6J.1—Proposed Additions to the CC List—FY 2018

Table 6J.2—Proposed Deletions to the CC List—FY 2018

Table 6P—Proposed ICD–10–CM and ICD–10–PCS Code Designations, MCE and MS–DRG Changes—FY 2018

Table 7A—Medicare Prospective Payment System Selected Percentile Lengths of Stay: FY 2016 MedPAR Update—December 2016

Table 7B—Medicare Prospective Payment System Selected Percentile Lengths of Stay: FY 2016 MedPAR Update—December 2016

Table 8A—Proposed FY 2018 Statewide Average Operating Cost-to-Charge Ratios (CCRs) for Acute Care Hospitals (Urban and Rural)

Table 8B—Proposed FY 2018 Statewide Average Capital Cost-to-Charge Ratios (CCRs) for Acute Care Hospitals

Table 10—Proposed New Technology Add-On Payment Thresholds for Applications for FY 2019

Table 16—Proposed Proxy FY 2018 Readmissions Adjustment Factors

Table 16—Proposed Proxy Hospital Value-Based Purchasing (VBP) Program Adjustment Factors for FY 2018

Uncompensated Care Payment Factor 3

The following LTPS tables for this FY 2018 proposed rule are available only through the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/LongTermCareHospitalPPS/index.html under the list item for Regulation Number CMS–1677–P.

Table 8C—Proposed FY 2018 Statewide Average Total Cost-to-Charge Ratios (CCRs) for LTCHs (Urban and Rural)

Table 11—Proposed MS–LTC–DRGs, Relative Weights, Geometric Average Length of Stay, and Short-Stay Outlier (SSO) Threshold for LTPS Discharges Occurring from October 1, 2017 through September 30, 2018

Table 12A—Proposed LTPS Wage Index for Urban Areas for Discharges Occurring from October 1, 2017 through September 30, 2018

Table 12B—Proposed LTPS Wage Index for Rural Areas for Discharges Occurring from October 1, 2017 through September 30, 2018

Table 13A—Proposed Composition of Low Volume Quintiles for MS–LTC–DRGs—FY 2018

Table 13B—Proposed No Volume MS LTC–DRG Crosswalk for FY 2018

**TABLE 1A—PROPOSED NATIONAL ADJUSTED OPERATING STANDARDIZED AMOUNTS, LABOR/NONLABOR**

<table>
<thead>
<tr>
<th>Hospital submitted quality data and is a meaningful EHR user (update = 1.75 percent)</th>
<th>Hospital submitted quality data and is NOT a meaningful EHR user (update = -0.425 percent)</th>
<th>Hospital did NOT submit quality data and is a meaningful EHR user (update = 1.025 percent)</th>
<th>Hospital did NOT submit quality data and is NOT a meaningful EHR user (update = -1.15 percent)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Labor</td>
<td>Nonlabor</td>
<td>Labor</td>
<td>Nonlabor</td>
</tr>
<tr>
<td>$3,822.07</td>
<td>$1,773.93</td>
<td>$3,740.37</td>
<td>$1,736.01</td>
</tr>
</tbody>
</table>
TABLE 1B—PROPOSED NATIONAL ADJUSTED OPERATING STANDARDIZED AMOUNTS, LABOR/NONLABOR

<table>
<thead>
<tr>
<th>Hospital submitted quality data and is a meaningful EHR user (update = 1.75 percent)</th>
<th>Hospital submitted quality data and is NOT a meaningful EHR user (update = -0.425 percent)</th>
<th>Hospital did not submit quality data and is a meaningful EHR user (update = 1.025 percent)</th>
<th>Hospital did NOT submit quality data and is NOT a meaningful EHR user (update = -1.15 percent)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Labor</td>
<td>Nonlabor</td>
<td>Labor</td>
<td>Nonlabor</td>
</tr>
<tr>
<td>$3,469.52</td>
<td>$2,126.48</td>
<td>$3,395.36</td>
<td>$2,981.02</td>
</tr>
</tbody>
</table>

TABLE 1C—PROPOSED ADJUSTED OPERATING STANDARDIZED AMOUNTS FOR HOSPITALS IN PUERTO RICO, LABOR/NONLABOR

<table>
<thead>
<tr>
<th>Standardized amount</th>
<th>Rates if wage index is greater than 1</th>
<th>Rates if wage index is less than or equal to 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>National</td>
<td>Labor</td>
<td>Nonlabor</td>
</tr>
</tbody>
</table>

1 For FY 2018, there are no CBSAs in Puerto Rico with a national wage index greater than 1.

TABLE 1D—PROPOSED CAPITAL STANDARD FEDERAL PAYMENT RATE

<table>
<thead>
<tr>
<th>Rate</th>
<th>[FY 2018]</th>
</tr>
</thead>
<tbody>
<tr>
<td>National</td>
<td>$451.37</td>
</tr>
</tbody>
</table>

TABLE 1E—PROPOSED LTCH PPS STANDARD FEDERAL PAYMENT RATE

<table>
<thead>
<tr>
<th>Rate</th>
<th>[FY 2018]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Full update (1 percent)</td>
<td>Reduced update* (-1.0 percent)</td>
</tr>
<tr>
<td>Standard Federal</td>
<td>$41,497.20</td>
</tr>
</tbody>
</table>

* For LTCHs that fail to submit quality reporting data for FY 2018 in accordance with the LTCH Quality Reporting Program (LTCH QRP), the annual update is reduced by 2.0 percentage points as required by section 1866(m)(5) of the Act.

Appendix A: Economic Analyses

I. Regulatory Impact Analysis

A. Introduction

We have examined the impacts of this proposed rule as required by Executive Order 12866 on Regulatory Planning and Review (September 30, 1993), Executive Order 13563 on Improving Regulation and Regulatory Review (January 18, 2011), the Regulatory Flexibility Act (RFA) (September 19, 1980, Pub. L. 96–354), section 1102(b) of the Social Security Act, section 202 of the Unfunded Mandates Reform Act of 1995 (March 22, 1995; Pub. L. 104–4), Executive Order 13132 on Federalism (August 4, 1999), the Congressional Review Act (5 U.S.C. 804(2)), and Executive Order 13771 on Reducing Regulation and Controlling Regulatory Costs (January 30, 2017). Executive Orders 12866 and 13563 direct agencies to assess all costs and benefits of available regulatory alternatives and, if regulation is necessary, to select regulatory approaches that maximize net benefits (including potential social benefits) of the rule, and minimize any significant and unjustified economic, environmental, public health and safety effects, distributive impacts, and equity. Section 3(f) of Executive Order 12866 defines a “significant regulatory action” as an action that is likely to result in a rule: (1) (Having an annual effect on the economy of $100 million or more in any 1 year, or adversely and materially affecting a sector of the economy, productivity, competition, jobs, the environment, public health or safety, or state, local or tribal governments or communities (also referred to as “economically significant”); (2) creating a serious inconsistency or otherwise interfering with an action taken or planned by another agency; (3) materially altering the budgetary impact of entitlement programs, user fees, or loan programs or the rights and obligations of recipients thereof; or (4) raising novel legal or policy issues arising out of legal mandates, the President’s priorities, or the principles set forth in the Executive Order.

We have determined that this proposed rule is a major rule as defined in 5 U.S.C. 804(2). We estimate that the proposed changes for FY 2018 acute care hospital operating and capital payments would redistribute amounts in excess of $100 million to acute care hospitals. The applicable percentage increase to the IPPS rates required by the statute, in conjunction with other proposed payment changes in this proposed rule, would result in an estimated $3.1 billion increase in FY 2018 proposed operating payments, including a $3.8 billion increase in FY 2018 proposed operating payments (or 1.7 percent change), an estimated $212 million increase in FY 2018 proposed capital payments (or 2.4 percent change), and an estimated $1.6 billion increase in proposed uncompensated care payments (or 1.2 percent change). As noted in section II.A. of this Appendix, all expenditures are classified as transfers to Medicare providers. These proposed changes are relative to payments made in FY 2017. The impact analysis of the proposed capital payments can be found in section II.I. of this Appendix. In addition, as described in section II.J. of this Appendix, LTCHs are expected to experience a decrease in payments by $173 million in FY 2018 relative to FY 2017.

Our operating payment impact estimate includes the proposed 1.75 percent hospital update to the standardized amount (which includes the estimated 2.9 percent market basket update less 0.4 percentage point for the proposed multifactor productivity adjustment and less 0.75 percentage point required under the Affordable Care Act). Our operating payment impact estimate also includes an adjustment factor of (1/1.006) to the FY 2018 rates to remove the 1.006 temporary one-time adjustment made in FY 2017 to address the effects of the 0.2 percent reduction in effect for FYs 2014 through 2016 as a result of the 2-midnight policy (we refer readers to section V.M. of the preamble of this proposed rule for an explanation of this adjustment). The estimates of IPPS operating payments to acute care hospitals do not reflect any changes in hospital admissions or real cased-mix intensity, which will also affect overall proposed payment changes.

The analysis in this Appendix, in conjunction with the remainder of this document, demonstrates that this proposed rule is consistent with the regulatory philosophy and principles identified in Executive Orders 12866 and 13563, the RFA, and section 1102(b) of the Act. This proposed rule would affect payments to a substantial number of small rural hospitals, as well as other classes of hospitals, and the effects on some hospitals may be significant. Finally, in accordance with the provisions of Executive Order 12866, the Executive Office of Management and Budget has reviewed this proposed rule.
B. Statement of Need

This proposed rule is necessary in order to make payment and policy changes under the Medicare IPPS for Medicare acute care hospital inpatient services for operating and capital-related costs as well as for certain hospitals and hospital units excluded from the IPPS. This proposed rule also is necessary to make payment and policy changes for Medicare hospitals under the LTCH PPS.

C. Objectives of the IPPS and the LTCH PPS

The primary objective of the IPPS and the LTCH PPS is to create incentives for hospitals to operate efficiently and minimize unnecessary costs while at the same time ensuring that payments are sufficient to adequately compensate hospitals for their legitimate costs in delivering necessary care to Medicare beneficiaries. In addition, we share national goals of preserving the Medicare Hospital Insurance Trust Fund.

We believe that the changes in this proposed rule would further each of these goals while maintaining the financial viability of the hospital industry and ensuring access to high quality health care for Medicare beneficiaries. We expect that these proposed changes will ensure that the outcomes of the prospective payment systems are reasonable and equitable while avoiding or minimizing unintended adverse consequences.

Because this proposed rule contains a range of proposed policies, we refer readers to the section of the proposed rule where each proposal is discussed. These sections include the rational for our decisions, including the need for the proposed policy.

D. Limitations of Our Analysis

The following quantitative analysis presents the projected effects of our proposed policy changes, as well as statutory changes effective for FY 2018, on various hospital groups. We estimate the effects of individual proposed policy changes by estimating payments per case while holding all other payment policies constant. We use the best data available, but generally, we do not attempt to make adjustments for future changes in such variables as admissions, lengths of stay, or case-mix. In addition, we discuss limitations of our analysis for specific proposals in the discussion of those proposals as needed.

E. Hospitals Included in and Excluded From the IPPS

The prospective payment systems for hospital inpatient operating and capital-related costs of acute care hospitals encompass most general short-term, acute care hospitals that participate in the Medicare program. There were 31 Indian Health Service hospitals in our database, which we excluded from the analysis due to the specifics of the prospective payment methodology for these hospitals. Among other short-term, acute care hospitals, hospitals in Maryland are paid in accordance with the Maryland All-Payer Model, and hospitals located outside the 50 States, the District of Columbia, and Puerto Rico (that is, 5 short-term acute care hospitals located in the U.S. Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa) receive payment for inpatient hospital services they furnish on the basis of reasonable costs, subject to a rate-of-increase ceiling.

As of March 2017, there were 3,292 IPPS acute care hospitals included in our analysis. This represents approximately 94 percent of all Medicare-participating hospitals. The majority of this impact analysis focuses on this set of hospitals. There also are approximately 1,385 CAHs. These small, limited service hospitals are paid on the basis of reasonable costs rather than under the IPPS. IPPS-excluded hospitals and units, which are paid under separate payment systems, include IPFs, IRFs, LTCHs, RNHCIs, children’s hospitals, 11 cancer hospitals, and 5 short-term acute care hospitals located in the Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa. With the exception of the IPFQR provisions presented in section IX.D. of the preamble of this proposed rule, changes in the prospective payment systems for IPFs and IRFs are made through separate rulemaking. Payment impacts of changes to the prospective payment systems for these IPPS-excluded hospitals and units are not included in this proposed rule. The impact of the proposed update and proposed policy changes to the LTCH PPS for FY 2018 is discussed in section I.J. of this Appendix.

F. Effects on Hospitals and Hospital Units Excluded From the IPPS

As of March 2017, there were 98 children’s hospitals, 11 cancer hospitals, 5 short-term acute care hospitals located in the Virgin Islands, Guam, the Northern Mariana Islands and American Samoa, and 18 RNHCIs being paid on a reasonable cost basis subject to the rate-of-increase limit since their base period, the lesser of: (1) 50 percent of its reasonable costs in excess of 110 percent of the limit; or (2) 10 percent of its limit. In addition, under § 413.40 of the regulations, the update would be the percentage increase in the proposed IPPS operating market basket to update the target amounts for FY 2018 and subsequent years for children’s hospitals, the 11 cancer hospitals, the 5 short-term acute care hospitals located in the U.S. Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa, and RNHCIs that are paid based on reasonable costs subject to the rate-of-increase limits. Consistent with current law, based on IHS Global Insight, Inc.’s 2016 fourth quarter forecast of the proposed 2014-based IPPS operating market basket increase, we are estimating the FY 2018 update to be 2.9 percent (that is, the estimate of the market basket rate-of-increase). We are proposing that if more recent data become available for the final rule, we would use them to calculate the IPPS operating market basket update for FY 2018. However, the Affordable Care Act requires an adjustment for multifactor productivity (currently estimated to be 0.4 percentage point for FY 2018 and a 0.75 percentage point reduction to the market basket update, resulting in a 1.75 percent applicable percentage increase for IPPS hospitals that submit quality data and are meaningful EHR users, as discussed in section IV.B. of the preamble of this proposed rule. Children’s hospitals, the 11 cancer hospitals, the 5 short-term acute care hospitals located in the Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa, and RNHCIs that continue to be paid based on reasonable costs subject to rate-of-increase limits under § 413.40 of the regulations are not subject to the reductions in the applicable percentage increase required under the Affordable Care Act. Therefore, for those hospitals paid under § 413.40 of the regulations, the update would be the percentage increase in the proposed 2014-based IPPS operating market basket for FY 2018, estimated at 2.9 percent, without the reductions described previously under the Affordable Care Act.

The impact of the proposed update in the rate-of-increase limit on those excluded hospitals depends on the cumulative cost increases experienced by each excluded hospital since its applicable base period. For excluded hospitals that have maintained their cost increases at a level below the rate-of-increase limits since their base period, the major effect is on the level of inpatient payments these excluded hospitals receive. Conversely, for excluded hospitals with cost increases above the cumulative update in their rate-of-increase limits, the major effect is the amount of excess costs that would not be paid.

We note that, under § 413.40(d)(3), an excluded hospital that continues to be paid under the TEFRA system and whose costs exceed 110 percent of its rate-of-increase limit receives its rate-of-increase limit plus the lesser of: (1) 50 percent of its reasonable costs in excess of 110 percent of the limit; or (2) 10 percent of its limit. In addition, under the various provisions set forth in § 413.40, hospitals can obtain payment adjustments for justifiable increases in operating costs that exceed the limit.
G. Quantitative Effects of the Proposed Policy Changes Under the IPPS for Operating Costs

1. Basis and Methodology of Estimates

In this proposed rule, we are announcing proposed payment rates and proposed payment rate updates for the IPPS for FY 2018 for operating costs of acute care hospitals. The proposed FY 2018 updates to the capital payments to acute care hospitals are discussed in section I.I. of this Appendix. Based on the overall percentage change in payments per case estimated using our payment simulation model, we estimate that total FY 2018 operating payments would increase by 1.7 percent compared to FY 2017. In addition to the applicable percentage increase, this amount reflects the FY 2018 adjustment required under section 15005 of the 21st Century Cures Act described in section II.D. of the preamble of this proposed rule of 0.4588 percent to the IPPS national standardized amounts. This amount also reflects the adjustment factor of (1/1.006) to remove the temporary one-time adjustment made in FY 2017 to address the effects of the 0.2 percent reduction in effect for FYs 2014 through 2016 related to the 2-midnight policy, which is discussed in section V.M. of the preamble of this proposed rule. The impacts do not reflect changes in the number of hospital admissions or real case-mix intensity, which would also affect overall proposed payment changes.

We have prepared separate impact analyses of the proposed changes to each system. This section deals with the proposed changes to the operating inpatient prospective payment system for acute care hospitals. Our payment simulation model relies on the most recent available data to enable us to estimate the impacts on payments per case of certain changes in this proposed rule. However, there are other proposed changes for which we do not have data available that would allow us to estimate the payment impacts using this model. For those proposed changes, we have attempted to predict the payment impacts based upon our experience and other more limited data.

The data used in developing the quantitative analyses of proposed changes in payments per case presented in this section are taken from the FY 2016 MedPAR file and the most current Provider-Specific File (PSF) that is used for payment purposes. Although the analyses of the proposed changes to the operating PPS do not incorporate cost data, data from the most recently available hospital cost reports were used to categorize hospitals. Our analysis has several qualifications. First, in this analysis, we do not make adjustments for future changes in such variables as admissions, lengths of stay, or underlying growth in real case-mix. Second, due to the interdependent nature of the IPPS payment components, it is very difficult to precisely quantify the impact associated with each proposed change. Third, we use various data sources to categorize hospitals in the tables. In some cases, particularly the number of beds, there is a fair degree of variation in the data from the different sources. We have attempted to construct these variables with the best available source overall. However, for individual hospitals, some miscategorizations are possible.

Using cases from the FY 2016 MedPAR file, we simulate payments under the operating IPPS given various combinations of payment parameters. As described previously, instead of operating inpatient hospital and hospital in Maryland were excluded from the simulations. The proposed impact of payments under the capital IPPS, or the impact of payments for costs other than inpatient operating costs, are not analyzed in this section. We estimated payment impacts of the capital IPPS for FY 2018 are discussed in section I.I. of this Appendix.

We discuss the following proposed changes:

• The effects of the proposed application of the adjustment required under section 15005 of the 21st Century Cures Act and the applicable percentage increase (including the proposed market basket update, the proposed multifactor productivity adjustment, and the applicable percentage increase in accordance with the Affordable Care Act) to the standardized amount and hospital-specific rates.
• The effects of the adjustment of (1/1.006) to remove the temporary one-time adjustment made in FY 2017 to address the effects of the 0.2 percent reduction in effect for FYs 2014 through 2016 related to the 2-midnight policy, as discussed in section V.M. of the preamble of this proposed rule.
• The effects of the proposed changes to the relative weights and MS–DRG GROUPER.
• The effects of the proposed changes in hospitals’ wage index values reflecting updated wage data from hospitals’ cost reporting periods beginning during FY 2014, compared to the FY 2013 wage data, to calculate the FY 2018 wage index.
• The effects of the geographic reclassifications by the MGCRB (as of publication of this proposed rule) that would be effective for FY 2018.
• The effects of the proposed rural floor with the application of the proposed national budget neutrality factor to the wage index.
• The effects of the proposed frontier State wage index adjustment under the statutory provision that requires that hospitals located in States that qualify as frontier States to not have a wage index less than 1.0. This provision is not budget neutral.
• The effects of the implementation of section 1886(d)(13) of the Act, as added by section 5001(a) of Public Law 109–171, as amended by section 4102(b)(1)(A) of the ARRA (Pub. L. 111–5) and by section 3401(a)(2) of the Affordable Care Act (Pub. L. 111–149), provides that, for FY 2007 and each subsequent year through FY 2014, the update factor will include a reduction of 2.0 percentage points for any subsection (d) hospital that does not submit data on measures in a form and manner and at a time specified by the Secretary. Beginning in FY 2015, the reduction is one-quarter of such applicable percentage increase determined without regard to section 1886(b)(3)(B)(vi) of the Act.
• The effects of the expiration of the special payment status for MDHs at the end of FY 2017 under current law as a result of which MDHs that currently receive the higher of payments made based on the Federal rate plus 75 percent of the difference between payments based on the Federal rate and the hospital-specific rate will be paid based on the Federal rate starting in FY 2018.
• The total estimated change in payments based on the proposed FY 2018 policies relative to payments based on FY 2017 policies that include the applicable percentage increase of 1.75 percent (or 2.9 percent market basket update with a proposed reduction of 0.4 percentage point for the multifactor productivity adjustment, and a 0.75 percentage point adjustment, as required under the Affordable Care Act).

To illustrate the impact of the proposed FY 2018 changes, our analysis begins with a FY 2017 baseline simulation model using: The FY 2017 applicable percentage increase of 1.75 percent and the FY 2016 reduction in reimbursement of 1.5 percent to the Federal standardized amount; the adjustment of (1/0.998) to permanently remove the –0.2 percent reduction to the rate put in place in FY 2014 to offset the estimated increase in IPPS expenditures as a result of the 2-midnight policy; the 1.006 temporary adjustment to address the effects of the 0.2 percent reduction in effect for FYs 2014 through 2016 related to the 2-midnight policy; the FY 2017 MS–DRG GROUPER (Version 34); the FY 2017 wage index; and no MGCRB reclassifications. Outlier payments are set at 5.1 percent of total operating MS–DRG and outlier payments for modeling purposes.

Section 1886(b)(3)(B)(vi) of the Act, as added by section 5001(a) of Public Law 109–171, as amended by section 4102(b)(1)(A) of the ARRA (Pub. L. 111–5) and by section 3401(a)(2) of the Affordable Care Act (Pub. L. 111–149), provides that, for FY 2007 and each subsequent year through FY 2014, the wage index for hospitals based on the OMB definitions from the 2010 Census; the FY 2017 wage index; and no MGCRB reclassifications. For purposes of the simulations shown later in this section, we modeled the proposed payment changes for FY 2018 using a reduced update for these hospitals.

For FY 2018, in accordance with section 1886(b)(3)(B)(ii) of the Act, a hospital that has been identified as not a meaningful EHR user would be subject to a reduction of three-quarters of such applicable percentage increase determined without regard to section 1886(b)(3)(B)(ix), (x), (xi), or (xii) of the Act, or one-quarter of the market basket update. Therefore, for FY 2018, we are proposing that hospitals that do not submit quality information data, meaningful EHR users under section 1886(b)(3)(B)(ix) of the Act would receive an applicable percentage increase of 1.025 percent. At the time that this impact was prepared, 62 hospitals are estimated to not receive the full market basket rate-of-increase for FY 2018 because they failed the quality data submission process or did not choose to participate but are meaningful EHR users. For purposes of the simulations shown later in this section, we modeled the proposed payment changes for FY 2018 using a reduced update for these hospitals.

For FY 2018, in accordance with section 1886(b)(3)(B)(ix) of the Act, a hospital that has been identified as not a meaningful EHR user would be subject to a reduction of three-quarters of such applicable percentage increase determined without regard to section 1886(b)(3)(B)(ix), (x), (xi), or (xii) of the Act. Therefore, for FY 2018, we are proposing that hospitals that are identified as not meaningful EHR users and do submit quality information under section 1886(b)(3)(B)(viii) of the Act would receive an applicable
percentage increase of –0.425 percent. At the time that this impact analysis was prepared, 103 hospitals are estimated to not receive the full market basket rate-of-increase for FY 2018 because they are identified as not meaningful EHR users that do submit quality information under section 1886(b)(3)(B)(ix) of the Act. For purposes of the simulations shown in this section, we modeled the proposed payment changes for FY 2018 using a reduced update for these hospitals. Hospitals that are identified as not meaningful EHR users under section 1886(b)(3)(B)(ix) of the Act and also do not submit quality data under section 1886(b)(3)(B)(viii) of the Act would receive an applicable percentage increase of –1.15 percent, which reflects a one-quarter reduction of the market basket update for failure to submit quality data and a three-quarter reduction of the market basket update for FY 2018 because they are identified as not meaningful EHR user.

Each proposed policy change, statutory or otherwise, is then added incrementally to this baseline, finally arriving at an FY 2018 model incorporating all of the proposed changes. This simulation allows us to isolate the effects of each proposed change.

Our final comparison illustrates the percent change in payments per case from FY 2017 to FY 2018. Two factors not discussed separately have significant impacts here. The first factor is the proposed update to the standardized amount. In accordance with section 1886(b)(3)(B)(i) of the Act, we are proposing to update the standardized amounts for FY 2018 using a proposed applicable percentage increase of 1.75 percent. This includes our forecasted IPPS operating hospital market basket increase of 2.9 percent with a 0.4 percentage point reduction for the multifactor productivity adjustment and a 0.75 percentage point reduction as required under the Affordable Care Act. Hospitals that fail to comply with the quality data submission requirements and are meaningful EHR users would receive a proposed update of 1.025 percent. This update includes a reduction of one-quarter of the market basket update for failure to submit these data. Hospitals that do comply with the quality data submission requirements but are not meaningful EHR users would receive an update of –0.425 percent, which includes a reduction of three-quarters of the market basket update. Furthermore, hospitals that do not comply with the quality data submission requirements and also are not meaningful EHR users would receive an update of –1.15 percent. Under section 1886(b)(3)(B)(iv) of the Act, the update to the hospital-specific amounts for SCHs is also equal to the applicable percentage increase, or 1.75 percent if the hospital submits quality data and is a meaningful EHR user.

A second significant factor that affects the proposed changes in hospitals’ payments per case from FY 2017 to FY 2018 is the change in hospitals’ geographic reclassification status from one year to the next. That is, payments may be reduced for hospitals reclassified in FY 2017 that are no longer reclassified in FY 2018. Conversely, payments may increase for hospitals not reclassified in FY 2017 that are reclassified in FY 2018.

2. Analysis of Table I

Table I displays the results of our analysis of the proposed changes for FY 2018. The table categorizes hospitals by various geographic and special payment consideration groups to illustrate the varying impacts on different types of hospitals. The top row of the table shows the overall impact on the 3,292 acute care hospitals included in the analysis.

The next four rows of Table I contain hospitals categorized according to their geographic location: All urban, which is further divided into large urban and other urban; and rural. There are 2,491 hospitals located in urban areas included in our analysis. Among these, there are 1,349 hospitals located in large urban areas (populations over 1 million), and 1,142 hospitals in other urban areas (populations of 1 million or fewer). In addition, there are 801 hospitals in rural areas. The next two groupings are by bed-size categories, shown separately for urban and rural hospitals. The final groupings by geographic location are by census divisions, also shown separately for urban and rural hospitals.

The second part of Table I shows hospital groups based on hospitals’ FY 2018 proposed payment classifications, including any reclassifications under section 1866(d)(10) of the Act. For example, the rows labeled urban, large urban, other urban, and rural show that the numbers of hospitals paid based on these categorizations after consideration of geographic reclassifications (including reclassifications under sections 1866(d)(8)(B) and 1866(d)(6)(E) of the Act that have implications for capital payments) are 2,391, 1,363, 1,028, and 901, respectively.

The next three groupings examine the impacts of the proposed changes on hospitals grouped by whether or not they have GME residency programs (teaching hospitals that receive an IME adjustment) or receive Medicare DSH payments, or some combination of these two payments. There are 2,211 nonteaching hospitals in our analysis, 835 teaching hospitals with fewer than 100 residents, and 246 teaching hospitals with 100 or more residents.

In the DSH categories, hospitals are grouped according to their DSH payment status, and whether they are considered urban or rural for DSH purposes. The next category groups together hospitals considered urban or rural, in terms of whether they receive the IME adjustment, the DSH adjustment, both, or neither.

The next three rows examine the impacts of the proposed changes on rural hospitals by special payment groups (SCHs, and RRCs).

The next series of groupings is based on the type of ownership and the hospital’s Medicare utilization expressed as a percent of total patient days. These data were taken from the FY 2014 or FY 2013 Medicare cost reports.

The next two groupings concern the geographic reclassification status of hospitals. The first grouping displays all urban hospitals that were reclassified by the MGCRB for FY 2018. The second grouping shows the MGCRB rural reclassifications.

### Table I—Impact Analysis of Proposed Changes to the IPPS for Operating Costs for FY 2018

<table>
<thead>
<tr>
<th>Number of hospitals</th>
<th>Proposed hospital rate update and adjustments</th>
<th>Proposed FY 2018 weights and DRG changes with application of recalibration budget neutrality</th>
<th>Proposed FY 2018 wage data with application of wage neutrality</th>
<th>FY 2018 SCH/MGCRB reclassifications</th>
<th>Proposed rural floor with application of national rural floor budget neutrality</th>
<th>Proposed application of the frontier wage index and out-migration adjustment</th>
<th>Expiration of MDH status</th>
<th>All proposed FY 2018 changes</th>
</tr>
</thead>
<tbody>
<tr>
<td>All Hospitals</td>
<td>3,292</td>
<td>1.5</td>
<td>0</td>
<td>0</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>1.7</td>
</tr>
<tr>
<td>By Geographic Location:</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Urban hospitals</td>
<td>2,491</td>
<td>1.6</td>
<td>0</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>1.8</td>
</tr>
<tr>
<td>Large urban areas</td>
<td>1,349</td>
<td>1.6</td>
<td>–0.1</td>
<td>0</td>
<td>0.4</td>
<td>-0.1</td>
<td>0.1</td>
<td>1.7</td>
</tr>
<tr>
<td>Other urban areas</td>
<td>1,142</td>
<td>1.6</td>
<td>0</td>
<td>0.3</td>
<td>0.2</td>
<td>2</td>
<td>0.2</td>
<td>1.8</td>
</tr>
<tr>
<td>Rural hospitals</td>
<td>801</td>
<td>1.3</td>
<td>0.3</td>
<td>0</td>
<td>0.2</td>
<td>0.2</td>
<td>0.2</td>
<td>0.8</td>
</tr>
<tr>
<td>Bed Size (Urban):</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0–99 beds</td>
<td>638</td>
<td>1.5</td>
<td>0.4</td>
<td>0.1</td>
<td>–0.6</td>
<td>0.1</td>
<td>0.2</td>
<td>0.7</td>
</tr>
<tr>
<td>100–199 beds</td>
<td>765</td>
<td>1.6</td>
<td>0.2</td>
<td>0.1</td>
<td>–0.1</td>
<td>0.2</td>
<td>0.2</td>
<td>0.1</td>
</tr>
<tr>
<td>200–299 beds</td>
<td>445</td>
<td>1.6</td>
<td>0.1</td>
<td>0</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>1.7</td>
</tr>
</tbody>
</table>
20198

Federal Register / Vol. 82, No. 81 / Friday, April 28, 2017 / Proposed Rules

TABLE I—IMPACT ANALYSIS OF PROPOSED CHANGES TO THE IPPS FOR OPERATING COSTS FOR FY 2018—Continued

asabaliauskas on DSK3SPTVN1PROD with PROPOSALS

Number of
hospitals 1

300–499 beds ...................
500 or more beds .............
Bed Size (Rural):
0–49 beds .........................
50–99 beds .......................
100–149 beds ...................
150–199 beds ...................
200 or more beds .............
Urban by Region:
New England .....................
Middle Atlantic ...................
South Atlantic ....................
East North Central ............
East South Central ............
West North Central ...........
West South Central ...........
Mountain ...........................
Pacific ................................
Puerto Rico .......................
Rural by Region:
New England .....................
Middle Atlantic ...................
South Atlantic ....................
East North Central ............
East South Central ............
West North Central ...........
West South Central ...........
Mountain ...........................
Pacific ................................
By Payment Classification:
Urban hospitals .................
Large urban areas ............
Other urban areas .............
Rural areas .......................
Teaching Status:
Nonteaching ......................
Fewer than 100 residents
100 or more residents .......
Urban DSH:
Non-DSH ...........................
100 or more beds .............
Less than 100 beds ..........
Rural DSH:
SCH ...................................
RRC ..................................
100 or more beds .............
Less than 100 beds ..........
Urban teaching and DSH:
Both teaching and DSH ....
Teaching and no DSH ......
No teaching and DSH .......
No teaching and no DSH ..
Special Hospital Types:
RRC ..................................
SCH ...................................
SCH and RRC ..................
Type of Ownership:
Voluntary ...........................
Proprietary .........................
Government ......................
Medicare Utilization as a Percent of Inpatient Days:
0–25 ..................................
25–50 ................................
50–65 ................................
Over 65 .............................
FY 2018 Reclassifications by
the Medicare Geographic
Classification Review Board:
All Reclassified Hospitals ..
Non-Reclassified Hospitals

VerDate Sep<11>2014

19:54 Apr 27, 2017

Proposed
hospital rate
update and
adjustments

Proposed
FY 2018
weights and
DRG
changes
with
application
of
recalibration
budget
neutrality

Proposed
FY 2018
wage data
with
application
of wage
budget
neutrality

(1) 2

(2) 3

(3) 4

FY 2018
MGCRB
reclassifications

Proposed
rural floor
with
application
of national
rural floor
budget
neutrality

Proposed
application
of the
frontier wage
index and
out-migration
adjustment

Expiration of
MDH status

All proposed
FY 2018
changes

(4) 5

(5) 6

(6) 7

(7) 8

(8) 9

431
212

1.6
1.5

0
¥0.3

0
0

¥0.1
¥0.2

0
¥0.1

0.1
0.1

0
0

1.8
1.7

313
285
117
46
40

1.2
1.3
1.3
1
1.4

0.5
0.3
0.3
0.2
0.1

0
0
0
0.1
0.2

0.4
0.6
1.3
1.8
2.9

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0
¥0.2
¥0.2

0.3
0.2
0.2
0.1
0

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0

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1.7
2

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404
385
147
160
378
162
375
51

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1.6
1.5
1.6
1.5
1.5
1.6

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¥0.1
¥0.4

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¥0.3
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0.4
¥0.4
¥0.2
¥0.3
¥0.8
¥0.5
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¥0.2
¥1

1
¥0.3
¥0.3
¥0.3
¥0.2
¥0.3
¥0.3
0.3
0.9
0.2

0
0.1
0
0
0
0.7
0
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0.1

¥0.2
0
¥0.1
0
0
¥0.1
¥0.1
0
0
0

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1.2
1.9
2
1.7
2
2.1
1
2
1.3

20
53
125
115
154
97
154
59
24

1.3
1.2
1.2
1.3
1.5
1.2
1.3
1
1.1

0.2
0.4
0.3
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0.1
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0.2

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0
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0
0.3
¥0.1
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1.2
2.4
0.1
1.7
0.2
1

¥0.3
¥0.2
¥0.2
¥0.2
¥0.3
0.2
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¥0.1
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0.2
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¥1.7
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¥0.3
¥0.7
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1.2
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1.3
1.4
1.2

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1,028
901

1.6
1.6
1.6
1.4

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1.4

0
¥0.1
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1.8
1.7
1.8
1.2

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835
246

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1.6
1.5

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0
0
¥0.1

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0.2
0

¥0.3
0
0

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1.6

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1.6
1.5

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0
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0.1

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¥0.1

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1.8
2.1

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271
41
240

1.1
1.4
1.6
1.5

0.2
0.1
0.2
0.7

0
0.2
0.3
0

0
1.7
1.6
0.6

0
0
¥0.1
¥0.3

0
0.3
0.1
0.7

0
¥0.3
0
¥4.7

1.2
1.9
1.7
¥3.2

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1.6
1.6
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¥0.1
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0

¥0.1
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¥0.4
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¥0.4

¥0.1
¥0.2
0.3
¥0.1

0.1
0.1
0.1
0.2

0
0
0
0

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1.2
2
1.9

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1.1

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0.1

2.1
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0.3

¥0.1
0.2
0

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0

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0

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1.3

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514

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1.6
1.5

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0

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0
0
0.2

0.1
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0.1

¥0.1
¥0.1
¥0.1

1.6
1.9
1.6

509
2,113
535
135

1.5
1.6
1.5
1.5

0
0
0.1
0.6

0
0
0.1
0.1

¥0.4
0
0.6
¥0.5

0.2
0
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0.4

0
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0.1
0.3

0
¥0.1
¥0.5
¥3.7

1.6
1.8
1.2
¥1.5

900
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1.5
1.6

0.1
0

0.1
0

1.9
¥0.9

¥0.1
0

0
0.2

¥0.1
¥0.1

1.7
1.7

Jkt 241001

PO 00000

Frm 00404

Fmt 4701

Sfmt 4702

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28APP2


a. Effects of the Proposed Hospital Update, Adjustment Required Under Section 15005 of the 21st Century Cures Act, and Other Adjustments (Column 1)

As discussed in section V.B. of the preamble of this proposed rule, this column includes the proposed hospital update, including the proposed 2.9 percent market basket update, the proposed reduction of 0.4 percentage point for the multifactor productivity adjustment, and the 0.75 percentage point reduction in accordance with the Affordable Care Act. In addition, as discussed in section I.D. of the preamble of this proposed rule, this column includes the FY 2018 adjustment of 0.4588 percent on the national standardized amount required under section 15005 of the 21st Century Cures Act and, as discussed in section V.M. of the preamble of this proposed rule, the adjustment factor of (1/1.006) to remove the 1.066 temporary one-time adjustment made in FY 2017 to address the effects of the 0.2 percent reduction in effect for FYs 2014 through 2016 related to the 2-midnight policy. As a result, we are proposing to make a 1.6 percent update to the national standardized amount. This column also includes the proposed update to the hospital-specific rates which includes the proposed 2.9 percent market basket update, the proposed reduction of 0.4 percentage point for the multifactor productivity adjustment, and the 0.75 percentage point reduction in accordance with the Affordable Care Act. As discussed in section V.M. of the preamble of this proposed rule, the adjustment factor of (1/1.006) to remove the 1.066 temporary one-time adjustment made in FY 2017 to address the effects of the 0.2 percent reduction in effect for FYs 2014 through 2016 related to the 2-midnight policy to both the national standardized amount and the hospital-specific rate. Hospitals that are paid under the hospital-specific rate would experience a 1.15 percent increase in payments; therefore, hospital categories containing hospitals paid under the hospital-specific rate would experience a lower than average increases in payments.

b. Effects of the Proposed Changes to the MS–DRG Reclassifications and Relative Cost-Based Weights With Recalibration Budget Neutrality (Column 2)

Column 2 shows the effects of the proposed changes to the MS–DRGs and relative weights with the application of the recalibration budget neutrality factor to the standardized amounts. Section 1886(d)(4)(C)(ii) of the Act requires us annually to make appropriate classification changes in order to reflect changes in treatment patterns, technology, and any other factors that may change the relative use of hospital resources. Consistent with section 1886(d)(4)(C)(iii) of the Act, we are calculating a recalibration budget neutrality factor to account for the changes in MS–
DRGs and relative weights to ensure that the overall payment impact is budget neutral.

As discussed in section II.E. of the preamble of this proposed rule, the FY 2018 MS–DRG relative weights would be 100 percent cost-based and 100 percent MS–DRGs. For FY 2018, the MS–DRGs are calculated using the FY 2016 MedPAR data grouped to the Version 35 (FY 2018) MS–DRGs. The methodology to calculate the relative weights and the reclassification changes to the GROUPER are described in more detail in section II.G. of the preamble of this proposed rule.

The “All Hospitals” line in Column 2 indicates that proposed changes due to the MS–DRGs and relative weights would result in a 0.0 percent change in payments with the application of the proposed recalibration budget neutrality factor of 0.997573 to the standardized amount. Hospital categories that generally treat more surgical cases than medical cases would experience a decrease in their payments under the proposed relative weights for reasons that include the proposals regarding operating room procedures described in section II.G. of the preamble of this proposed rule. Rural hospitals would experience a 0.3 percent increase in payments in part because rural hospitals tend to treat fewer surgical cases than medical cases, while teaching hospitals with more than 100 residents would experience a decrease in payments by 0.3 percent in part because those hospitals treat more surgical cases than medical cases.

c. Effects of the Proposed Wage Index Changes (Column 3)

Column 3 shows the impact of updated wage data using FY 2014 cost report data, with the application of the wage budget neutrality factor. The wage index is calculated and assigned to hospitals on the basis of the labor market area in which the hospital is located. Under section 1886(d)(3)(E) of the Act, beginning with FY 2005, we delineate hospital labor market areas based on the Core Based Statistical Areas (CBSAs) established by OMB. The current statistical standards used in FY 2018 are based on OMB standards published on February 28, 2013 (75 FR 37246 and 37252), and 2010 Decennial Census data (OMB Bulletin No. 13–01), as updated in OMB Bulletin No. 15–01. (We refer readers to the FY 2015 IPPS/LTC PPS final rule (79 FR 49951 through 49963) for a full discussion on our adoption of the OMB labor market area delineations based on the 2010 Decennial Census data, effective beginning with the FY 2015 IPPS wage index, and to section III.A.2. of the preamble of the FY 2017 IPPS/LTC PPS final rule (81 FR 56913) for a discussion of our adoption of the CBSA updates in OMB Bulletin No. 15–01, which were effective beginning with the FY 2017 wage index.)

Section 1886(d)(3)(E) of the Act requires that, beginning October 1, 1993, we annually update the wage data used to calculate the wage index. In accordance with this requirement, the proposed wage index for acute care hospitals for FY 2018 is based on data submitted for hospital cost reporting periods beginning on or after October 1, 2013 and before October 1, 2014. The estimated impact of the updated wage data using the FY 2014 cost report data and the OMB labor market area delineations on hospital payments is isolated in Column 3 by holding the other parameter constants in this simulation. That is, Column 3 shows the percentage change in payments when going from a model using the FY 2017 wage index, based on FY 2013 wage data, the labor-related share of 69.6 percent, under the OMB delineations and having a 100-percent occupational mix adjustment applied, to a model using the FY 2018 pre-reclassification wage index based on FY 2014 wage data with the labor-related share of 68.3 percent, under the OMB delineations, also having a 100-percent occupational mix adjustment applied, while holding other proposed payment parameter changes such as use of the Version 35 MS–DRG GROUPER constant.

The proposed FY 2018 occupational mix adjustment is based on the CY 2013 occupational mix survey.

In addition, the column shows the impact of the proposed application of the wage budget neutrality to the national standardized amount. In FY 2010, we began calculating separate wage budget neutrality and recalibration budget neutrality factors, in accordance with section 1886(d)(3)(E) of the Act, which specifies that budget neutrality to account for wage index changes or updates made under that subparagraph must be made without regard to the 62 percent labor-related share guaranteed under section 1886(d)(3)(E)(ii) of the Act. Therefore, for FY 2018, we begin calculating the wage budget neutrality factor to ensure that payments under updated wage data and the labor-related share of 68.3 percent are budget neutral without regard to the lower labor-related share of 62 percent applied to hospitals with a wage index less than or equal to 1.0. In other words, the wage budget neutrality is calculated under the assumption that all hospitals receive the higher labor-related share of the standardized amount. The proposed FY 2018 wage budget neutrality factor is 1.000465, and the overall proposed payment change is 0.0 percent.

Column 3 shows the impacts of updating the wage data using FY 2014 cost reports. Overall, the new wage data and the labor-related share, combined with the proposed wage budget neutrality adjustment, would lead to no change for all hospitals as shown in Column 3.

In looking at the wage data itself, the proposed national average hourly wage would increase 1.02 percent compared to FY 2017. Therefore, the only manner in which to maintain or exceed the previous year’s wage index was to match or exceed the 1.02 percent increase in the national average hourly wage. Of the 3,287 hospitals with wage data for both FYs 2017 and 2018, 1,698 or 51.7 percent would experience an average hourly wage increase of 1.02 percent or more.

The following chart compares the shifts in wage index values for hospitals due to proposed changes in the average hourly wage data for FY 2018 relative to FY 2017. Among urban hospitals, 10 would experience a decrease of 10 percent or more, and 2 urban hospitals would experience an increase of 10 percent or more. One hundred and one urban hospitals would experience an increase or decrease of at least 5 percent or more but less than 10 percent. Among rural hospitals, none would experience an increase of at least 5 percent or more, but 12 rural hospitals would experience a decrease of greater than or equal to 5 percent but less than 10 percent. Three rural hospitals would experience decreases of 10 percent or more. Seven rural hospitals would experience increases or decreases of less than 5 percent, while 2,384 urban hospitals would experience increases or decreases of less than 5 percent. No urban hospitals and no rural hospitals experience no change to their wage index. These figures reflect proposed changes in the “pre-reclassified, occupational mix-adjusted wage index,” that is, the proposed wage index before the application of proposed geographic reclassification, the proposed rural floor, the proposed out-migration adjustment, and other proposed wage index exceptions and adjustments. (We refer readers to sections III.C. through III.L. of the preamble of this proposed rule for a complete discussion of the exceptions and adjustments to the wage index.) We note that the proposed “post-reclassified wage index” or proposed “payment wage index,” which is the proposed wage index that includes all such exceptions and adjustments (as reflected in Tables 2 and 3 associated with this proposed rule, which are available via the Internet on the CMS Web site) is used to adjust the labor-related share of a hospital’s standardized amount, either 68.3 percent or 62 percent, depending upon whether a hospital’s wage index is greater than 1.0 or less than or equal to 1.0. Therefore, the proposed pre-reclassified wage index and the proposed wage index values in the following chart may illustrate a somewhat larger or smaller change than would occur in a hospital’s proposed payment wage index and total payment.

The following chart shows the projected impact of proposed changes in the area wage index values for urban and rural hospitals.

<table>
<thead>
<tr>
<th>Increase 10 percent or more</th>
<th>Increase greater than or equal to 5 percent and less than 10 percent</th>
<th>Increase or decrease less than 5 percent</th>
<th>Decrease greater than or equal to 5 percent and less than 10 percent</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>54</td>
<td>2,384</td>
<td>47</td>
</tr>
<tr>
<td>0</td>
<td>775</td>
<td>775</td>
<td>12</td>
</tr>
</tbody>
</table>
d. Effects of MGCRB Reclassifications (Column 4)

Our impact analysis to this point has assumed acute care hospitals are paid on the basis of their actual geographic location (with the exception of ongoing policies that provide that certain hospitals receive payments on bases other than where they are geographically located). The proposed changes in Column 4 reflect the per case payment impact of moving from this baseline to a simulation incorporating the MGCRB decisions for FY 2018.

By spring of each year, the MGCRB makes reclassification determinations that would be effective for the next fiscal year, which begins on October 1. The MGCRB may approve a hospital’s reclassification request for the purpose of using another area’s wage index value. Hospitals may appeal denials of MGCRB decisions to the CMS Administrator. Further, hospitals have 45 days from publication of the IPPS proposed rule in the Federal Register to decide whether to approve a hospital’s reclassification request or lose due to the application of the proposed rural floor. We are proposing to extend the imputed floor policy.

Table 2 listed in section VI. of the Addendum to this proposed rule and available via the Internet on the CMS Web site reflects the proposed reclassifications for FY 2018.

e. Effects of the Proposed Rural Floor, Including Application of National Budget Neutrality (Column 5)

As discussed in section III.B. of the preamble of the FY 2009 IPPS final rule, the FY 2010 IPPS/RY 2010 LTCH PPS final rule, the FYs 2011, 2012, 2013, 2014, 2015, 2016, and 2017 IPPS/LTCH PPS final rules, and this proposed rule, section 4410 of Public Law 105–33 established the rural floor by requiring that the wage index for a hospital in any urban area cannot be less than the wage index received by rural hospitals in the same State. We would apply a uniform budget neutrality adjustment to the wage index. As discussed in section III.H. of the preamble of this proposed rule, we are not proposing to extend the imputed floor policy. Therefore, column 6 shows the effects of the proposed rural floor only.

The Affordable Care Act requires that we apply one rural floor budget neutrality factor to the wage index nationally. We have calculated a proposed FY 2018 rural floor budget neutrality factor to be applied to the wage index of 0.993672, which would reduce wage indexes by 0.63 percent. Column 5 shows the projected impact of the proposed rural floor with national budget neutrality. The column compares the proposed post-reclassification FY 2018 wage index of providers before the proposed rural floor adjustment and the proposed post-reclassification FY 2018 wage index of providers with the proposed rural floor adjustment based on the OMB labor market area delineations. Only urban hospitals can benefit from the rural floor. Because the provision is budget neutral, all other hospitals (that is, all rural hospitals and those urban hospitals to which the adjustment is not made) would experience a decrease in payments due to the budget neutrality adjustment that is applied nationally to their wage index.

We estimate that 392 hospitals would receive the proposed rural floor in FY 2018. All IPPS hospitals in our model would have their wage index reduced by the rural floor budget neutrality adjustment of 0.993672. We project that, in aggregate, rural hospitals would experience a 0.63 percent decrease in payments as a result of the application of the proposed rural floor budget neutrality because the rural hospitals do not benefit from the rural floor, but have their wage indexes downwardly adjusted to ensure that the application of the rural floor is budget neutral overall. We project hospitals located in urban areas would experience no change in payments because increases in payments by hospitals benefiting from the rural floor offset decreases in payments by non-rural floor urban hospitals whose wage index is downwardly adjusted by the rural floor budget neutrality factor. Urban hospitals in the New England region would experience a 1.0 percent increase in payments primarily due to the application of the proposed rural floor in Massachusetts. Thirty-six urban providers in Massachusetts are expected to receive the proposed rural floor wage index value, including the rural floor budget neutrality adjustment, increasing payments overall to Massachusetts by an estimated $44 million. We estimate that Massachusetts hospitals would receive approximately a 1.3 percent increase in IPPS payments due to the application of the proposed rural floor in FY 2018.

Urban Puerto Rico hospitals are expected to experience a 0.2 percent increase in payments as a result of the application of the proposed rural floor.

In response to a public comment addressed in the FY 2012 IPPS/LTCH PPS final rule (76 FR 51593), we are providing the payment impact of the proposed rural floor with budget neutrality at the State level. Column 1 of the following table displays the number of IPPS hospitals located in each State. Column 2 displays the number of hospitals in each State that would receive the proposed rural floor wage index for FY 2018. Column 3 displays the percentage of total payments each State would receive or contribute to fund the rural floor with national budget neutrality. The column compares the proposed post-reclassification FY 2018 wage index of providers before the proposed rural floor adjustment and the proposed post-reclassification FY 2018 wage index of providers with the proposed rural floor adjustment. Column 4 displays the estimated payment amount that each State would gain or lose due to the application of the proposed rural floor with national budget neutrality.

<table>
<thead>
<tr>
<th>Number of hospitals</th>
</tr>
</thead>
<tbody>
<tr>
<td>Urban</td>
</tr>
<tr>
<td>10</td>
</tr>
<tr>
<td>0</td>
</tr>
</tbody>
</table>
### Proposed FY 2018 IPPS Estimated Payments Due to Rural Floor with National Budget Neutrality

<table>
<thead>
<tr>
<th>State</th>
<th>Number of hospitals</th>
<th>Proposed number of hospitals that would receive the rural floor</th>
<th>Proposed percent change in payments due to application of rural floor with budget neutrality</th>
<th>Proposed difference (in $ millions)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alabama</td>
<td>84</td>
<td>3</td>
<td>0.2</td>
<td>3.05</td>
</tr>
<tr>
<td>Alaska</td>
<td>6</td>
<td>4</td>
<td>1.4</td>
<td>26.2</td>
</tr>
<tr>
<td>Arizona</td>
<td>57</td>
<td>44</td>
<td>0.9</td>
<td>17.47</td>
</tr>
<tr>
<td>Arkansas</td>
<td>44</td>
<td>1</td>
<td>-0.3</td>
<td>-3.39</td>
</tr>
<tr>
<td>California</td>
<td>299</td>
<td>177</td>
<td>1.3</td>
<td>136.28</td>
</tr>
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<td>Colorado</td>
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<td>4</td>
<td>0.4</td>
<td>4.97</td>
</tr>
<tr>
<td>Connecticut</td>
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<td>10</td>
<td>0.4</td>
<td>6.31</td>
</tr>
<tr>
<td>Delaware</td>
<td>6</td>
<td>0</td>
<td>-0.3</td>
<td>-1.61</td>
</tr>
<tr>
<td>Washington, D.C.</td>
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<td>0</td>
<td>-0.3</td>
<td>-1.67</td>
</tr>
<tr>
<td>Florida</td>
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<td>17</td>
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<td>-14.93</td>
</tr>
<tr>
<td>Georgia</td>
<td>103</td>
<td>0</td>
<td>-0.3</td>
<td>-8.07</td>
</tr>
<tr>
<td>Hawaii</td>
<td>12</td>
<td>0</td>
<td>-0.3</td>
<td>-0.83</td>
</tr>
<tr>
<td>Idaho</td>
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<td>0</td>
<td>-0.2</td>
<td>-0.77</td>
</tr>
<tr>
<td>Illinois</td>
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<td>-15.87</td>
</tr>
<tr>
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<td>-5.92</td>
</tr>
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<td>Iowa</td>
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<td>-0.3</td>
<td>-2.94</td>
</tr>
<tr>
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<td>-2.62</td>
</tr>
<tr>
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</table>

The effects of the Application of the Proposed Frontier State Wage Index and Out-Migration Adjustment (Column 6)

This column shows the combined effects of the application of section 1886(d)(13) of the Affordable Care Act, which requires that we establish a minimum post-reclassified wage-index of 1.00 for all hospitals located in “frontier States,” and the effects of section 505 of Public Law 108–173, which provides for an increase in the wage index for hospitals located in certain counties that have a relatively high percentage of hospital employees who reside in the county, but work in a different area with a higher wage index. These two wage index provisions are not budget neutral and increase payments overall by 0.1 percent compared to the provisions not being in effect.

The term “frontier States” is defined in the statute as States in which at least 50 percent of counties have a population density less than 6 persons per square mile. Based on these criteria, 5 States (Montana, Nevada, North Dakota, South Dakota, and Wyoming)
are considered frontier States and 48 hospitals located in those States would receive a frontier wage index of 1.0000. Overall, this provision is not budget neutral and is estimated to increase IPPS operating payments by approximately $65 million. Rural hospitals located in the West North Central region would experience an increase in payments by 0.3 and 0.7 percent, respectively, because many of the hospitals located in this region are frontier State hospitals.

In addition, section 1886(d)(13) of the Act, as added by section 505 of Public Law 108–173, provides for an increase in the wage index for hospitals located in certain counties that have a relatively high percentage of hospital employees who reside in the county, but work in a different area with a higher wage index. Hospitals located in counties that qualify for the payment adjustment are to receive an increase in the wage index that is equal to a weighted average of the difference between the wage index of the resident county, post-reclassification and the higher wage index work area(s), weighted by the overall percentage of workers who are employed in an area with a higher wage index. There are an estimated 429 providers that could receive the out-migration wage adjustment in FY 2018. Rural hospitals generally qualify for the adjustment, resulting in a 0.2 percent increase in payments. This provision appears to benefit section 401 hospitals and RRCs in that they would experience a 0.3 percent and 0.4 percent increase in payments, respectively. This out-migration wage adjustment also is not budget neutral, and we estimate the impact of these providers receiving the out-migration increase would be approximately $39 million.

g. Effects of the Expiration of MDH Special Payment Status (Column 7)

Column 7 shows our estimate of the changes in payments due to the expiration of MDH status, a nonbudget neutral payment provision. Section 205 of the Medicare Access and CHIP Reauthorization Act of 2015 (MACRA) (Pub. L. 114–10, enacted on April 16, 2015) extended the MDH program (which, under previous law, was to be in effect for discharges on or before March 31, 2015 only) for discharges occurring on or after April 1, 2015, through FY 2017 (that is, for discharges occurring on or before September 30, 2017). Therefore, under current law, the MDH program will expire at the end of FY 2017. Hospitals that qualified to be MDHs receive the higher of payments made based on the Federal rate or the payments made based on the Federal rate amount plus 75 percent of the difference between payments based the Federal rate and payments based the hospital-specific rate (a hospital-specific cost-based rate). Because this provision was not budget neutral, the expiration of this payment provision results in a 0.1 percent decrease in payments overall. There are currently 158 MDHs, of which we estimate 96 would have been paid under the blended payment of the Federal rate and hospital-specific rate if the MDH program had not expired. Because those 96 MDHs will no longer receive the blended payment and will be paid only under the Federal rate in FY 2018, it is estimated that those hospitals would experience an overall decrease in payments of approximately $119 million.

MDHs were generally rural hospitals, so the expiration of the MDH program will result in an overall decrease in payments to rural hospitals of 0.9 percent. Rural New England hospitals can expect a decrease in payments of 2.1 percent because 6 out of the 20 rural New England hospitals are MDHs that will lose this special payment status under the expiration of the program at the end of FY 2017. MDHs can expect a decrease in payments of 0.4 percent.

h. Effects of All FY 2018 Proposed Changes (Column 8)

Column 8 shows our estimate of the proposed changes in payments per discharge from FY 2017 to FY 2018, resulting from all proposed changes reflected in this proposed rule for FY 2018. It includes combined effects of the year to year change of the previous columns in the table. The proposed average increase in payments under the IPPS for all hospitals is approximately 1.7 percent for FY 2018 relative to FY 2017 and for this row is primarily driven by the changes reflected in Column 8. Column 8 includes the proposed annual hospital update of 1.6 percent to the national standardized amount. This proposed annual hospital update includes the 2.9 percent market basket update, the proposed reduction of 0.4 percentage point for the multifactor productivity adjustment, and the 0.75 percentage point reduction under section 3401 of the Affordable Care Act. As discussed in section II.D. of the preamble of this proposed rule, this column also includes the proposed FY 2018 adjustment of 0.4588 percent on the national standardized amount. In addition, this column includes the adjustment factor of (1/1.006) to remove the 1.006 temporary one-time adjustment made in FY 2017 to address the effects of the 0.2 percent reduction in effect for FYs 2014 through 2016 related to the 2-midnight policy, which is discussed in section V.M. of the preamble of this proposed rule. Hospitals paid under the hospital-specific rate would receive a 1.15 percent proposed hospital update. As described in Column 1, the proposed annual hospital update with the proposed adjustment of 0.4588 percent for hospitals paid under the national standardized amount, the adjustment of (1/1.006) to remove the 1.006 temporary one-time adjustment made in FY 2017 to address the effects of the 0.2 percent reduction in effect for FYs 2014 through 2016, which is discussed in section V.M. of the preamble of this proposed rule, combined with the proposed annual hospital update for hospitals paid under the hospital-specific rates would result in a 1.7 percent increase in payments in FY 2018 relative to FY 2017. There are also interactive effects among the various factors comprising the payment system that we are not able to isolate which contribute to our estimate of the proposed changes in payments per discharge from FY 2017 and FY 2018 in Column 8.

Overall payments to hospitals paid under the IPPS due to the proposed applicable percentage increase and proposed changes to policies related to MS–DRGs, geographic adjustments, and outliers are estimated to increase by 1.7 percent for FY 2018. Hospitals in urban areas would experience a 1.8 percent increase in payments per discharge in FY 2018 compared to FY 2017. Hospital payments per discharge in rural areas are estimated to increase by 0.8 percent in FY 2018.

3. Impact Analysis of Table II

Table II presents the projected impact of the proposed changes for FY 2018 for urban and rural hospitals and for the different categories of hospitals shown in Table I. It compares the estimated average payments per discharge for FY 2017 with the proposed estimated average payments per discharge for FY 2018, as calculated under our models. Therefore, this table presents, in terms of the average dollar amounts paid per discharge, the combined effects of the proposed changes presented in Table I. The proposed estimated percentage changes shown in the last column of Table II equal the estimated percentage changes in average payments per discharge from Column 8 of Table I.

<table>
<thead>
<tr>
<th>Number of hospitals</th>
<th>Estimated average FY 2017 payment per discharge</th>
<th>Estimated average FY 2018 payment per discharge</th>
<th>Proposed FY 2018 changes</th>
</tr>
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<td>All Hospitals</td>
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<td>12,041</td>
<td>1.7</td>
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<td>By Geographic Location:</td>
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<tr>
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<tr>
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<td>Number of hospitals</td>
<td>Estimated average FY 2017 payment per discharge</td>
<td>Estimated average FY 2018 payment per discharge</td>
</tr>
<tr>
<td>-------------------</td>
<td>--------------------</td>
<td>-----------------------------------------------</td>
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</tr>
<tr>
<td></td>
<td>(1)</td>
<td>(2)</td>
<td>(3)</td>
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<tr>
<td>Large urban areas</td>
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<td>12,953</td>
<td>13,174</td>
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<tr>
<td>Other urban areas</td>
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<td>11,516</td>
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<td>8,975</td>
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<td>Bed Size (Urban):</td>
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<tr>
<td>0–99 beds</td>
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<tr>
<td>100–199 beds</td>
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<td>10,423</td>
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<tr>
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<td></td>
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<td>50–99 beds</td>
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<tr>
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<td>11,374</td>
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<tr>
<td>Type of Ownership:</td>
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TABLE II—IMPACT ANALYSIS OF PROPOSED CHANGES FOR FY 2018 ACUTE CARE HOSPITAL OPERATING PROSPECTIVE PAYMENT SYSTEM—Continued
[Payments per discharge]

<table>
<thead>
<tr>
<th>Number of hospitals</th>
<th>Estimated average FY 2017 payment per discharge</th>
<th>Estimated average FY 2018 payment per discharge</th>
<th>Proposed FY 2018 changes</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(1)</td>
<td>(2)</td>
<td>(3)</td>
</tr>
<tr>
<td>Voluntary</td>
<td>1,914</td>
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<td>12,223</td>
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<td>Proprietary</td>
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<td>10,585</td>
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<tr>
<td>Government</td>
<td>514</td>
<td>12,805</td>
<td>13,012</td>
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<tr>
<td>Medicare Utilization as a Percent of Inpatient Days:</td>
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<td></td>
</tr>
<tr>
<td>0–25</td>
<td>509</td>
<td>15,200</td>
<td>15,448</td>
</tr>
<tr>
<td>25–50</td>
<td>2,113</td>
<td>11,775</td>
<td>11,983</td>
</tr>
<tr>
<td>50–65</td>
<td>535</td>
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<tr>
<td>Over 65</td>
<td>135</td>
<td>7,473</td>
<td>7,364</td>
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<td>FY 2018 Reclassifications by the Medicare Geographic Classification Review Board:</td>
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<tr>
<td>All Reclassified Hospitals</td>
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<td>11,720</td>
<td>11,914</td>
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<td>Non-Reclassified Hospitals</td>
<td>2,392</td>
<td>11,900</td>
<td>12,101</td>
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<td>Rural Hospitals Reclassified Full Year</td>
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<tr>
<td>All Section 401 Reclassified Hospitals</td>
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<td>12,157</td>
<td>12,376</td>
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<td>Other Reclassified Hospitals (Section 1886(d)(8)(B))</td>
<td>48</td>
<td>8,080</td>
<td>8,169</td>
</tr>
</tbody>
</table>

H. Effects of Other Proposed Policy Changes

In addition to those proposed policy changes discussed previously that we are able to model using our IPPS payment simulation model, we are proposing to make various other changes in this proposed rule. Generally, we have limited or no specific data available with which to estimate the impacts of these proposed changes. Our estimates of the likely impacts associated with these other proposed changes are discussed in this section.

1. Effects of Proposed Policy Relating to New Medical Service and Technology Add-On Payments

In section II.H. of the preamble to this proposed rule, we discuss six technologies for which we received applications for add-on payments for new medical services and technologies for FY 2018, as well as the status of the new technologies that were approved to receive new technology add-on payments in FY 2017. We note that three applicants withdrew their applications prior to the issuance of this proposed rule. As explained in the preamble to this proposed rule, add-on payments for new medical services and technologies under section 1886(d)(5)(K) of the Act are not required to be budget neutral. As discussed in section II.H.6 of the preamble of this proposed rule, we have not yet determined whether any of these six technologies for which we received applications for consideration for new technology add-on payments for FY 2018 will meet the specified criteria. Consequently, it is premature to estimate the potential payment impact of these six technologies for any potential new technology add-on payments for FY 2018. We note that if any of the six technologies are found to be eligible for new technology add-on payments for FY 2018, in the FY 2018 IPPS/LTCPPS final rule, we would discuss the estimated payment impact for FY 2018. In section II.H.5 of the preamble of this proposed rule, we are proposing to discontinue new technology add-on payments for Bimatumab (BLINCYTO™), CardioMEMSTM HF (Heart Failure) Monitoring System, the LUTONIX™ Drug Coated Balloon (DCB) Percutaneous Transluminal Angioplasty (PTA) and IN.PACT™ Admiral™ Paclitaxel Coated Percutaneous Transluminal Angioplasty (PTA) Balloon Catheter, and the MAGEC® Spinal Bracing and Distraction System (MAGEC® Spine) for FY 2018 because these technologies will have been on the U.S. market for 3 years. We also are proposing to continue to make new technology add-on payments for Defitelio® (Defibrotide), GORE® EXCLUDER® Iliac Branch Endoprostheses (IBE), Idarucizumab and Vistogard™ (Uridine Triacetate) in FY 2018 because these technologies would still be considered new. We note that new technology add-on payments for each case are limited to the lesser of (1) 50 percent of the costs of the new technology or (2) 50 percent of the amount by which the costs of the case exceed the standard MS–DRG payment for the case. Because it is difficult to predict the actual new technology add-on payment for each case, our estimates below are based on the increase in new technology add-on payments for FY 2018 as if every claim that would qualify for a new technology add-on payment would receive the maximum add-on payment. For Defitelio®, based on the applicant’s estimate from FY 2017, we currently estimate that new technology add-on payments for Defitelio® would increase overall FY 2018 payments by $5,161,200 (maximum add-on payment of $75,900 * 68 patients). Based on the applicant’s estimate for FY 2017, we currently estimate that new technology add-on payments for the GORE® EXCLUDER® IBE would increase overall FY 2018 payments by $5,683,750 (maximum add-on payment of $5,250 * 1,083 patients). Based on the applicant’s estimate for FY 2017, we currently estimate that new technology add-on payments for Idarucizumab would increase overall FY 2018 payments by $14,766,500 (maximum add-on payment of $1,750 * 8,438 patients). Based on the applicant’s estimate for FY 2017, we currently estimate that new technology add-on payments for Vistogard™ would increase overall FY 2018 payments by $3,242,500 (maximum add-on payment of $37,500 * 75 patients).

2. Effects of Proposed Changes to MS–DRGs Subject to the Postacute Care Transfer Policy and the MS–DRG Special Payment Policy

In section V.A. of the preamble of this proposed rule, we discuss our proposed changes to the list of MS–DRGs subject to the postacute care transfer policy and the DRG special payment policy. As reflected in Table 5 listed in section VI. of the Addendum to this proposed rule (which is available via the Internet on the CMS Web site), using criteria set forth in regulations at 42 CFR 412.4, we evaluated MS–DRG charge, discharge, and transfer data to determine which MS–DRGs qualify for the postacute care transfer and MS–DRG special payment policies. We note that we are not proposing to make any changes in these payment policies in this FY 2018 proposed rule. As a result of our proposals to revise the MS–DRG classifications for FY 2018, which are discussed in section II.F of the preamble of this proposed rule, we are proposing to add three MS–DRGs to the list of MS–DRGs subject to the postacute care transfer policy.
and the MS-DRG special payment policy. Column 4 of Table I in this Appendix A shows the effects of the proposed changes to the MS–DRGs and the relative payment weights and the application of the recalibration budget neutrality factor to the standard methodology in Section 1886(d)(4)(C)(ii) of the Act requires us annually to make appropriate DRG classification changes in order to reflect changes in treatment patterns, technology, and any other factors that may change the relative resources. The analysis and methods for determining the proposed changes due to the MS–DRGs and relative payment weights account for and include changes as a result of the proposed changes to the MS–DRGs subject to the MS–DRG postacute care transfer and MS–DRG special payment policies. We refer readers to section I.G. of this Appendix A for a detailed discussion of payment impacts due to the proposed MS–DRG reclassification policies for FY 2018.

3. Effects of the Proposed Changes to the Volume Decrease Adjustment for Sole Community Hospitals (SCHs)

In section V.C. of the preamble of this proposed rule, we discuss our proposal to modify the methodology used to calculate volume decrease adjustments for SCHs. We are proposing to prospectively require that the MACs compare Medicare revenue allocable to fixed costs from the cost reporting period when the hospital experienced the volume decrease to the hospital’s fixed costs from that same cost reporting period when calculating a volume decrease adjustment. We also are proposing that the cap will no longer be applied to the volume decrease adjustment calculation methodology in future periods. In addition, we are proposing to prospectively modify the volume decrease adjustment process to no longer require that a hospital explicitly demonstrate that it appropriately adjusted the number of staff in inpatient areas of the hospital based on the decrease in the number of inpatient days and to no longer require the MACs to adjust the volume decrease adjustment payment amount for excess staffing. We estimate that these proposed changes to the volume decrease adjustment would increase aggregate volume decrease adjustment payments by a total of approximately $15 million for cost reporting periods beginning in FY 2018. Given that the volume decrease adjustment is only available to SCHs and is predicated on the unanticipated nature of the volume decrease, it is difficult to predict how many hospitals will qualify for the adjustment in FY 2018. We assumed 20 hospitals would qualify for the adjustment in FY 2018 that the additional amount of the volume decrease adjustment payment based on our proposed methodology would be $750,000 per hospital.

4. Effects of Proposed Changes to Low-Volume Hospital Payment Adjustment Policy

In section V.E. of the preamble of this proposed rule, we discuss the expiration of the temporary changes to the low-volume hospital payment policy originally provided for by the Affordable Care Act and extended through FY 2017 by subsequent legislation. Effective for FY 2018 and subsequent years, qualifying hospitals must have less than 200 combined Medicare and non-Medicare discharges (instead of 1,600 Medicare discharges) and must be located more than 25 road miles from another subsection (d) hospital (instead of 15 road miles from another subsection (d) hospital). In this same section, we discuss our proposed parallel low-volume hospital payment adjustment regarding hospitals operated by the IHS or a Tribe. Under this proposal, an IHS hospital would be able to qualify for a low-volume hospital adjustment based on distance to the nearest IHS hospital, and a non-IHS hospital would be able to qualify to receive a low-volume hospital adjustment based on its distance to the nearest non-IHS hospital. Based upon the best available data at this time, we estimate the expiration of the temporary changes to the low-volume hospital payment and the proposed change to the low-volume payment adjustments would decrease aggregate low-volume payment adjustments from $315 million in FY 2017 to $4 million in FY 2018. This $311 million decrease is based on an estimated $314 million decrease in payments from the expiration of the temporary changes to the low-volume hospital definition and payment adjustment methodology together with an estimated increase of $3 million in payments made to hospitals that are expected to qualify under our proposed parallel low-volume hospital payment adjustment. These payment estimates were determined by identifying providers that, based on the best available data, are expected to qualify under the criteria that will apply in FY 2018 (that is, are located at least 25 miles from the nearest subsection (d) hospital and have less than 200 total discharges, and were determined from the same data used in developing the quantitative analyses of proposed changes in payments per case discussed previously in section I.C. of this Appendix A.

5. Effects of the Proposed Changes to Medicare DSH and Uncompensated Care Payments for FY 2018

As discussed in section V.G. of the preamble of this proposed rule, under section 333 of the Affordable Care Act, hospitals that are eligible to receive Medicare DSH payments will receive 25 percent of the amount they previously would have received under the statutory formula for Medicare DSH payments under section 1886(d)(5)(F) of the Act. The remainder, equal to an estimate of 75 percent of what formerly would have been paid as Medicare DSH payments (Factor 1), reduced to reflect changes in the percentage of uninsured individuals and additional statutory adjustments (Factor 2), is available to make additional payments to each hospital that qualifies for Medicare DSH payments and that has uncompensated care. Each hospital eligible for Medicare DSH payments will receive payments based on its estimated share of the total amount of uncompensated care for all hospitals eligible for Medicare DSH payments. The uncompensated care payment methodology has redistributive effects based on the proportion of a hospital’s uncompensated care relative to the uncompensated care for all hospitals eligible for Medicare DSH payments (Factor 3). For FY 2018, we are proposing a Factor 2 of 58.01 percent determined using the uninsured estimates produced by CMS’ Office of the Actuary (OACT) as part of the development of the National Health Expenditure Accounts (NHEA). We also are proposing to continue to use low-income insured patient days as a proxy for uncompensated care in combination with data on uncompensated care costs from Worksheet S–10 in the calculation of Factor 3. The uncompensated care payment methodology has redistributive effects based on the proportion of a hospital’s uncompensated care relative to the total uncompensated care for all hospitals eligible for Medicare DSH payments. The reduction to Medicare DSH payments under section 333 of the Affordable Care Act is not budget neutral.

In this proposed rule, we are proposing to establish the amount to be distributed as uncompensated care payments to SCH eligible hospitals, which for FY 2018 is $6,962,310,946.63, or 75 percent of what otherwise would have been paid for Medicare DSH payment adjustments adjusted by a proposed Factor 2 of 58.01 percent. For FY 2017, the amount available to be distributed for uncompensated care was $5,977,483,146.86, or 75 percent of what otherwise would have been paid for Medicare DSH payment adjustments adjusted by a Factor 2 of 55.36 percent. To calculate Factor 3 for FY 2018, we are proposing to use an average of data contained in the Medicaid days from hospitals’ 2012 and 2013 cost reports from the March 2017 update of the HCRIS database, uncompensated care costs from hospitals’ 2014 cost reports from the same extract of HCRIS, Medicaid days from 2012 cost report data submitted to CMS by IHS hospitals, and SSI days from the FY 2014 and FY 2015 SSI ratios. For each eligible hospital, we are proposing to calculate an individual Factor 3 for cost reporting years beginning during FYS 2012, 2013, and 2014. We will then add the individual amounts and divide the sum by three in order to calculate an average Factor 3 for FY 2018. For purposes of this proposed rule, we are using data from the December 2016 update of the HCRIS database for the Medicaid days component of the Factor 3 calculation as well as for the Worksheet S–10 uncompensated care cost component. For modeling purposes, as the FY 2015 SSI ratios are not yet available, we are using SSI days from the FY 2013 and FY 2014 SSI ratios, which are the most recent available SSI ratios. We expect the March 2017 update of the HCRIS database as well as the FY 2015 SSI ratios to be available in time for calculating Factor 3 for the FY 2018 IPPS/LTCF PFS final rule. The proposed FY 2018 policy of using data from hospitals’ FY 2012, FY 2013, and FY 2014 cost reports will be determined Factor 3 is based on our FY 2017 final policy (81 FR 56943 through 56973), which is in contrast to the methodology used in FY 2016, when we used Medicaid days from the more recent of a hospital’s full year 2012 or 2011 cost report from the March 2013 update of the HCRIS database, Medicaid days from
2012 cost report data submitted to CMS by IHS hospitals, and SSI days from the FY 2013 SSI ratios to calculate Factor 3. In addition, as explained in section V.G.4.c. of the preamble of this proposed rule, we are proposing to make several additional modifications to the Factor 3 methodology: (1) To annualize Medicaid data if a hospitals’ cost report does not equal 12 months of data; (2) to apply a scaling factor to the uncompensated care payment amount calculated for each DSH eligible hospital so that total uncompensated care payments are consistent with the estimated amount available to make uncompensated care payments for FY 2018; (3) to apply statistical trims to the CCRs on Worksheet S–10 that are considered anomalies to ensure reasonable CCRs are used to convert charges to costs for purposes of determining uncompensated care costs, and (4) to calculate Factor 3 for Puerto Rico hospitals and Indian Health Service and Tribal hospitals by substituting data regarding low-income insured days for FY 2013 for Worksheet S–10 data from FY 2014 cost reports. We also are proposing to continue the policies that were finalized in the FY 2015 IPPS/LTCH PPS final rule (79 FR 50020 through 50022) to address several issues concerning the process and data to be employed in determining Factor 3 in the case of hospital mergers for FY 2018 and subsequent years, as well as proposing to continue the policies finalized in the FY 2017 IPPS/LTCH PPS final rule concerning the methodology for calculating each hospital’s relative share of uncompensated care, such as combining data from multiple cost reports beginning in the same fiscal year and averaging the sum of three individual Factor 3s by the number of cost reporting periods with data.

To estimate the impact of the combined effect of changes in Factors 1 and 2, as well as the changes to the data used in determining Factor 3, on the calculation of Medicare DSH payments, including both empirically justified Medicare DSH payments and uncompensated care payments, we compared total DSH payments estimated in the FY 2017 IPPS/LTCH PPS final rule to total DSH payments estimated in this FY 2018 IPPS/LTCH PPS proposed rule. For FY 2017, for each hospital, we calculated the sum of: (1) 25 Percent of the estimated amount of what would have been paid as Medicare DSH payments in FY 2017 in the absence of section 3133 of the Affordable Care Act; and (2) 75 percent of the estimated amount of what would have been paid as Medicare DSH payments in the absence of section 3133 of the Affordable Care Act, adjusted by a Factor 2 of 55.36 percent and multiplied by a Factor 3 calculated as described in the FY 2017 IPPS/LTCH PPS final rule. For FY 2018, we calculated the sum of: (1) 25 Percent of the estimated amount of what would be paid as Medicare DSH payments in FY 2018 absent section 3133 of the Affordable Care Act; and (2) 75 percent of the estimated amount of what would be paid as Medicare DSH payments absent section 3133 of the Affordable Care Act, adjusted by a Factor 2 of 58.01 percent and multiplied by a Factor 3 calculated using the methodology described above.

Our analysis included 2,418 hospitals that are projected to be eligible for DSH in FY 2018. It did not include hospitals that had terminated their participation in the Medicare program as of February 23, 2017, Maryland hospitals, and SCHs that are expected to be paid based on their hospital-specific rates. In addition, data from merged or acquired hospitals were combined under the surviving hospital’s CCN, and the nonsurviving CCN was excluded from the analysis. The estimated impact of the proposed changes to Factors 1, 2, and 3 across all hospitals projected to be eligible for DSH payments in FY 2018, by hospital characteristic, is presented in the following table.

### Modeled Disproportionate Share Hospital Proposed Payments for Estimated FY 2018 DSHs by Hospital Type: Model DSH $ (in millions) From FY 2017 to FY 2018

<table>
<thead>
<tr>
<th>Number of estimated DSHs (FY 2018)</th>
<th>FY 2017 final rule estimated DSH $ (in millions)</th>
<th>FY 2018 proposed rule estimated DSH $ (in millions)</th>
<th>Dollar difference: FY 2017–FY 2018 $ (in millions)</th>
<th>Percent change **</th>
</tr>
</thead>
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<tr>
<td>Total</td>
<td></td>
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</tr>
<tr>
<td></td>
<td>2,418</td>
<td>$9,553</td>
<td>$10,931</td>
<td>$1,378</td>
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<tr>
<td>By Geographic Location:</td>
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<tr>
<td>Urban Hospitals</td>
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</tr>
<tr>
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<td>1,921</td>
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<td>10,355</td>
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<td>Large Urban Areas</td>
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<tr>
<td></td>
<td>1,037</td>
<td>5,717</td>
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<tr>
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<td></td>
<td>884</td>
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<td>Rural Hospitals</td>
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<tr>
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<tr>
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<td>240</td>
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<tr>
<td>100 to 249 Beds</td>
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<tr>
<td>250+ Beds</td>
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<td>100 to 249 Beds</td>
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<tr>
<td></td>
<td>114</td>
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<td>250+ Beds</td>
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<td></td>
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<td>72</td>
<td>140</td>
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Changes in projected FY 2018 DSH payments from DSH payments in FY 2017 are primarily driven by (1) proposed changes to Factor 1, which increased from $10.797 billion to $12.002 billion; (2) proposed changes to Factor 2, which increased from 55.36 percent to 58.01 percent; and (3) proposed changes to the data used to determine Factor 3. The proposed impact analysis found that, across all projected DSH eligible hospitals, FY 2018 DSH payments are estimated at approximately $10.930 billion, or an increase of approximately 14.4 percent from FY 2017 DSH payments (approximately $9.553 billion). While these proposed changes result in a net increase in the amount available to be distributed in uncompensated care payments, DSH payments to select hospital types are expected to decrease. This redistribution of DSH payments is caused by changes in the data used to determine Factor 3.

As seen in the above table, percent changes in DSH payments of less than 14.4 percent indicate that hospitals within the specified category are projected to experience a smaller increase in DSH payments, on average, compared to the universe of projected FY 2018 DSH hospitals. Conversely, percent changes in DSH payments that are greater than 14.4 percent indicate a hospital type is projected to have a larger increase than the overall average. The variation in the distribution of DSH payments by hospital characteristic is largely dependent on the change in a given hospital’s number of Medicare days and SSI days for purposes of the low-income insured days proxy between FY 2017 and FY 2018, as well as on its uncompensated care costs as reported on Worksheet S-10, used in the Factor 3 computation.

Many rural hospitals, grouped by geographic location, payment classification, and bed size, are projected to experience a larger increase in DSH payments than their urban counterparts. Overall, urban hospitals are projected to receive a 13.6 percent increase in DSH payments, and rural hospitals are projected to receive a 31.2 percent increase in DSH payments. However, only smaller rural hospitals are projected to receive larger than average increases in DSH payments, with rural hospitals that have 0–99 beds projected to experience a 52.7 percent payment increase, and larger rural hospitals with 250+ beds projected to experience a 46.7 percent payment increase. This trend is consistent with urban hospitals, in which the smallest urban hospitals (0–99 beds) are projected to receive an increase in DSH payments of 30.2 percent. Larger hospitals (100–250 beds and 250+ beds) are projected to receive increases of 14.3 and 13.0 percent in DSH payments, respectively, which are generally consistent with the overall average.

By region, projected DSH payment increases for urban hospitals are smallest in Pacific, Middle Atlantic, New England, and East South Central regions. The West South Central, Puerto Rico, and South Atlantic region hospitals are projected to receive a larger than average increase in DSH payments. Increases in remaining urban hospital regions are generally consistent with the overall average percent increase of 14.4. Regionally, rural hospitals are projected to receive a wider range of increases. Rural hospitals in the Pacific region are expected to receive a decrease in DSH payments (due to the reduction in the number of DSH hospitals in the region) while rural hospitals in the Middle Atlantic region are expected to receive virtually no change in in DSH payments, despite an estimated increase in the overall amount of DSH payments. Increases are projected to be substantially larger than the overall average in most regions, including West South Central, Mountain, New England, East North Central, South Atlantic, and West North Central regions.

Teaching hospitals with 100 or more residents are projected to receive relatively larger increases than teaching hospitals with fewer than 100 residents, although all are fairly consistent with the national average. Government hospitals are projected to receive larger than average increases, while voluntary hospitals are expected to receive increases generally consistent with the overall average. Proprietary hospitals are expected to receive smaller increases in DSH payments. Hospitals with 25 to 50 percent
Medicare utilization are projected to receive increases in DSH payments slightly below the overall average, while all other hospitals are projected to receive larger increases.

6. Effects of Proposed Reduction Under the Hospital Readmissions Reduction Program

In section V.I of the preamble of this proposed rule, we discuss our proposals for the FY 2018 Hospital Readmissions Reduction Program (established under section 3025 of the Affordable Care Act), which requires a reduction to a hospital’s base operating MS–DRG payments to account for excess readmissions. In this proposed rule, we estimate that 2,591 hospitals would have their base operating MS–DRG payments reduced by their proposed proxy FY 2018 hospital-specific readmissions adjustment. As a result, we estimate that the Hospital Readmissions Reduction Program would save approximately $564 million in FY 2018, an increase of $27 million over the estimated FY 2017 savings. This estimate is based on the same data used in developing the quantitative analyses of proposed changes in payments per case discussed previously in section I.G. of this Regulatory Impact Analysis, in conjunction with the FY 2017 hospital-specific readmissions adjustment factors and the proposed proxy FY 2018 hospital-specific readmissions adjustment factors found in Table 15 of this proposed rule (available only through the Internet as described in section VI of the Addendum to this proposed rule).

7. Effects of Proposed Changes Under the FY 2018 Hospital Value-Based Purchasing (VBP) Program

In section V.J. of the preamble of this proposed rule, we discuss the Hospital VBP Program under which the Secretary makes value-based incentive payments to hospitals based on their performance on measures during the performance period with respect to a fiscal year. These incentive payments will be funded for FY 2018 through a reduction to the FY 2018 base operating DRG payment amounts for all discharges for participating hospitals for such fiscal year, as required by section 1866(o)(6)(B) of the Act. The applicable percentage for FY 2018 and subsequent years is 2 percent. The total amount available for value-based incentive payments must be equal to the total amount of reduced payments for all hospitals for the fiscal year, as estimated by the Secretary.

In section V.I.1.b. of the preamble of this proposed rule, we estimate the available pool of funds for value-based incentive payments in the FY 2018 program year, which, in accordance with section 1886(o)(7)(C)(v) of the Act, will be 2.00 percent of base operating DRG payments, or a total of approximately $1.9 billion. We intend to update this estimate for the FY 2018 IPPS/LTCPPS final rule using the March 2017 update of the FY 2016 MedPAR file.

The proposed estimated impacts of the FY 2018 program year by hospital characteristic, found in the table below, are based on historical TPSs. We used the FY 2017 program year’s TPSs to calculate the proxy adjustment factors used for this impact analysis. These are the most recently available scores that hospitals were given an opportunity to review and correct. The proxy adjustment factors use estimated annual base operating DRG payment amounts derived from the December 2016 update to the FY 2016 MedPAR file. The proxy adjustment factors can be found in Table 16 associated with this proposed rule (available via the Internet on the CMS Web site at: http://www.cms.gov/Medicare/Medicare-Fee-For-Service-Payment/AcuteInpatientPPS/index.html).

The impact analysis shows that, for the FY 2018 program year, the number of hospitals that would receive an increase in their base operating DRG payment amounts is higher than the number of hospitals that would receive a decrease. Among urban hospitals, those in the New England, South Atlantic, East North Central, East South Central, West North Central, West South Central, Mountain, and Pacific regions would have an increase, on average, in their base operating DRG payment amounts. Urban hospitals in the Middle Atlantic region would receive an average decrease in their base operating DRG payment amounts. Among rural hospitals, those in all regions would have an increase, on average, in their base operating DRG payment amounts. On average, hospitals that receive a higher (over 65 percent of DSH payments) would receive decreases in base operating DRG payment amounts. With respect to hospitals’ Medicare utilization as a percent of inpatient days (MCR), those hospitals with an MCR above 65 percent would have the largest average increase in base operating DRG payment amounts. Nonteaching hospitals would have an average increase, and teaching hospitals would experience an average decrease in base operating DRG payment amounts.

### Impact Analysis of Base Operating DRG Payment Amount Proposed Changes Resulting from the FY 2018 Hospital VBP Program

<table>
<thead>
<tr>
<th>By Geographic Location:</th>
<th>Number of hospitals</th>
<th>Average percentage change</th>
</tr>
</thead>
<tbody>
<tr>
<td>All Hospitals</td>
<td>2,955</td>
<td>0.183</td>
</tr>
<tr>
<td>Large Urban</td>
<td>1,227</td>
<td>0.094</td>
</tr>
<tr>
<td>Other Urban</td>
<td>1,048</td>
<td>0.152</td>
</tr>
<tr>
<td>Rural Area</td>
<td>680</td>
<td>0.392</td>
</tr>
<tr>
<td>Urban hospitals</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0–99 beds</td>
<td></td>
<td></td>
</tr>
<tr>
<td>100–199 beds</td>
<td>2,275</td>
<td>0.121</td>
</tr>
<tr>
<td>200–299 beds</td>
<td>486</td>
<td>0.685</td>
</tr>
<tr>
<td>300–499 beds</td>
<td>721</td>
<td>0.082</td>
</tr>
<tr>
<td>500 or more beds</td>
<td>434</td>
<td>-0.039</td>
</tr>
<tr>
<td>Rural hospitals</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0–99 beds</td>
<td>680</td>
<td>0.392</td>
</tr>
<tr>
<td>100–199 beds</td>
<td>207</td>
<td>0.612</td>
</tr>
<tr>
<td>200–299 beds</td>
<td>276</td>
<td>0.398</td>
</tr>
<tr>
<td>300–499 beds</td>
<td></td>
<td></td>
</tr>
<tr>
<td>500 or more beds</td>
<td>114</td>
<td>0.243</td>
</tr>
<tr>
<td>By Region:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Urban By Region</td>
<td></td>
<td></td>
</tr>
<tr>
<td>New England</td>
<td>2,275</td>
<td>0.121</td>
</tr>
<tr>
<td>Middle Atlantic</td>
<td>110</td>
<td>0.072</td>
</tr>
<tr>
<td>South Atlantic</td>
<td>297</td>
<td>-0.119</td>
</tr>
<tr>
<td>East North Central</td>
<td>387</td>
<td>0.025</td>
</tr>
<tr>
<td>East South Central</td>
<td>364</td>
<td>0.217</td>
</tr>
<tr>
<td>West North Central</td>
<td>135</td>
<td>0.009</td>
</tr>
<tr>
<td>West South Central</td>
<td>152</td>
<td>0.451</td>
</tr>
<tr>
<td>Mountain</td>
<td>320</td>
<td>0.194</td>
</tr>
<tr>
<td></td>
<td>156</td>
<td>0.058</td>
</tr>
</tbody>
</table>
Program for FY 2018, including adoption of finalized changes to the HAC Reduction rule. In the FY 2017 IPPS/LTCH PPS final proposed measures and scoring system for the estimated cumulative effect of the changes to the HAC Reduction Program for 20210 Federal Register

Actual FY 2018 program year’s TPSs will not be reviewed and corrected by hospitals until after the FY 2018 IPPS/LTCH PPS final rule has been published. Therefore, the same historical universe of eligible hospitals and corresponding TPSs from the FY 2017 program year will be used for the updated impact analysis in that final rule.

8. Effects of Proposed Changes to the HAC Reduction Program for FY 2018

In section V.K. of the preamble of this proposed rule, we discuss the proposed changes to the HAC Reduction Program for FY 2018. The table and analysis below show the estimated cumulative effect of the proposed measures and scoring system for the HAC Reduction Program in this proposed rule. In the FY 2017 IPPS/LTCH PPS final rule (81 FR 57013 through 57025), we finalized changes to the HAC Reduction Program for FY 2018, including adoption of the modified PSI 90 Composite, defining the applicable time period, and changes to the scoring methodology (adoption of the z-score method for calculating measure scores). Based on this z-score methodology, the table below presents data on the estimated proportion of hospitals in the worst-performing quartile of the Total HAC Scores by hospital characteristic. We note that because scores will undergo a 30-day review and correction period by the hospitals that will not conclude until after the publication of the FY 2018 IPPS/LTCH PPS final rule, we are not providing hospital-level data or a hospital-level payment impact in conjunction with this FY 2018 IPPS/LTCH PPS proposed rule.

To estimate the impact of the FY 2018 HAC Reduction Program, we used, as previously finalized, AHRQ PSI 90 measure results based on Medicare FFS discharges from July 2014 through September 2015 and version 6.0 (recalibrated) of the AHRQ software. For the CLABSI, CAUTI, Colon and Abdominal Hysterectomy SSI, MRSA Bacteremia, and CDI measure results, we used standardized infection ratios (SIRs) calculated with hospital surveillance data reported to the NHSN for infections occurring between January 1, 2014 and December 31, 2015. We noted that actual FY 2018 HACRP results will be calculated using CDC NHSN data from CYs 2015 and 2016 and will use the re-baselined values and expansion to non-ICU wards but could not be presented here due to data timelines.

We note that, at this time, we are unable to provide the estimated impact of the FY 2018 HAC Reduction Program due to an error in the version 6.0 (recalibrated) AHRQ software. We anticipate that we will be able to provide this information in the FY 2018 final rule. We have provided the final impacts from the FY 2017 final rule as estimated impacts for FY 2018.

**ESTIMATED PROPORTION OF HOSPITALS IN THE WORST-PERFORMING QUARTILE (>75TH PERCENTILE) OF THE TOTAL HAC SCORE FOR THE FY 2017 HAC REDUCTION PROGRAM**

<table>
<thead>
<tr>
<th>Hospital characteristic</th>
<th>Number of hospitals</th>
<th>Number of hospitals in the worst-performing quartile</th>
<th>Percent of hospitals in the worst-performing quartile</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total</td>
<td>3,215</td>
<td>771</td>
<td>24.0</td>
</tr>
<tr>
<td>By Geographic Location:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All hospitals:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Urban</td>
<td>2,404</td>
<td>653</td>
<td>27.2</td>
</tr>
<tr>
<td>Rural</td>
<td>796</td>
<td>107</td>
<td>13.4</td>
</tr>
<tr>
<td>Urban hospitals:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1–99 beds</td>
<td>592</td>
<td>91</td>
<td>15.4</td>
</tr>
</tbody>
</table>

*By hospital characteristic*
### ESTIMATED PROPORTION OF HOSPITALS IN THE WORST-PERFORMING QUARTILE (>75TH PERCENTILE) OF THE TOTAL HAC SCORE FOR THE FY 2017 HAC REDUCTION PROGRAM—Continued

[By hospital characteristic]

<table>
<thead>
<tr>
<th>Hospital characteristic</th>
<th>Number of hospitals</th>
<th>Number of hospitals in the worst-performing quartile</th>
<th>Percent of hospitals in the worst-performing quartile</th>
</tr>
</thead>
<tbody>
<tr>
<td>100–199 beds</td>
<td>734</td>
<td>166</td>
<td>22.6</td>
</tr>
<tr>
<td>200–299 beds</td>
<td>440</td>
<td>134</td>
<td>30.5</td>
</tr>
<tr>
<td>300–399 beds</td>
<td>276</td>
<td>101</td>
<td>36.6</td>
</tr>
<tr>
<td>400–499</td>
<td>150</td>
<td>61</td>
<td>40.7</td>
</tr>
<tr>
<td>500 or more beds</td>
<td>212</td>
<td>100</td>
<td>47.2</td>
</tr>
</tbody>
</table>

Rural hospitals:

<p>| | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1–49 beds</td>
<td>303</td>
<td>48</td>
<td>15.8</td>
</tr>
<tr>
<td>50–99 beds</td>
<td>289</td>
<td>29</td>
<td>10.0</td>
</tr>
<tr>
<td>100–149 beds</td>
<td>118</td>
<td>11</td>
<td>9.3</td>
</tr>
<tr>
<td>150–199 beds</td>
<td>45</td>
<td>9</td>
<td>20.0</td>
</tr>
<tr>
<td>200 or more beds</td>
<td>41</td>
<td>10</td>
<td>24.4</td>
</tr>
</tbody>
</table>

By Region:

<table>
<thead>
<tr>
<th>Region</th>
<th>Number of hospitals</th>
<th>Number of hospitals in the worst-performing quartile</th>
<th>Percent of hospitals in the worst-performing quartile</th>
</tr>
</thead>
<tbody>
<tr>
<td>New England</td>
<td>134</td>
<td>42</td>
<td>31.3</td>
</tr>
<tr>
<td>Mid-Atlantic</td>
<td>365</td>
<td>131</td>
<td>35.9</td>
</tr>
<tr>
<td>South Atlantic</td>
<td>519</td>
<td>133</td>
<td>25.6</td>
</tr>
<tr>
<td>East North Central</td>
<td>494</td>
<td>96</td>
<td>19.4</td>
</tr>
<tr>
<td>East South Central</td>
<td>295</td>
<td>45</td>
<td>15.3</td>
</tr>
<tr>
<td>West North Central</td>
<td>259</td>
<td>38</td>
<td>14.7</td>
</tr>
<tr>
<td>West South Central</td>
<td>511</td>
<td>104</td>
<td>20.4</td>
</tr>
<tr>
<td>Mountain</td>
<td>226</td>
<td>55</td>
<td>24.3</td>
</tr>
<tr>
<td>Pacific</td>
<td>397</td>
<td>116</td>
<td>29.2</td>
</tr>
</tbody>
</table>

By DSH Percent:

<table>
<thead>
<tr>
<th>DSH Percent</th>
<th>Number of hospitals</th>
<th>Number of hospitals in the worst-performing quartile</th>
<th>Percent of hospitals in the worst-performing quartile</th>
</tr>
</thead>
<tbody>
<tr>
<td>0–24</td>
<td>1,387</td>
<td>321</td>
<td>23.1</td>
</tr>
<tr>
<td>25–49</td>
<td>1,454</td>
<td>324</td>
<td>22.3</td>
</tr>
<tr>
<td>50–64</td>
<td>181</td>
<td>58</td>
<td>32.0</td>
</tr>
<tr>
<td>65 and over</td>
<td>178</td>
<td>57</td>
<td>32.0</td>
</tr>
</tbody>
</table>

By Teaching Status:

<table>
<thead>
<tr>
<th>Teaching Status</th>
<th>Number of hospitals</th>
<th>Number of hospitals in the worst-performing quartile</th>
<th>Percent of hospitals in the worst-performing quartile</th>
</tr>
</thead>
<tbody>
<tr>
<td>Non-teaching</td>
<td>2,160</td>
<td>381</td>
<td>17.6</td>
</tr>
<tr>
<td>Fewer than 100 residents</td>
<td>790</td>
<td>237</td>
<td>30.0</td>
</tr>
<tr>
<td>100 or more residents</td>
<td>250</td>
<td>142</td>
<td>56.8</td>
</tr>
</tbody>
</table>

By Type of Ownership:

<table>
<thead>
<tr>
<th>Type of Ownership</th>
<th>Number of hospitals</th>
<th>Number of hospitals in the worst-performing quartile</th>
<th>Percent of hospitals in the worst-performing quartile</th>
</tr>
</thead>
<tbody>
<tr>
<td>Voluntary</td>
<td>1,868</td>
<td>478</td>
<td>25.6</td>
</tr>
<tr>
<td>Proprietary</td>
<td>825</td>
<td>154</td>
<td>18.7</td>
</tr>
<tr>
<td>Government</td>
<td>485</td>
<td>121</td>
<td>24.9</td>
</tr>
</tbody>
</table>

By MCR Percent:

<table>
<thead>
<tr>
<th>MCR Percent</th>
<th>Number of hospitals</th>
<th>Number of hospitals in the worst-performing quartile</th>
<th>Percent of hospitals in the worst-performing quartile</th>
</tr>
</thead>
<tbody>
<tr>
<td>0–24</td>
<td>472</td>
<td>148</td>
<td>31.4</td>
</tr>
<tr>
<td>25–49</td>
<td>2,106</td>
<td>481</td>
<td>22.8</td>
</tr>
<tr>
<td>50–64</td>
<td>518</td>
<td>104</td>
<td>20.1</td>
</tr>
<tr>
<td>65 and over</td>
<td>80</td>
<td>18</td>
<td>22.5</td>
</tr>
</tbody>
</table>


- 9. Effects of Implementation of the Additional 5-Year Extension of the Rural Community Hospital Demonstration Program

In section V.L. of the preamble of this proposed rule, we discuss our proposed implementation of section 410A of Public Law 108–173, as amended by sections 3123 and 10313 of Public Law 111–148, and more recently, by section 15003 of Public Law 114–255, which requires the Secretary to conduct a demonstration that tests the feasibility and advisability of establishing “rural community” hospitals to furnish covered inpatient hospital services to Medicare beneficiaries. The demonstration makes payments under a reasonable cost methodology for covered inpatient hospital services furnished to Medicare beneficiaries by up to 30 rural hospitals. Section 15003 of Public Law 114–255, enacted December 13, 2016, requires a 10-year extension period (in place of the 5-year extension required by Public Law 111–148) for the demonstration. Therefore, the Secretary is required to conduct the demonstration for an additional 5-year period. Section 15003 of Public Law 114–255 also requires that, no later than 120 days after enactment of Public Law 114–255, the Secretary issue a solicitation for applications to select additional hospitals to
participate in the demonstration program for the second 5 years of the 10-year extension period so long as the maximum number of 30 hospitals stipulated by Public Law 111–148 is not exceeded. Section 410A(c)(2) of Public Law 108–173 requires that, in conducting the demonstration program under this section, the Secretary shall ensure that the aggregate payments made by the Secretary do not exceed the amount which the Secretary would have paid if the demonstration program under this section was not implemented and budget neutrality was maintained.

In this FY 2018 IPPS/LTC PPS proposed rule, we describe our proposals for implementation of the extension under section 15003 of Public Law 114–255, the proposed budget neutrality methodology for the extension period authorized by the legislation, and the proposed reconciliation of actual and estimated costs of the demonstration for previous years (2011 through 2016). Our proposal for budget neutrality would adopt the general methodology and approaches from previous years for the demonstration. As discussed in section V.L. of the preamble of this proposed rule, in the IPPS final rules from FYs 2005 through 2016, we have estimated the additional payments for each of the participating hospitals as a result of the demonstration. In order to achieve budget neutrality, we have adjusted the national IPPS rates by an amount sufficient to account for the added costs of this demonstration. In other words, we have applied budget neutrality across the payment system as a whole rather than across the participating hospitals.

The language of the statutory budget neutrality requirement permits the agency to implement the budget neutrality provision in this manner. The statutory language requires that aggregate payments made by the Secretary do not exceed the amount which the Secretary would have paid if the demonstration was not implemented, but does not identify the range across which aggregate payments must be held equal.

Section 15003 of Public Law 114–255 requires us to conduct the Rural Community Hospital Demonstration for a 10-year extension period (in place of the 5-year extension period required by Public Law 108–173), beginning on the date immediately following the last day of the initial 5-year period under section 410A(a)(5) of Public Law 108–173. Specifically, section 15003 of Public Law 114–255 amended section 410Ag(4) of Public Law 108–173 to require that, for hospitals participating in the demonstration as of the last day of the initial 5-year period, the Secretary shall provide for continued participation of such rural community hospitals in the demonstration during the 10-year extension period, unless the hospital makes an election to discontinue participation. Furthermore, section 15003 of Public Law 114–255 added subsection (g)(5) to section 410Ag(4) of Public Law 108–173 which provides for participation under the demonstration during the second 5 years of the 10-year extension period for hospitals that are not described in section 410Ag(4) of Public Law 108–173, but that were participating in the demonstration as of December 30, 2014, unless the hospital makes an election to discontinue participation.

We are proposing to implement the second 5 years of the 10-year extension period in a way that recognizes a gap in participation for the previously participating hospitals between participation in the second 5 years and the start of the second 5 years of the extension period, and that provides for alignment of the periods of performance under the extension among all participating hospitals. Thus, for each previously participating hospital that decides to participate in the second 5 years of the 10-year extension period, we are proposing that the start date for the period of performance under the second 5-year extension period would be the start of the first cost reporting period on or after October 1, 2017 following upon the announcement of the selection of the additional hospitals for the demonstration. Our goal is to finalize this selection by June 2017, in time to include in the FY 2018 IPPS final rule an estimate of the costs of the demonstration during FY 2018 and the fiscal year. We have calculated this offset amount for these newly participating hospitals, as well as for those hospitals among the previously participating hospitals that decide to participate in the second 5 years of the 10-year extension period.

We are proposing that if the selection of the additional hospitals under the solicitation is not announced by June 2017, we would include the estimated costs of the demonstration for all participating hospitals for FY 2018 in the budget neutrality offset amount to be calculated in the FY 2019 IPPS/LTC PPS proposed and final rule.

In section V.L. of the preamble of this proposed rule, we also describe an alternative approach that we considered, under which each previously participating hospital would begin the second 5 years of the 10-year extension period on the date immediately after the date of the period of performance under the first 5-year extension period ended. In addition, we describe the methodology that we considered for calculating the budget neutrality offset amount under this alternative approach. We are inviting public comments on this alternative approach and calculation methodology.

In previous years, we have incorporated a second component into the budget neutrality offset amounts identified in the final IPPS rules. As finalized cost reports became available, we determined the amount by which the actual costs of the demonstration for an earlier, given year differed from the estimated costs for the demonstration set forth in the final IPPS rule for the corresponding fiscal year, and we incorporated that amount into the budget neutrality offset amount for the upcoming fiscal year. We have calculated this difference for FYs 2005 through 2010 between the actual costs of the demonstration as determined from cost reports once available, and estimated costs of the demonstration as identified in the applicable IPPS final rules for these years.

With the extension of the demonstration for another 5-year period, as authorized by section 15003 of Public Law 114–255, we are proposing to continue this general procedure. Specifically, we are proposing that when finalized cost reports for FYs 2011, 2012, and 2013 are available, we would include this difference for these years in the budget neutrality offset adjustment to be applied to the national IPPS rates in a future final rule. We indicate that this will be effective in FY 2019. We also are proposing that when finalized cost reports for FYs 2014 through 2016 are available, we would include the difference between the actual costs as reflected on these cost reports and the estimated amounts included in the budget neutrality offset amounts for these fiscal years in a future final rule.

As discussed in section V.L. of the preamble of this proposed rule, depending on when the selection of additional hospitals as authorized by section 15003 of Public Law 114–225 is finalized, the estimate of the cost of the demonstration for FY 2018 will be formulated and included in the budget neutrality offset amount in either the FY 2018 final rule or the FY 2019 proposed and final rules. Therefore, the FY 2018 IPPS/LTC PPS proposed rule sets forth our proposed budget neutrality offset methodology, it does not include a specific budget neutrality offset amount.

10. Effects of the Proposed Changes Relating to Provider-Based Status of Indian Health Service and Tribal Facilities and Organizations

In section V.N. of the preamble of this proposed rule, we discuss our proposals relating to provider-based status of Indian Health Service (IHS) and tribal facilities and organizations. Regulations at § 413.65(m) currently grandfather facilities from provider-based regulations if they meet certain criteria, including on or before April 7, 2000, having furnished only services that were billed as if they had been furnished by a department of a hospital operated by the IHS or a Tribe. We have also issued subregulatory guidance on circumstances that would not result in a facility or organization losing its grandfathered status. After consideration of the special and legally recognized relationship between Indian Tribes and the U.S. Government, as well as current IHS policies and procedures, we are proposing to remove the date limitation in § 413.65(m) that restricted the grandfathering provision to IHS or Tribal facilities and organizations furnishing services on or before April 7, 2000. We also are proposing to make a technical change to make the regulation text more consistent with our current rules that require these facilities to comply with all applicable Medicare conditions of participation that apply to the main provider.

We do not expect any significant payment impact because these proposals are in line with current guidance, and we believe that IHS policies and procedures regarding the planning, operation, and funding of such facilities are resulting in appropriate Medicare payments.

11. Effects of the Proposed Changes Relating to Hospital-within-Hospital (HwH) Policy

In section VII.B. of the preamble of this proposed rule, we discuss our proposal to revise the regulations applicable to HwHs so that the separateness and control
requirements would only apply to IPPS-excluded HwHs that are co-located with IPPS hospitals beginning in FY 2018. This proposal is premised on the belief that the policy concerns that underlie our existing HwH regulations (that is, inappropriate patient routing and hospitals acting as illegal de facto units) are sufficiently moderated in situations where IPPS-excluded hospitals are co-located with other but not IPPS hospitals, in large part due to the payment system changes that have occurred over the intervening years for IPPS-excluded hospitals. In addition, we are proposing to revise the HwH requirements to no longer require the provisions that outline performance of basic hospital functions in order to maintain IPPS-exclusion beginning in FY 2018. This proposed revision would not result in a practical change to how HwHs are currently operated because the performance of basic hospital functions that are required under the HwH regulations are currently addressed under CMS\’ interpreting guidelines for the hospital conditions of participation. We do not expect any significant payment impact because these proposals are primarily administrative in nature or in line with current guidance.

12. Effects of Continued Implementation of the Frontier Community Health Integration Project (FCHIP) Demonstration

In section VIII.C.2. of the preamble of this proposed rule, we discuss the implementation of the FCHIP demonstration, which allows eligible entities to develop and test new models for the delivery of health care services in eligible counties in order to improve access to and better integrate the delivery of acute care, extended care, and other health care services to Medicare beneficiaries in no more than four States. Section 123(g)(1)(B) of Public Law 110–275 requires that the demonstration be budget neutral. Specifically this provision states that, in each demonstration, the Secretary shall ensure that the aggregate payments made by the Secretary do not exceed the amount which the Secretary estimates would have been paid if the demonstration were not implemented. Furthermore, section 123(j) of Public Law 110–275 states that the Secretary may waive the requirements of Titles XVIII and XIX of the Act as may be necessary and appropriate for the purpose of carrying out the demonstration, thus allowing the waiver of Medicare payment rules encompassed in the demonstration. Budget neutrality estimates for the demonstration will be based on the demonstration period of August 1, 2016 through July 31, 2019. The demonstration includes three intervention phases, on which specific provisions of the Medicare payment rules will allow for enhanced payment: Telehealth, skilled nursing facility/nursing facility services, and ambulance services. These waivers are being implemented with the goal of improving access to care without a net increase in costs. (We initially addressed this demonstration in the FY 2017 IPPS/LTC PP S final rule (81 FR 57064 through 57065).)

We specified waivers and payment enhancements for the demonstration and selected CAHs for participation with the goal of maintaining the budget neutrality of the demonstration on its own terms (that is, the demonstration will produce savings from reduced transfers and admissions to other health care providers, thus offsetting any increase in payments resulting from the demonstration). However, because of the small size of this demonstration and uncertainty associated with projected Medicare utilization and costs, in the FY 2017 IPPS/LTC PPS final rule (81 FR 57064 through 57065), we adopted a contingency plan to ensure that the neutrality requirement in section 123 of Public Law 110–275 is met. Accordingly, if analysis of claims data for the Medicare beneficiaries receiving services at each of the participating CAHs, as well as of other data sources, including cost reports, shows that increases in Medicare payments under the demonstration during the 3-year period are not sufficiently offset by reductions elsewhere, we will recoup the additional expenditures attributable to the demonstration through a reduction in payments to all CAHs nationwide. The demonstration is projected to impact payments to participating CAHs under both Medicare Part A and Part B. Thus, in the event that we determine that aggregate payments under the demonstration exceed the payments that would otherwise have been made, CMS will recoup payments through reductions of Medicare payments to all CAHs under both Medicare Part A and Part B. Because of the small scale of the demonstration, it will be feasible to implement budget neutrality by reducing payments only to the participating CAHs. Therefore we will make the reduction to payments to all CAHs, not just those participating in the demonstration, because the FCHIP demonstration is specifically designed to test innovations that affect delivery of services by this provider category. As we explained in the FY 2017 IPPS/LTC PPS final rule (81 FR 57065), we believe that the language of the statutory budget neutrality requirement in section 123(g)(1)(B) of the Act permits the agency to implement the budget neutrality provision in this manner. The statutory language merely refers to ensuring that aggregate payments made by the Secretary do not exceed the amount which the Secretary estimates would have been paid if the demonstration project was not implemented, and does not identify the range across which aggregate payments must be held equal.

Given the 3-year period of performance of the FCHIP demonstration, the time needed to conduct the budget neutrality analysis, in the event the demonstration is found not to have been budget neutral, we plan to recoup any excess costs over a period of three cost report periods, beginning in CY 2020. Therefore, this policy has no impact for any national payment system for FY 2018.

1. Effects of Proposed Changes in the Capital IPPS

1. General Considerations

For the impact analysis presented below, we used data from the December 2016 update of the FY 2016 MedPAR file and the December 2016 update of the provider-

Specific File (PSF) that is used for payment purposes. Although the analyses of the proposed changes to the capital prospective payment system do not incorporate cost data, we used the December 2016 update of the most recently available hospital cost report data (FYs 2013 and 2014) to categorize hospitals. Our analysis has several qualifications. We use the best data available and make assumptions about case-mix and beneficiary enrollment as described later in this section. Due to the interdependent nature of the IPPS, it is very difficult to precisely quantify the impact associated with each change. In addition, we draw upon various sources for the data used to categorize hospitals in the tables. In some cases (for instance, the number of beds), there is a fair degree of variation in the data from different sources. We have attempted to construct these variables with the best available sources overall. However, it is possible that some individual hospitals are placed in the wrong category.

Using cases from the December 2016 update of the FY 2016 MedPAR file, we simulated payments under the capital IPPS for FY 2017 and proposed payments for FY 2018 for a comparison of total payments per case. Any short-term, acute care hospitals not paid under the general IPPS (for example, hospitals in Maryland) are excluded from the simulations.

The methodology for determining a capital IPPS payment is set forth at § 412.312. The basic methodology for calculating the proposed capital IPPS payments in FY 2018 is as follows: (Standard Federal Rate) × (DRG weight) × (GAF) × (COLA for hospitals located in Alaska and Hawaii) × (1 + DSH Adjustment Factor + IME adjustment factor, if applicable). In addition to the other adjustments, hospitals may receive outlier payments for those cases that qualify under the threshold established for each fiscal year. We modeled payments for each hospital by multiplying the capital Federal rate by the GAF and the hospital’s case-mix. We then added estimated payments for indirect medical education, disproportionate share, and outliers, if applicable. For purposes of this impact analysis, the model includes the following assumptions:

- We estimate that the Medicare case-mix index will increase by 0.5 percent in both FYs 2017 and 2018.
- We estimate that Medicare discharges will be approximately 11.1 million in FY 2016 and 11.3 million in FY 2018.
- The capital Federal rate was updated beginning in FY 1996 by an analytical framework that considers changes in the prices associated with capital-related costs and adjustments to account for forecast error, changes in the case-mix index, allowable changes in intensity, and other factors. As discussed in section III.A.1.a. of the Addendum to this proposed rule, the proposed update is 1.2 percent for FY 2018.
- In addition to the proposed FY 2018 update factor, the proposed FY 2018 capital Federal rate was calculated based on a proposed GAF/DRG budget neutrality
adjustment factor of 0.9992, a proposed outlier adjustment factor of 0.9434, and an adjustment to remove the one-time prospective adjustment of 1.006 made in FY 2017 to address the effect of the 0.2 percent reduction to the national capital Federal rates in effect for FY 2014, FY 2015, and FY 2016 relating to the 2-midnight policy. The 2-midnight adjustment that was finalized in the FY 2017 IPPS/LTCH PPS final rule (81 FR 57294) is discussed in section V.C. of the preamble of this proposed rule as it relates to the capital Federal rate. As also discussed in section V.C. of the preamble of this proposed rule, we are not proposing to make an additional MS–DRG documentation and coding adjustment to the capital IPPS Federal rate for FY 2018.

2. Results

We used the actuarial model previously described in section I.I. of Appendix A of this proposed rule to estimate the potential impact of our proposed changes for FY 2018 on total capital payments per case, using a universe of 3,292 hospitals. As previously described, the individual hospital payment parameters are taken from the best available data, including the December 2016 update of the FY 2016 MedPAR file, the December 2016 update to the PSF, and the most recent cost report data from the December 2016 update of HCRIS. In Table III, we present a comparison of estimated total payments per case for FY 2017 and estimated proposed total payments per case for FY 2018 based on the proposed payment policies. Column 2 shows estimates of payments per case under our model for FY 2017. Column 3 shows estimates of proposed payments per case under our model for FY 2018. Column 4 shows the total percentage change in payments from FY 2017 to FY 2018. The change represented in Column 4 includes the proposed 1.2 percent update to the capital Federal rate and other proposed changes in the adjustments to the capital Federal rate. The comparisons are provided by: (1) Geographic location; (2) region; and (3) payment classification.

The simulation results show that, on average, proposed capital payments per case in FY 2018 are expected to increase as compared to capital payments per case in FY 2017. This expected increase overall is due to the proposed approximately 1.2 percent update to the capital Federal rate for FY 2018, as well as the proposed outlier adjustment of 0.9434 which is a 0.51 percent change from the FY 2017 outlier adjustment of 0.9386. The change in the outlier adjustment is expected to increase capital payments per case for most hospitals to a lesser or greater extent, as are changes in the DRGs. The expected increase in capital payments per case as a result of MS–DRG changes is somewhat larger for hospitals in rural areas than for hospitals in rural areas. (For a discussion of the determination of the capital Federal rate and adjustments, we refer readers to section III.A. of the Addendum to this proposed rule.) Over all hospitals, the proposed changes to the GAFs have no effect on capital payments per case. However, by region, hospitals within both rural and urban regions may experience an increase or a decrease in capital payments per case due to proposed changes in the GAFs. These regional effects of the proposed changes to the GAFs on capital payments are consistent with the projected changes in payments due to proposed changes in the wage index (and policies affecting the wage index) as shown in Table I in section I.G. of this Appendix A.

The net impact of these proposed changes is an estimated 2.4 percent change in capital payments per case from FY 2017 to FY 2018 as compared to FY 2017. Capital IPPS payments per case for hospitals in large urban areas would increase by an estimated 2.8 percent, while hospitals in rural areas, on average, are expected to experience a 2.6 percent increase in capital payments per case from FY 2017 to FY 2018. Capital IPPS payments per case for other urban hospitals are estimated to increase 1.7 percent.

The comparisons by region show that the estimated increases in capital payments per case from FY 2017 to FY 2018 in urban areas would range from a 3.6 percent increase for the West South Central urban region to a 1.1 percent increase for the Mountain urban region. For rural regions, the New England rural region is projected to experience the largest increase in capital IPPS payments per case of 5.0 percent, while the South Atlantic rural region is projected to experience an increase in capital IPPS payments per case of 1.4 percent.

Hospitals of all types of ownership (that is, voluntary hospitals, government hospitals, and proprietary hospitals) are expected to experience an increase in capital payments per case from FY 2017 to FY 2018. The proposed increase in capital payments for voluntary hospitals is estimated to be 2.0 percent and for government hospitals, the increase is estimated to be 3.4 percent. Proprietary hospitals are expected to experience an increase in capital IPPS payments of 3.0 percent.

Section 1886(d)(10) of the Act established the MGCRB. Hospitals may apply for reclassification for purposes of the wage index for FY 2018. Reclassification for wage index purposes also affects the GAFs because that factor is constructed from the hospital wage index. To present the effects of the hospitals being reclassified as of the publication of this proposed rule for FY 2018, we show the average capital payments per case for reclassified hospitals for FY 2018. Urban reclassified hospitals are expected to experience an increase in capital payments of 1.6 percent; urban nonreclassified hospitals are expected to experience an increase in capital payments of 2.8 percent. The estimated percentage increase for rural reclassified hospitals is 2.4 percent, and for rural nonreclassified hospitals, the estimated increase is 2.6 percent. Hospitals reclassified under section 401 are among the few groups of hospitals not expected to experience an increase in capital payments—it is expected that these hospitals would experience a decrease in capital payments of 1.8 percent, while capital payments for other reclassified hospitals are expected to increase an estimated 6.1 percent.

<table>
<thead>
<tr>
<th>Number of hospitals</th>
<th>Average FY 2017 payments/case</th>
<th>Average FY 2018 payments/case</th>
<th>Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>By Geographic Location:</td>
<td></td>
<td></td>
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<tr>
<td>All hospitals</td>
<td>3,292</td>
<td>921</td>
<td>943</td>
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<td>Large urban areas (populations over 1 million)</td>
<td>1,349</td>
<td>1,016</td>
<td>1,044</td>
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<tr>
<td>Other urban areas (populations of 1 million of fewer)</td>
<td>1,142</td>
<td>886</td>
<td>902</td>
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<tr>
<td>Rural areas</td>
<td>801</td>
<td>625</td>
<td>642</td>
</tr>
<tr>
<td>Urban hospitals</td>
<td>2,491</td>
<td>955</td>
<td>977</td>
</tr>
<tr>
<td>0–99 beds</td>
<td>638</td>
<td>769</td>
<td>799</td>
</tr>
<tr>
<td>100–199 beds</td>
<td>765</td>
<td>826</td>
<td>845</td>
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<tr>
<td>200–299 beds</td>
<td>445</td>
<td>877</td>
<td>923</td>
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<tr>
<td>300–499 beds</td>
<td>431</td>
<td>969</td>
<td>993</td>
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<td>500 or more beds</td>
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<td>Rural hospitals</td>
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<td>642</td>
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<tr>
<td>0–49 beds</td>
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<td>522</td>
<td>541</td>
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<tr>
<td>50–99 beds</td>
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<td>585</td>
<td>598</td>
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<tr>
<td>100–149 beds</td>
<td>117</td>
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<tr>
<td>150–199 beds</td>
<td>46</td>
<td>668</td>
<td>689</td>
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### TABLE III—COMPARISON OF TOTAL PAYMENTS PER CASE—Continued

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<tr>
<th>By Region:</th>
<th>Number of hospitals</th>
<th>Average FY 2017 payments/case</th>
<th>Average FY 2018 payments/case</th>
<th>Change</th>
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<tbody>
<tr>
<td>200 or more beds</td>
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<tr>
<td>By Region:</td>
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</tr>
<tr>
<td>Urban by Region</td>
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<tr>
<td>New England</td>
<td>114</td>
<td>1,037</td>
<td>1,052</td>
<td>1.4</td>
</tr>
<tr>
<td>Middle Atlantic</td>
<td>345</td>
<td>1,059</td>
<td>1,082</td>
<td>2.1</td>
</tr>
<tr>
<td>South Atlantic</td>
<td>404</td>
<td>850</td>
<td>867</td>
<td>2.0</td>
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<tr>
<td>East North Central</td>
<td>365</td>
<td>918</td>
<td>937</td>
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<tr>
<td>East South Central</td>
<td>187</td>
<td>801</td>
<td>813</td>
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<tr>
<td>West North Central</td>
<td>160</td>
<td>932</td>
<td>953</td>
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<tr>
<td>West South Central</td>
<td>387</td>
<td>863</td>
<td>895</td>
<td>3.6</td>
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<tr>
<td>Mountain</td>
<td>162</td>
<td>1,005</td>
<td>1,016</td>
<td>1.1</td>
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<tr>
<td>Pacific</td>
<td>375</td>
<td>1,211</td>
<td>1,254</td>
<td>3.5</td>
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<td>Rural by Region</td>
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<tr>
<td>New England</td>
<td>801</td>
<td>625</td>
<td>642</td>
<td>2.6</td>
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<tr>
<td>Rural areas</td>
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<td>861</td>
<td>905</td>
<td>5.0</td>
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<td>Urban by Region</td>
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<td>Teaching Status:</td>
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<td>Non-teaching</td>
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<td>779</td>
<td>799</td>
<td>2.6</td>
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<tr>
<td>Fewer than 100 Residents</td>
<td>835</td>
<td>893</td>
<td>910</td>
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<td>100 or more Residents</td>
<td>246</td>
<td>1,288</td>
<td>1,321</td>
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<tr>
<td>Urban DSH:</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>100 or more beds</td>
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<td></td>
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</tr>
<tr>
<td>Less than 100 beds</td>
<td>1,563</td>
<td>982</td>
<td>1,007</td>
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<tr>
<td>Rural DSH:</td>
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<tr>
<td>Sole Community (SCH/EACH)</td>
<td>259</td>
<td>623</td>
<td>632</td>
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<tr>
<td>Referral Center (RRC/EACH)</td>
<td>271</td>
<td>772</td>
<td>775</td>
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<tr>
<td>Other Rural:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>100 or more beds</td>
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<td></td>
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<tr>
<td>Less than 100 beds</td>
<td>240</td>
<td>507</td>
<td>516</td>
<td>1.8</td>
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<tr>
<td>Urban teaching and DSH:</td>
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<tr>
<td>Both teaching and DSH</td>
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<tr>
<td>Teaching and no DSH</td>
<td>870</td>
<td>1,054</td>
<td>1,082</td>
<td>2.6</td>
</tr>
<tr>
<td>No teaching and DSH</td>
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<td>928</td>
<td>941</td>
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<tr>
<td>No teaching and no DSH</td>
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<td>844</td>
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<td>Rural Hospital Types:</td>
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<td>Non special status hospitals</td>
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<td>833</td>
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<tr>
<td>RRC/EACH</td>
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<td>SCH, RRC and EACH</td>
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<td>716</td>
<td>731</td>
<td>2.0</td>
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<td>Hospitals Reclassified by the Medicare Geographic Classification Review Board:</td>
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<tr>
<td>FY 2018 Reclassifications:</td>
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<tr>
<td>All Urban Reclassified</td>
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<td>956</td>
<td>971</td>
<td>1.6</td>
</tr>
<tr>
<td>All Urban Non-Reclassified</td>
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<td>956</td>
<td>983</td>
<td>2.8</td>
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<tr>
<td>All Rural Reclassified</td>
<td>271</td>
<td>660</td>
<td>675</td>
<td>2.4</td>
</tr>
<tr>
<td>All Rural Non-Reclassified</td>
<td>482</td>
<td>850</td>
<td>855</td>
<td>2.6</td>
</tr>
<tr>
<td>All Section 401 Reclassified Hospitals</td>
<td>148</td>
<td>873</td>
<td>857</td>
<td>-1.8</td>
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<tr>
<td>Other Reclassified Hospitals (Section 1886(d)(8)(B))</td>
<td>42</td>
<td>600</td>
<td>637</td>
<td>6.1</td>
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<tr>
<td>Type of Ownership:</td>
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<tr>
<td>Voluntary</td>
<td>1,914</td>
<td>938</td>
<td>957</td>
<td>2.0</td>
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<tr>
<td>Proprietary</td>
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<td>823</td>
<td>848</td>
<td>3.0</td>
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<tr>
<td>Government</td>
<td>514</td>
<td>960</td>
<td>993</td>
<td>3.4</td>
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<td>Medicare Utilization as a Percent of Inpatient Days:</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>0–25</td>
<td>509</td>
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<td>1,129</td>
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<tr>
<td>25–50</td>
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<td>948</td>
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<td>50–65</td>
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<td>756</td>
<td>772</td>
<td>2.1</td>
</tr>
<tr>
<td>Over 65</td>
<td>135</td>
<td>582</td>
<td>639</td>
<td>9.8</td>
</tr>
</tbody>
</table>

[FY 2017 payments compared to FY 2018 payments]
There are 415 LTCHs included in this impacts analysis, which includes data for 72 nonprofit (voluntary ownership control) LTCHs, 328 proprietary LTCHs, and 15 LTCHs that are government-owned and operated. (We currently approximately 425 LTCHs, for purposes of this impact analysis, we excluded the data of all-inclusive rate providers consistent with the development of the proposed FY 2018 MS–LTC–DRG relative weights (discussed in section VIII.B.3.c. of the preamble of this proposed rule). Moreover, in the claims data use for this proposed rule, 3 of these 415 LTCHs only have claims for site neutral payment rate cases and are therefore not included in our impact analysis for LTCH PPS standard Federal payment rate cases. In the impact analysis, we used the proposed payment rate, factors, and policies presented in this proposed rule, which include the rolling end to transition to the site neutral payment rate required by section 1886(m)(6)(A) of the Act (as described below), the proposed 1.0 percent annual update to the LTCH PPS standard Federal payment rate required by section 411 of Public Law 114–10, the proposed update to the MS–LTC–DRG classification relative weights, the proposed update to the wage index values and labor-related share, the proposed change to the SSO payment methodology (discussed in VII.B, of the preamble of this proposed rule), our proposal to adopt a regulatory delay of the full implementation of the 25-percent threshold policy for FY 2018, and our proposals to implement certain provisions of the 21st Century Cures Act, and the best available claims and CCR data to estimate the proposed change in payments for FY 2018.

Under the dual rate LTCH PPS payment structure, payment for LTCH discharges that meet the criteria for exclusion from the site neutral payment rate (that is, LTCH PPS standard Federal payment rate cases) is based on the LTCH PPS standard Federal payment rate. Consistent with the statute, the site neutral payment rate is the lower of the IPPS comparable per diem amount as determined under §412.529(d)(4), including any applicable outlier payments as specified in §412.525(a); or 100 percent of the estimated cost of the case as determined under existing §412.529(d)(2). In addition, there are two separate HCO targets—one for LTCH PPS standard Federal payment rate cases and one for site neutral payment rate cases. The statute also establishes a transitional payment method for LTCH discharges occurring in cost reporting periods beginning during FY 2016 and FY 2017. For FY 2018, the applicability of this transitional payment method for site neutral payment rate cases is dependent upon both, the discharge date and the start date of the LTCH’s FY 2018 cost reporting period. Specifically, the transitional payment method only applies to those site neutral payment rate cases that occur in cost reporting periods that begin before October 1, 2017. The transitional payment amount for site neutral payment rate cases is a blended payment rate, which is calculated as 50 percent of the applicable site neutral payment rate amount for the discharge as determined under §412.523, while site neutral payment rate cases in cost reporting periods beginning on or after October 1, 2017 are paid the site neutral payment rate amount determined under §412.522(c)(1). Based on the best available data for the 415 LTCHs in our database that were considered in the analyses used for this proposed rule, we estimate that overall LTCH PPS payments in FY 2018 would decrease by approximately 5.2 percent (or approximately $238 million) based on the proposed rates and factors presented in section VIII. of the preamble and section V. of the Addendum to this proposed rule. (We note that this estimate does not reflect our proposal to adopt a 1-year regulatory delay of the full implementation of the 25-percent threshold policy for FY 2018 and, with the exception of changes to the HCO payment policy, does not reflect our proposals regarding the implementation of certain provisions of the 21st Century Cures Act. As discussed in greater detail below, our actuaries estimate these proposals would increase spending by approximately $65 million in FY 2018.) This projection takes into account estimated payments for LTCH cases in our database that met or would have met the patient-level criteria and were paid the LTCH PPS standard Federal payment rate if those criteria had been in effect at the time of the discharge, and estimated payments for LTCH cases that did not meet or would not have met the patient-level criteria and were paid under the site neutral payment rate if that rate had been in effect at the time of the discharge, as described in the following paragraph.

The statutory transitional payment method for cases that are paid the site neutral payment rate for LTCH discharges occurring in cost reporting periods that begin during FY 2016 or FY 2017 uses a blended payment rate, which is determined as 50 percent of the site neutral payment rate amount for the discharge and 50 percent of the standard Federal prospective payment rate amount for the discharge [§412.522(c)(3)]. The transitional blended payment rate uses the same blend percentages (that is, 50 percent) for both years of the 2-year transition period. Therefore, when estimating FY 2017 LTCH PPS payments for site neutral payment rate cases for this impact analysis, the transitional blended payment rate was applied to all such cases because all LTCHs are either in the hospital’s cost reporting period that began during FY 2016 or in the hospital’s cost reporting period that will begin during FY 2017. However, when estimating FY 2018 LTCH PPS payments for site neutral payment rate cases for this impact analysis, because the statute specifies that the site neutral payment rate effective date (and 2-year transitional period) for a given LTCH is based on the date that LTCH’s cost reporting period begins during FY 2018, we included an adjustment to account for this rolling effective date, consistent with the general approach used for the LTCH PPS impact analysis presented in the FY 2016 IPPS/LTCH PPS final rule (80 FR 49831).

For purposes of this analysis, to estimate proposed total FY 2018 LTCH PPS payments for site neutral payment rate cases, we used the same general approach as was used in the FY 2016 IPPS/LTCH PPS final rule with modifications to account for the rolling end date to the transitional site neutral payment rate in FY 2018 instead of the rolling effective date for implementation of the transitional site neutral payment rate in FY 2016. In summary, under this approach, we grouped LTCHs based on the first quarter their cost reporting periods would begin during FY 2018. For example, LTCHs with cost reporting periods that begin during October through December 2017 begin during the first quarter of FY 2018. For LTCHs grouped in each quarter of FY 2018, we modeled those LTCHs’ estimated FY 2018 site neutral payment rate payments under the transitional blended payment rate based on the quarter in which the LTCHs in each group would continue to be paid the transitional payment method for the site neutral payment rate cases.

For purposes of this estimate, then, we assume the cost reporting period is the same for all LTCHs in each of the quarterly groups and that this cost reporting period begins on the first day of that quarter. (For example, our first group consists of 41 LTCHs whose cost reporting period will begin in the first quarter of FY 2018 so that, for purposes of this estimate, we assume all 41 LTCH will begin their FY 2018 cost reporting period on October 1, 2017.) Second, we estimated the proportion of FY 2018 site neutral payment rate cases in each of the quarterly groups, and we then assume this proportion is applicable.
for example, as discussed in more detail below, we estimate the first quarter group will discharge 6.3 percent of all FY 2018 site neutral payment rate cases and therefore, we estimate that group of LTCHs will discharge 6.3 percent of site neutral payment rate cases in each quarter of FY 2018. Then, we modeled estimated FY 2018 payments on a quarterly basis under the LTCH PPS standard Federal payment rate based on the assumptions described above. We continue to believe that this approach is a reasonable means of taking the rolling effective date into account when estimating FY 2018 payments.

Based on the fiscal year begin date information in the December 2016 update of the PSF and the LTCH claims from the December 2016 update of the FY 2016 MedPAR files for the 415 LTCHs in our database used for this proposed rule, we found the following: 6.3 percent of site neutral payment rate cases are from 41 LTCHs whose cost reporting periods will begin during the first quarter of FY 2018; 23.7 percent of site neutral payment rate cases are from 106 LTCHs whose cost reporting periods will begin in the second quarter of FY 2018; 9.3 percent of site neutral payment rate cases are from 55 LTCHs whose cost reporting periods will begin in the third quarter of FY 2018; and 60.7 percent of site neutral payment rate cases are from 213 LTCHs whose cost reporting periods will begin in the fourth quarter of FY 2018.

Therefore, the following percentages apply in the approach described above:

- First Quarter FY 2018: 6.3 percent of site neutral payment rate cases (that is, the percentage of discharges from LTCHs whose FY 2018 cost reporting will begin in the first quarter of FY 2018) are no longer eligible for the transitional payment method while the remaining 93.7 percent of site neutral payment rate discharges are eligible to be paid under the transitional payment method.
- Second Quarter FY 2018: 30.0 percent of site neutral payment rate second quarter discharges (that is, the percentage of discharges from LTCHs whose FY 2018 cost reporting will begin in the first or second quarter of FY 2018) are no longer eligible for the transitional payment method while the remaining 70.0 percent of site neutral payment rate second quarter discharges are eligible to be paid under the transitional payment method.
- Third Quarter FY 2018: 39.3 percent of site neutral payment rate third quarter discharges (that is, the percentage of discharges from LTCHs whose FY 2018 cost reporting will begin in the second, third, or fourth quarter of FY 2018) are no longer eligible for the transitional payment method while the remaining 60.7 percent of site neutral payment rate third quarter discharges are eligible to be paid under the transitional payment method.
- Fourth Quarter FY 2018: 100.0 percent of site neutral payment rate fourth quarter discharges (that is, the percentage of discharges from LTCHs whose FY 2018 cost reporting will begin in the first, second, third, or fourth quarter of FY 2018) are no longer eligible for the transitional payment method so that no site neutral payment rate discharge cases are eligible to be paid under the transitional payment method.

Based on the FY 2016 LTCH cases that were used for the analyses in this proposed rule, approximately 42 percent of those cases were or would have been classified as site neutral payment rate cases if the site neutral payment rate had been in effect at the time of the discharge (that is, 42 percent of LTCH cases did not or would not have met the patient-level criteria for exclusion from the site neutral payment rate cases). Our Office of the Actuary estimates that the percent of LTCH PPS cases that will be paid at the site neutral payment rate in FY 2018 will not change significantly from the historical data. Taking into account the transitional blended payment rate and other changes that would apply to the site neutral payment rate cases in FY 2018, we estimate that aggregate LTCH PPS payments for these site neutral payment rate cases would decrease by approximately 22 percent (or approximately $252 million). Therefore, the LTCH cases are expected to meet the patient-level criteria for exclusion from the site neutral payment rate payment method in FY 2018, and would be based on the proposed LTCH PPS standard Federal payment rate for the full year. We estimate that total LTCH PPS payments for these LTCH PPS standard Federal payment rate cases in FY 2018 would increase approximately 0.4 percent (or approximately $15 million). This estimated increase in LTCH PPS payments for LTCH PPS standard Federal payment rate cases in FY 2018 is primarily due to the estimated effects of the 1.0 percent annual update to the LTCH PPS standard Federal payment rate for FY 2018 required by section 411 of Public Law 114–10 (discussed in section V.A. of the Addendum to this proposed rule) and an estimated decrease in HCO payments for these cases (discussed in section V.D. of the Addendum to this proposed rule). (We note that because our proposed SSO payment methodology discussed in VIII.E. of the preamble of this proposed rule) does not incorporate the factors described as well as the proposed estimated payments for LTCH PPS standard Federal payment rate are applied to the FY 2017 LTCH PPS standard Federal payment rate set forth under § 412.523(c)(3)(xiv) (that is, $42,476.41).

Table IV shows the estimated impact for LTCH PPS standard Federal payment rate cases. The estimated change attributable solely to the annual update of 1.0 to the LTCH PPS standard Federal payment rate is projected to result in an increase in 0.9 percent in payments per discharge for LTCH PPS standard Federal payment rate cases from FY 2017 to FY 2018, on average, for all LTCHs (Column 6). In addition to the proposed annual update to the LTCH PPS standard Federal payment rate for FY 2018, the estimated increase of 0.9 percent shown in Column 6 of Table IV also includes the estimated payments for SSO cases that would be paid using special methodologies that are not affected by the proposed annual update to the LTCH PPS standard Federal payment rate (without incorporating our proposed SSO payment methodology as discussed in VIII.E. of the preamble of this proposed rule),
as well as the proposed reduction that is applied to the annual update of LTCHs that do not submit the required LTCH QRP data. Therefore, for all hospital categories, the projected increase in payments based on the proposed LTCH PPS standard Federal payment rate cases for FY 2018 required under section 411 of Public Law 114–10.

For FY 2018, we are proposing to update the wage index values based on the most recent available data, and we are proposing to continue to use labor market areas based on the OMB CBASA delineations (as discussed in section V.B. of the Addendum to this proposed rule). In addition, we are proposing to reduce the labor-related share from 66.5 percent to 66.3 percent under the LTCH PPS for FY 2018, based on the most recent available data on the relative importance of the labor-related share of operating and capital costs of the 2013-based LTCH market basket. We are proposing to apply a proposed area wage level budget neutrality factor of 1.000077 to ensure that the proposed changes to the wage data and labor-related share do not result in a change in estimated aggregate LTCH PPS payments to LTCH PPS standard Federal payment rate cases.

As we discuss in VIII.E. of the preamble of this proposed rule, we are proposing to simplify our SSO payment methodology in order to alleviate potential incentives to improperly hold patients beyond the SSO threshold. We also note that we do not believe aggregate payments to LTCHs should increase or decrease as a result of our policy, and thus, we are proposing to apply a proposed budget neutrality factor of 0.9672 to ensure the proposed changes to the SSO payment methodology does not result in a change in estimated aggregate LTCH PPS payments to LTCH PPS standard Federal payment rate cases.

We currently estimate total HCO payments for LTCH PPS standard Federal payment rate cases would increase from FY 2017 to FY 2018. Based on the FY 2016 LTCH cases that were used for the analyses in this proposed rule, we estimate that the FY 2017 HCO threshold of $21,943 (as established in the FY 2017 IPPS/LTCPPS final rule) would result in estimated HCO payments for LTCH PPS standard Federal payment rate cases in FY 2017 that are above the estimated 8 percent target. Specifically, we currently estimate that HCO payments for LTCH PPS standard Federal payment rate cases would be approximately 8.6 percent of the estimated total LTCH PPS standard Federal payment rate payments in FY 2017. Combined with our estimate that proposed FY 2018 HCO payments for LTCH PPS standard Federal payment rate cases would be 7.975 percent of estimated total LTCH PPS standard Federal payment rate cases, we estimate that the proposed FY 2018 HCO payments for LTCH PPS standard Federal payment rate cases would be approximately $15 million in FY 2018. Specifically, the proposed change to the site neutral payment rate formula (§ 412.529) provides for an estimated aggregate increase in Medicare spending for FY 2018 of $10 million. The remaining estimated increase of $5 million in Medicare spending comes from the temporary exception to the site neutral payment rate for certain spinal cord hospitals provided for under section 15009 (as discussed in section VIII.E. of the preamble of this proposed rule). Our actuaries estimate the remaining provisions of the 21st Century Cures Act applicable to LTCHs (that is, sections 15007, 15008, and 15010, discussed in sections VIII.I., VIII.J., and VIII.F., respectively, of the preamble of this proposed rule) will have negligible impact on aggregate Medicare spending in FY 2018. (We note that section 15007, which provides for an additional delay in the full implementation of the 25-percent threshold policy (discussed in VIII.G. of the preamble of this proposed rule), does not impact FY 2018 LTCH PPS payments.) In addition, if adopted, our actuaries estimate that our proposal to further delay the full implementation of the 25-percent threshold policy for FY 2018 would increase aggregate Medicare spending by $50 million.

As discussed in section VIII.E. of the preamble of this proposed rule, section 15009 of the 21st Century Cures Act provides for a temporary exception to the site neutral payment rate for certain spinal cord specialty hospitals for discharges occurring in cost reporting periods beginning during FY 2018 and FY 2019. To qualify for this temporary exception, an LTCH must, among other things, meet the “significant out-of-state admissions criterion” at section 1886(m)(6)(F)(iii) of the Act. The statute further provides authority for the Secretary to implement the significant out-of-state admissions criterion at section 1886(m)(6)(F)(iii) of the Act by program instruction or otherwise, and exempts the policy initiatives from any information collection requirements under the Paperwork Reduction Act. Although exempt from these information collection requirements, we estimate that each application will require 2.5 hours of work from each LTCH (to review the billing addresses of the hospital’s Medicare and non-Medicare inpatients). This information will be collected on a one-time basis. Based on the best information available to CMS, we estimate that only two hospitals meet the other requirements for this exception. Therefore, we estimate that the total number of hours associated with this request will be 5 (2.5 hours per hospital for 2 hospitals). We estimate a current, average salary of $29 per hour, which provides for fringe benefits ($58 per hour). Therefore, we estimate the total costs associated with this information collection will be $290 (5 hours at $58 per hour).

4. Anticipated Effects of Proposed LTCH PPS Payment Rate Changes and Policy Changes
   a. Budgetary Impact

Section 123(a)(1) of the BBRA requires that the PPS developed for LTCHs “maintain budget neutrality.” We believe that the statute’s mandate for budget neutrality applies only to the first year of the implementation of the LTCH PPS (that is, FY 2005). Therefore, in calculating the FY 2003
section I.J. of this Appendix.)

As discussed in more detail in section I.J. of this Appendix, some LTCH discharges in FY 2018 will still be eligible to be paid based on the blended payment.
COLA for LTCHs located in Alaska and Hawaii. Specifically, for modeling FY 2017 LTCH PPS payments, we used the current FY 2017 labor-related share (66.5 percent); the wage index values established in the Tables 12A and 12B listed in the Addendum to the FY 2017 IPPS/LTCH PPS final rule (which are available via the Internet on the CMS Web site); the FY 2017 HCO fixed-loss amount for LTCH PPS standard Federal payment rate cases of $21,943 (as discussed in section V.D. of the Addendum to that final rule) and the FY 2017 COLA factors (shown in the table in section V.C. of the Addendum to that final rule) to adjust the FY 2017 nonlabor-related share (33.5 percent) for LTCHs located in Alaska and Hawaii. Similarly, for modeling proposed FY 2018 LTCH PPS payments, we used the proposed FY 2018 LTCH PPS labor-related share (66.3 percent), the proposed FY 2018 wage index values from Tables 12A and 12B listed in section VI. of the Addendum to this proposed rule (which are available via the Internet on the CMS Web site), the proposed FY 2018 fixed-loss amount for LTCH PPS standard Federal payment rate cases of $30,081 (as discussed in section V.D.3 of the Addendum to this proposed rule), and the proposed FY 2018 COLA factors (shown in the table in section V.C. of the Addendum to this proposed rule) to adjust the FY 2018 nonlabor-related share (33.7 percent) for LTCHs located in Alaska and Hawaii.

As previously discussed, our impact analysis reflects an estimated change in payments for SSO cases (including our proposed changes to the SSO payment methodology), as well as an estimated decrease in HCO payments for LTCH PPS standard Federal payment rate cases (as described previously in section I.J.1. of this Appendix). In modeling payments for SSO cases prior to accounting for our proposed SSO payment methodology and for HCO cases for LTCH PPS standard Federal payment rate cases, we applied a proposed inflation factor of 5.6 percent (determined by the Office of the Actuary) to update the 2016 costs of each case.

The impacts that follow reflect the estimated “losses” or “gains” among the various classifications of LTCHs from FY 2017 to FY 2018 based on the proposed payment rules and proposed policy changes applicable to LTCH PPS standard Federal payment rate cases presented in this proposed rule. Table IV illustrates the estimated aggregate impact of the proposed change in LTCH PPS payments for LTCH PPS standard Federal payment rate cases among various classifications of LTCHs. (As discussed previously, these impacts do not include LTCH PPS site neutral payment rate cases.)

- The first column, LTCH Classification, identifies the type of LTCH.
- The second column lists the number of LTCHs of each classification type.
- The third column identifies the number of LTCH cases expected to meet the LTCH PPS standard Federal payment rate criteria.
- The fourth column shows the estimated FY 2017 payment per discharge for LTCH cases expected to meet the LTCH PPS standard Federal payment rate criteria (as described previously).
- The fifth column shows the estimated FY 2018 payment per discharge for LTCH cases expected to meet the LTCH PPS standard Federal payment rate criteria (as described previously).
- The sixth column shows the proposed percent change in estimated payments per discharge for LTCH PPS standard Federal payment rate cases from FY 2017 to FY 2018 due to the proposed annual update to the standard Federal rate (as discussed in section V.A.2. of the Addendum to this proposed rule).
- The seventh column shows the percentage change in estimated payments per discharge for LTCH PPS standard Federal payment rate cases from FY 2017 to FY 2018 for proposed changes to the area wage level adjustment (that is, the wage indexes and the labor-related share), including the application of the proposed area wage level budget neutrality factor (as discussed in section V.B. of the Addendum to this proposed rule).
- The eighth column shows the percentage change in estimated payments per discharge for LTCH PPS standard Federal payment rate cases for changes resulting from our proposed SSO payment methodology and associated budget neutral adjustment to the LTCH PPS standard Federal payment rate (column 7).
- The ninth column shows the percentage change in estimated payments per discharge for LTCH PPS standard Federal payment rate cases from FY 2017 (Column 4) to FY 2018 (Column 5) for all proposed changes (and includes the effect of estimated changes to HCO and SSO payments).

**Table IV—Impact of Proposed Payment Rate and Proposed Policy Changes to LTCH PPS Payments for Standard Payment Rate Cases for FY 2018**

<table>
<thead>
<tr>
<th>LTCH classification</th>
<th>Number of LTCHs</th>
<th>Number of LTCHs standard payment rate cases</th>
<th>Average FY 2017 LTCH PPS payment per standard payment rate</th>
<th>Average FY 2018 LTCH PPS payment per standard payment rate</th>
<th>Proposed percent change due to change to proposed annual update to the standard federal rate</th>
<th>Proposed percent change due to proposed changes to area wage adjustment with wage budget neutrality</th>
<th>Proposed percent change due to proposed changes to the short stay outlier payment methodology change</th>
<th>Proposed percent change due to all proposed standard payment rate changes</th>
</tr>
</thead>
<tbody>
<tr>
<td>All Providers</td>
<td>415</td>
<td>73,231</td>
<td>$46,947</td>
<td>$47,149</td>
<td>0.9</td>
<td>0.0</td>
<td>0.0</td>
<td>0.4</td>
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<td>By Location:</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Rural</td>
<td>21</td>
<td>2,214</td>
<td>37,951</td>
<td>37,702</td>
<td>0.9</td>
<td>-0.3</td>
<td>0.1</td>
<td>-0.7</td>
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<tr>
<td>Urban</td>
<td>394</td>
<td>71,017</td>
<td>47,227</td>
<td>47,443</td>
<td>0.9</td>
<td>0.0</td>
<td>0.0</td>
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<tr>
<td>Large</td>
<td>200</td>
<td>40,843</td>
<td>49,951</td>
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<tr>
<td>Other</td>
<td>194</td>
<td>131,177</td>
<td>43,541</td>
<td>43,645</td>
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<td>-0.1</td>
<td>-0.2</td>
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<td>By Participation Date:</td>
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<td></td>
<td></td>
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<tr>
<td>Before Oct. 1983</td>
<td>16</td>
<td>2,509</td>
<td>42,228</td>
<td>43,135</td>
<td>0.9</td>
<td>-0.5</td>
<td>2.0</td>
<td>2.1</td>
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<tr>
<td>Oct. 1983–Sept. 1993</td>
<td>45</td>
<td>9,580</td>
<td>52,603</td>
<td>52,668</td>
<td>0.8</td>
<td>-0.1</td>
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<td>Oct. 1993–Sept. 2002</td>
<td>169</td>
<td>30,469</td>
<td>45,835</td>
<td>46,061</td>
<td>0.9</td>
<td>0.0</td>
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<td>After October 2002</td>
<td>185</td>
<td>30,673</td>
<td>46,671</td>
<td>46,834</td>
<td>0.9</td>
<td>0.0</td>
<td>0.0</td>
<td>0.3</td>
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<td>By Ownership Type:</td>
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<tr>
<td>Voluntary</td>
<td>72</td>
<td>9,536</td>
<td>49,476</td>
<td>49,458</td>
<td>0.9</td>
<td>-0.1</td>
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<td>Proprietary</td>
<td>328</td>
<td>62,236</td>
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<td>54,034</td>
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<td>By Region:</td>
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<tr>
<td>New England</td>
<td>12</td>
<td>2,748</td>
<td>44,003</td>
<td>44,457</td>
<td>0.9</td>
<td>-0.3</td>
<td>0.3</td>
<td>1.0</td>
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<td>Middle Atlantic</td>
<td>25</td>
<td>5,845</td>
<td>51,781</td>
<td>52,133</td>
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<td>-0.2</td>
<td>0.4</td>
<td>0.7</td>
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<tr>
<td>South Atlantic</td>
<td>66</td>
<td>13,245</td>
<td>46,739</td>
<td>47,089</td>
<td>0.9</td>
<td>-0.1</td>
<td>0.4</td>
<td>0.7</td>
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<tr>
<td>East North Central</td>
<td>68</td>
<td>11,419</td>
<td>46,589</td>
<td>46,717</td>
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<td>0.0</td>
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<tr>
<td>East South Central</td>
<td>34</td>
<td>5,209</td>
<td>43,878</td>
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<tr>
<td>West North Central</td>
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<td>4,325</td>
<td>46,735</td>
<td>45,380</td>
<td>0.9</td>
<td>0.2</td>
<td>-1.2</td>
<td>-0.8</td>
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</table>
TABLE IV—IMPACT OF PROPOSED PAYMENT RATE AND PROPOSED POLICY CHANGES TO LTCH PPS PAYMENTS FOR STANDARD PAYMENT RATE CASES FOR FY 2018—Continued

<table>
<thead>
<tr>
<th>LTCH classification</th>
<th>Number of LTCHs</th>
<th>Number of LTCH PPS standard payment rate cases</th>
<th>Average FY 2017 LTCH PPS payment per standard payment rate</th>
<th>Average proposed FY 2018 LTCH PPS payment per standard payment rate</th>
<th>Proposed payment rate due to proposed annual update to the LTCH PPS standard Federal payment rate</th>
<th>Proposed payment rate due to proposed policy changes to the short stay outlier payment methodology change</th>
<th>Proposed payment rate due to all proposed policy changes to the LTCH PPS standard Federal payment rate</th>
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<tbody>
<tr>
<td>West South Central</td>
<td>127</td>
<td>18,398</td>
<td>41,960</td>
<td>41,299</td>
<td>0.8</td>
<td>0.2</td>
<td>0.0</td>
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<tr>
<td>Mountain</td>
<td>31</td>
<td>4,184</td>
<td>49,112</td>
<td>49,256</td>
<td>0.0</td>
<td>0.2</td>
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<td>Pacific</td>
<td>25</td>
<td>7,858</td>
<td>58,479</td>
<td>59,128</td>
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<td>47,363</td>
<td>0.8</td>
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1 Estimated FY 2018 LTCH PPS payments for LTCH PPS standard Federal payment rate criteria based on the proposed payment rate and factor changes applicable to such cases presented in the preamble of and the Addendum to this proposed rule.

2 Percent change in estimated payments per discharge for LTCH PPS standard Federal payment rate cases from FY 2017 to FY 2018 for the proposed annual update to the LTCH PPS standard Federal payment rate.

3 Percent change in estimated payments per discharge for LTCH PPS standard Federal payment rate cases from FY 2017 to FY 2018 for proposed changes to the area wage level factor under §412.522(c) (as discussed in section V.B. of the Addendum to this proposed rule).

4 Percent change in estimated payments per discharge for LTCH PPS standard Federal payment rate cases from FY 2017 to FY 2018 for proposed change to the SSO payment methodology.

5 Percent change in estimated payments per discharge for LTCH PPS standard Federal payment rate cases from FY 2017 to FY 2018 (shown in Column 4) to FY 2018 (shown in Column 5), including all of the proposed changes to the rates and factors applicable to such cases presented in the preamble and the Addendum to this proposed rule. We note that this column, which shows the proposed percent change in estimated payments per discharge for all proposed changes, does not equal the sum of the proposed percent changes in estimated payments per discharge for the proposed annual update to the LTCH PPS standard Federal payment rate (Column 6) and the proposed changes to the area wage level adjustment with budget neutrality (Column 7) due to the effect of estimated changes in both estimated payments to SOS cases (prior to accounting for the proposed change to the SSO payment methodology) and aggregate HCO payments for LTCH PPS standard Federal payment rate cases (as discussed in this impact analysis), as well as other interactive effects that cannot be isolated.

d. Results

Based on the FY 2016 LTCH cases (from 415 LTCHs) that were used for the analyses in this proposed rule, we have prepared the following summary of the impact (as shown in Table IV) of the proposed LTCH PPS payment rate and proposed policy changes for LTCH PPS standard Federal payment rate cases presented in this proposed rule. The impact analysis in Table IV shows that estimated payments per discharge for LTCH PPS standard Federal payment rate cases are projected to increase 0.4 percent, on average, for all LTCHs from FY 2017 to FY 2018 as a result of the proposed payment rate and proposed policy changes applicable to LTCH PPS standard Federal payment rate cases presented in this proposed rule. This estimated 0.4 percent increase in LTCH PPS payments per discharge was determined by comparing estimated FY 2018 LTCH PPS payments (using the proposed payment rates and factors discussed in this proposed rule) to estimated FY 2017 LTCH PPS payments for LTCH discharges which will be LTCH PPS standard Federal payment rate cases if the dual rate LTCH PPS payment structure was or had been in effect at the time of the discharge (as described in section 13.4. of this Appendix).

As stated previously, we are proposing to update the LTCH PPS standard Federal payment rate for FY 2018 by 1.0 percent as required by statute. For LTCHs that fail to submit quality data under the requirements of the LTCH QRP, as required by section 1886(m)(5)(C) of the Act, a 2.0 percentage point reduction is applied to the annual update to the LTCH PPS standard Federal payment rate. Consistent with §412.523(d)(4), we are also proposing to apply an area wage level budget neutrality factor to the proposed FY 2018 LTCH PPS standard Federal payment rate of 1.000077, based on the best available data at this time, to ensure that any proposed changes to the area wage level adjustment (that is, the proposed annual update of the wage index values and labor-related share) would not result in any change (increase or decrease) in estimated aggregate LTCH PPS standard Federal payment rate payments. Finally, we are proposing a budget neutrality adjustment of 0.9672 for our proposed changes to the SSO payment methodology (discussed in VIII.E.2.d. of the preamble of this proposed rule). As we also explained earlier in this section, for most categories of LTCHs (as shown in Table IV, Column 7), the estimated payment increase due to the proposed 1.0 percent annual update to the LTCH PPS standard Federal payment rate is projected to result in approximately a 0.9 percent increase in estimated payments per discharge for LTCH PPS standard Federal payment rate cases for all LTCHs from FY 2017 to FY 2018. This is because our estimate of the proposed changes in payments due to the proposed update to the LTCH PPS standard Federal payment rate also reflects estimated payments for SOS cases that are paid using special methodologies that are not affected by the update to the LTCH PPS standard Federal payment rate (prior to accounting for the proposed change to the SSO payment methodology). Consequently, for certain hospital categories, we estimate that payments to LTCH PPS standard Federal payment rate cases may increase by less than 1.0 percent due to the proposed annual update to the LTCH PPS standard Federal payment rate for FY 2018.

(1) Location

Based on the most recent available data, the vast majority of LTCHs are located in urban areas. Only approximately 5 percent of the LTCHs are identified as being located in a rural area, and approximately 3 percent of all LTCHs are rural LTCHs. Urban LTCHs are expected to be treated in these rural hospitals. The impact analysis presented in Table IV shows that the proposed average percent increase in estimated payments per discharge for LTCH PPS standard Federal payment rate cases from FY 2017 to FY 2018 is 0.4 percent. However, for rural LTCHs, the proposed overall percent change for LTCH PPS standard Federal payment rate cases is estimated to be a 0.7 percent decrease. This projected decrease is primarily driven by a projected decrease resulting from changes to the proposed changes to the LTCH PPS standard Federal payment rate, which is 0.7 percent. For urban LTCHs, we estimate an increase of 0.5 percent. The proposed changes to the short stay outlier payment methodology are not expected to affect the estimated payment rates for rural LTCHs.
percent from FY 2017 to FY 2018. Among the urban LTCHs, large urban LTCHs are projected to experience an increase of 0.6 percent in estimated payments per discharge for LTCH PPS standard Federal payment rate cases from FY 2017 to FY 2018, and the rural LTCHs are projected to experience an increase of 0.2 percent in estimated payments per discharge for LTCH PPS standard Federal payment rate cases from FY 2017 to FY 2018, as shown in Table IV.

(2) Participation Date

LTCHs are grouped by participation date into four categories: (1) Before October 1983; (2) between October 1983 and September 1993; (3) between October 1993 and September 2002; and (4) October 2002 and after. Based on the most recent available data, the categories of LTCHs with the largest expected percentage of LTCH PPS standard Federal payment rate cases (approximately 42 percent) are in LTCHs that began participating in the Medicare program after September 2002, and they are projected to experience a 0.3 percent increase in estimated payments per discharge for LTCH PPS standard Federal payment rate cases from FY 2017 to FY 2018, as shown in Table IV.

Approximately 4 percent of LTCHs began participating in the Medicare program before October 1983, and these LTCHs are projected to experience an average percent increase of 2.1 percent in estimated payments per discharge for LTCH PPS standard Federal payment rate cases from FY 2017 to FY 2018, as shown in Table IV with a large portion of this increase among this small group of LTCHs to be a projected 2.0 percent increase resulting from our proposed SSO payment method. Approximately 11 percent of LTCHs began participating in the Medicare program between October 1983 and September 1993, and these LTCHs are projected to experience an average percent increase in estimated payments for LTCH PPS standard Federal payment rate cases from FY 2017 to FY 2018. LTCHs that began participating in the Medicare program between October 1993 and October 1, 2002, which treat approximately 42 percent of all LTCH PPS standard Federal payment rate cases, are projected to experience a 0.5 percent increase in estimated payments from FY 2017 to FY 2018. Lastly, LTCHs that began participating in Medicare program after October, 2002 also treat approximately 42 percent of all LTCH PPS standard Federal payment rate cases and are projected to experience a 0.3 percent increase in estimated payments from FY 2017 to FY 2018.

(3) Ownership Control

LTCHs are grouped into four categories based on ownership control type: Voluntary, proprietary, government and unknown. Based on the most recent available data, approximately 79 percent of LTCHs are identified as voluntary (Table IV). The majority (approximately 79 percent) of LTCHs are identified as proprietary, while government owned and operated LTCHs represent approximately 4 percent of LTCHs. Based on ownership type, voluntary LTCHs are expected to experience no change in payments to LTCH PPS standard Federal payment rate cases, while proprietary LTCHs are expected to experience an average increase of 0.5 percent in payments to LTCH PPS standard Federal payment rate cases.

Government owned and operated LTCHs, meaning LTCHs that experience a 1.0 percent decrease in payments to LTCH PPS standard Federal payment rate cases from FY 2017 to FY 2018.

(4) Census Region

Estimated payments per discharge for LTCH PPS standard Federal payment rate cases for FY 2017 are projected to experience a decrease from FY 2017 for LTCHs located in the West South Central and West North Central regions, while LTCHs located in all other regions are projected to experience an increase in estimated payments per discharge in comparison to FY 2017. Of the 9 census regions, we project that the increase in estimated payments per discharge to LTCH PPS standard Federal payment rate cases would have the largest positive impact on LTCHs in the Pacific and New England regions (1.1 percent and 1.0 percent, respectively, as shown in Table IV), which is largely attributable to the proposed changes in the proposed SSO payment method. In contrast, LTCHs located in the East North Central and Mountain regions are projected to experience the smallest increase in estimated payments per discharge for LTCH PPS standard Federal payment rate cases from FY 2017 to FY 2018.

(5) Bed Size

LTCHs are grouped into six categories based on bed size: 0–24 beds; 25–49 beds; 50–74 beds; 75–124 beds; 125–199 beds; and greater than 200 beds. We project that LTCHs with 0–24 beds would experience a decrease in payments for LTCH PPS standard Federal payment rate cases of 0.3 percent, and LTCHs with 25–49 beds would experience an increase in payments for LTCH PPS standard Federal payment rate cases of 0.3 percent. LTCHs with 50–74 beds would experience an increase in payments for LTCH PPS standard Federal payment rate cases of 0.1 percent. We project the largest increases in payments to occur in LTCHs with at least 75 beds. In particular, we project LTCHs with 75–124 beds would experience an increase in payments for LTCH PPS standard Federal payment rate cases of 0.8 percent while LTCHs with 125–199 beds would experience an increase in payments for LTCH PPS standard Federal payment rate cases of 0.4 percent. Finally, LTCHs with 200 or more beds would experience the largest increase in payments for LTCH PPS standard Federal payment rate cases of 1.6 percent mostly due to estimated increase in payments from proposed changes to the FY 2018 MS–LTCH–DRG classifications and relative weights and our proposed SSO payment method.

4. Effect on the Medicare Program

As stated previously, we project that the provisions of this proposed rule would result in an increase in estimated aggregate LTCH PPS payments to LTCH PPS standard Federal payment rate cases in FY 2018 relative to FY 2017 of approximately $15 million (or approximately 0.4 percent) for the 415 LTCHs in our database. Although, as stated previously, the hospital-level impacts do not include LTCH PPS site neutral payment rate cases, we estimate that the provisions of this proposed rule would result in a decrease in estimated aggregate LTCH PPS payments to site neutral payment rate cases in FY 2018 relative to FY 2017 of approximately $252 million (or approximately 22 percent) for the 415 LTCHs in our database. Therefore, we project that the provisions of this proposed rule would result in a decrease in estimated aggregate LTCH PPS payments to site neutral payment rate cases in FY 2018 relative to FY 2017 of approximately $238 million (or approximately 5.2 percent) for the 415 LTCHs in our database. Furthermore, as stated previously, our Office of the Actuary estimates an additional estimated increase in aggregate FY 2018 LTCH PPS payments of approximately $65 million for our proposal to delay full implementation of the 25-percent threshold policy for FY 2018 and our proposed implementation of certain provisions of the 21st Century Cures Act. Therefore, in total, we project an overall decrease in LTCH PPS payments of approximately $173 million ($238 million decrease + $65 million increase) or approximately a 3.75 percent decrease in LTCH PPS payments in FY 2018 as compared to FY 2017.

5. Effect on Medicare Beneficiaries

Under the LTCH PPS, hospitals receive payment based on the average resources consumed by patients for each diagnosis. We do not expect any changes in the quality of care or access to services for Medicare beneficiaries as a result of this proposed rule, but we continue to expect that paying prospectively for LTCH services will enhance the efficiency of the Medicare program.

K. Effects of Proposed Requirements for the Hospital Inpatient Quality Reporting (IQR) Program

1. Background

In section IX.A. of the preamble of this proposed rule, we discuss our requirements for hospitals to report quality data under the Hospital IQR Program in order to receive the full annual percentage increase for the FY 2020 payment determination.

In this proposed rule, we are proposing to: (1) Update the electronic clinical quality measure (eCQM) reporting requirements with regard to the number of eCQMs and quarters of data for the FY 2019 and FY 2020 payment determinations; (2) update the eCQM certification requirements for the FY 2019 and FY 2020 payment determinations; (3) update our previously finalized eCQM validation processes for the FY 2020 payment determination and subsequent years; (4) allow hospitals to use an educational review process to correct incorrect validation results for the first three quarters of validation for chart-abstracted measures beginning with the FY 2020 payment determination and for subsequent years; (5) begin voluntary reporting on the new Hybrid Hospital-Wide 30-Day Readmission measure for the CY 2018 reporting period; (6) refine the Hospital Consumer Assessment of Healthcare Providers and Systems (HCAHPS) survey
measure to replace the questions on pain management for the FY 2020 payment determination and subsequent years; (7) refine the Hospital 30-Day Mortality Following Acute Ischemic Stroke Hospitalization measure to include NIH stroke scale for the FY 2023 payment determination and subsequent years; (8) provide confidential reports of measure data stratified by dual eligible status for the Hospital 30-day, All-Cause, Risk-Standardized Readmission Rate Following Pneumonia Hospitalization and Hospital 30-day, All-Cause, Risk Standardized Mortality Rate (RSRR) for Pneumonia measures; and (9) update the Extraordinary Circumstances Exceptions (ECE) Policy for the FY 2020 payment determination and subsequent years.

As further explained in section XIII.B.6. of the preamble of this proposed rule, we believe that there will be an overall decrease in burden for hospitals due to the proposals discussed above. We refer readers to section XIII.B.6. of the preamble of this proposed rule for a summary of our burden estimates.

2. Impact of the Proposed Updates to the eCQM Reporting Requirements

(a) Impact for the CY 2017 Reporting Period/ FY 2019 Payment Determination

In the FY 2017 IPPS/LTC PPS final rule, we finalized policies to require hospitals to submit a full year (four quarters) of data (81 FR 57159) for at least eight eCQMs (81 FR 57157) for both the FY 2019 and FY 2020 payment determinations. In section IX.A.8. of the preamble of this proposed rule, we are proposing the following changes to this finalized policy: (1) Revise the CY 2017 reporting period/FY 2019 payment determination eCQM reporting requirements, such that hospitals are required to report six eCQMs and to submit two, self-selected, calendar quarters of data; and (2) revise the CY 2018 reporting period/FY 2020 payment determination eCQM reporting requirements such that hospitals are required to report six eCQMs for the first three quarters of CY 2018. As described in section XIII.B.6.b. of the preamble of this proposed rule, we believe that the reduction in the required number of eCQMs for the CY 2017 reporting period/FY 2019 payment determination will result in a reduction of 200 minutes per hospital per year, or 3 hours and 20 minutes per hospital per year, for the FY 2019 payment determination.

In total, for the FY 2019 payment determination, we expect our proposal to require hospitals to report data on six eCQMs for two quarters (as compared to our previously finalized requirements to report data on eight eCQMs for four quarters) to represent an annual burden reduction of 11,000 hours across all 3,300 IPPS hospitals participating in the Hospital IQR Program. Using the wage estimate described in section XIII.B.6.a. of the preamble of this proposed rule, we expect this to represent a cost reduction of $361,240 across all 3,300 IPPS hospitals participating in the Hospital IQR Program.

(b) Impact for the CY 2017 Reporting Period/ FY 2019 Payment Determination

Using the same estimate as described above of 10 minutes per record per quarter, we note that if our proposed updates to the CY 2018 reporting period/FY 2020 payment determination are finalized as proposed, we anticipate our proposal to require: (1) Reporting on six of the available eCQMs; and (2) submission of the first three quarters of CY 2018 eCQM data, will result in a burden reduction of 2 hours and 20 minutes (140 minutes) per hospital for the FY 2020 payment determination as compared to the previously finalized requirements to report eight eCQMs for four quarters for the FY 2020 payment determination (81 FR 57157 through 57159). In total, this would represent an annual burden reduction of 7,700 hours across all 3,300 IPPS hospitals participating in the Hospital IQR Program and a cost reduction of $252,868 ($32.84 hourly wage × 7,700 annual hours reduction) across all 3,300 IPPS hospitals.

3. Impact of the Proposed Modifications to eCQM Certification Requirements for the FY 2019 and FY 2020 Payment Determinations and Subsequent Years

In section IX.10.d. of the preamble of this proposed rule, we discuss our proposal to make changes to the Hospital IQR Program eCQM submission requirements to align with the Medicare EHR Incentive Program for eligible hospitals and CAHs. Specifically, for the CY 2017 reporting period/FY 2019 payment determination, we are proposing that: (1) A hospital using EHR technology certified to the 2014 or 2015 Edition, but for which such EHR technology is not certified to all available eCQMs, would be required to have its EHR technology certified to all eCQMs that are available to report; and (2) EHR technology that is certified to all available eCQMs does not need to be recertified each time it is updated to a more recent version of the eCQM specifications. For the CY 2018 reporting period/FY 2020 payment determination, we are proposing that: (1) A hospital using EHR technology certified to the 2015 Edition of such EHR technology is not certified to all available eCQMs, would be required to have its EHR technology certified to all of the eCQMs that are available to report; and (2) an EHR certified for all available eCQMs under the 2015 Edition of CHERT would not need to be recertified each time it is updated to a more recent version of the eCQM specifications. Further, we are proposing that: (1) For the CY 2017 reporting period/FY 2019 payment determination, hospitals would be required to use the most recent version of the eCQM electronic specifications; Spring 2016 version of the eCQM specifications and any applicable addenda; and (2) for the CY 2018 reporting period/FY 2020 payment determination, hospitals be required to use the most recent version of the eCQM electronic specifications; Spring 2017 version of the eCQM specifications and any applicable addenda. Because the use of certified EHR technology is already required for the Medicare EHR Incentive Program for eligible hospitals and CAHs, we believe that these proposals will have no effect on burden for hospitals under the Hospital IQR Program.

4. Impact of the Proposed Modifications to the Existing Validation Processes for the FY 2020 Payment Determination and Subsequent Years

(a) Impact of the Proposed Modifications to the Validation of eCQM Data for the FY 2020 Payment Determination and Subsequent Years

In section IX.A.11. of the preamble of this proposed rule, we discuss our proposal to adopt a modification to the existing eCQM data validation process for the Hospital IQR Program data beginning with validation for the FY 2020 payment determination. First, we are proposing to require eight cases to be submitted per quarter for eCQM validation for the FY 2020 payment determination and subsequent years. We are making this proposal in conjunction with our proposal to require two quarters of data for the CY 2017 eCQM reporting period and our proposal to require three quarters of data for the CY 2018 eCQM reporting period. Accordingly, if those eCQM reporting proposals are finalized, we are proposing that the number of required case files for validation would be 16 records (eight cases per quarter over two quarters) for the FY 2020 payment determination and subsequent years. We are proposing that the number of required case files for validation would be 24 records (eight cases per quarter over three quarters) for the FY 2021 payment determination. Second, we are proposing to add additional exclusion criteria to our hospital and case selection process for eCQM validation for the CY 2018 and subsequent years. We are proposing to extend to the FY 2021 payment determination our previously finalized medical record submission policy for eCQM validation, as finalized in the FY 2017 IPPS/LTC PPS final rule (81 FR 57181), requiring submission of at least 75 percent of sampled eCQM measure medical records in a timely and complete manner. We are proposing to extend to the FY 2021 payment determination our previously finalized medical record submission policy for eCQM validation, as finalized in the FY 2017 IPPS/LTC PPS final rule (81 FR 57181), that the accuracy of eCQM data submitted for validation would not affect a hospital’s validation score. We are also proposing to clarify our finalized policy.

We believe the updates to the exclusions and maintaining previously finalized medical record submission requirements will have no effect on burden for hospitals. We believe that the changes associated with the proposed eCQM validation process will result in a burden reduction of approximately 4,333 hours across up to 200 hospitals selected for eCQM validation. Using the estimated hourly labor cost of $32.84, we estimate an annual cost reduction of $142,296 ($32.84 × 8,981.24 per year) across the 200 hospitals selected for eCQM validation due to our proposal to decrease the number of records collected for validation from 32 records to 16 records for the FY 2020 payment determination. We refer readers to section XIII.B.6.d.(1) of the preamble of this proposed rule for more detail on these calculations.
(b) Impact of the Proposed Modifications to the Validation of eCQM Data for the FY 2021 Payment Determination and Subsequent Years

Applying the time per individual submission of 1 hour and 20 minutes (or 80 minutes) per record for the 24 records we are proposing that hospitals submit for eCQM validation for the FY 2021 payment determination, we estimate a burden reduction of approximately 2,133 hours across up to 200 hospitals selected for eCQM validation for the FY 2021 payment determination. Using the estimated hourly labor cost of $32.84, we estimate an annual cost reduction of $70,048 (2,133 hours × $32.84 per hour) across the 200 hospitals selected for eCQM validation due to our proposal to reduce the number of records collected from 32 records as finalized in the FY 2017 IPPS/LTCH PPS Final rule to 24 records for the FY 2021 payment determination. We refer readers to section XIII.B.6.d.(2) of the preamble of this proposed rule for more detail on these calculations.

(c) Impact of the Proposed Modifications to the Validation Exclusions for the FY 2020 Payment Determination and Subsequent Years

In section IX.A.11.b. of the preamble of this proposed rule, we are proposing a new eCQM validation exclusion criterion. Specifically, hospitals that do not have at least five discharges for at least one reported measure (among the six required eCQMs proposed for the CY 2017 and CY 2018 eCQM reporting periods) included in their QRDA I file submissions would be excluded from the random sample of up to 200 hospitals selected for eCQM validation for the FY 2020 payment determination and subsequent years. We also are proposing, for the FY 2020 payment determination and subsequent years, to exclude hospitals meeting the newly proposed exclusion criterion discussed above and/or either of the two exclusion criteria finalized in the FY 2017 IPPS/LTCH PPS Final rule (81 FR 57178). Lastly, we are proposing that the three exclusions would be applied before the random selection of 200 hospitals for eCQM validation, such that hospitals meeting any of these exclusions would not be eligible for selection.

In section IX.A.11.b. of the preamble of this proposed rule, we also are proposing to exclude the following cases from validation for those hospitals that are chosen to participate in eCQM validation: (1) Episodes of care that are longer than 120 days; and (2) cases with a zero denominator for each measure, for the FY 2020 payment determination and subsequent years.

We do not believe that these proposals will impact the burden experienced by hospitals because, while they influence which hospitals and cases would be selected, they would not change the number of hospitals that must participate in eCQM validation, the number of records that would be collected for validation, or the validation reporting requirements for the hospitals selected.

(d) Impact of the Proposed Modifications to the Medical Record Submission Requirements for the FY 2021 Payment Determination and Subsequent Years

In section IX.A.11.b. of the preamble of this proposed rule, we are proposing that for hospitals participating in eCQM validation we: (1) Require submission of at least 75 percent of sampled eCQM measure medical records in a timely and complete manner; and (2) that the accuracy of eCQM data submitted for validation would not affect a hospital’s validation score (81 FR 57180). We do not expect these proposals to impact the burden experienced by hospitals, as we are continuing existing policies.

(e) Impact of the Proposed Educational Review Process for Chart-Abstracted Measures for the FY 2020 Payment Determination and Subsequent Years

In section IX.A.11.c. of the preamble of this proposed rule, we are proposing to formalize the process of allowing hospitals to use an educational review process to correct validation results for the first three quarters of validation for chart-abstracted measures. Second, we are proposing to update the process to specify that if the results of an educational review indicate that we incorrectly scored a hospital, the corrected score would be used to compute the hospital’s final validation score whether or not the hospital submits a reconsideration request. As stated in the FY 2016 IPPS/LTCH PPS Final rule (80 FR 40762), we estimate a burden of 15 minutes per hospital to report structural measure data and to complete all forms, including the reconsideration request form and the educational review form. We refer readers to the FY 2017 IPPS/LTCH PPS Final rule for more detailed information on the burden associated with the chart-abstracted validation requirements (81 FR 57260). Although this proposal may allow hospitals to avoid the formal reconsideration process, we do not expect this proposal to change our previously finalized burden estimates for the chart-abstracted measures validation process or add any additional burden, as it would not change the requirements for selecting hospitals for validation of chart-abstracted measures nor change the chart-abstracted validation reporting requirements for the selected hospitals.

5. Impact of the Proposed Voluntary Reporting on the Hybrid Hospital-Wide 30-Day Readmission Measure for the CY 2018 Reporting Period

In section IX.A.7.a. of the preamble of this proposed rule, we are proposing voluntary reporting on the Hybrid Hospital-Wide 30-Day Readmission measure for the CY 2018 reporting period. This measure uses both claims-based data as well as a set of 13 core clinical data elements from patient electronic health records (EHRs) and linking variables. We do not expect this proposal to add any additional burden to hospitals to report the claims-based portion of this measure because these data are already reported to the Medicare program for payment purposes.

As described in section IX.A.7.b. of the preamble of this proposed rule, we are proposing that hospitals submit the 13 core clinical data elements and the six data elements required for linking with claims data for this measure using the same submission process required for eCQM reporting, specifically, that these data be reported using QRDA I files submitted to the CMS data receiving system. Accordingly, we expect the burden associated with voluntarily reporting this measure to be similar to our estimates for eCQM reporting (that is 10 minutes per measure, per quarter). We anticipate that approximately 100 hospitals would voluntarily report the Hybrid Hospital-Wide 30-Day Readmission measure. As such, this proposal represents an annual burden increase of 67 hours across up to 100 hospitals voluntarily participating. Using the wage estimate described above, we estimate this to represent a cost increase of $2,200 ($32.84 hourly wage × 67 annual hours) across up to 100 hospitals voluntarily reporting data for this measure. We refer readers to section XIII.B.6.e. of the preamble of this proposed rule for more detail on these burden calculations.

6. Impact of the Proposed Refined Measurement of the HCAHPS Survey Measure for the FY 2020 Payment Determination and Subsequent Years

In section IX.A.6.a. of the preamble of this proposed rule, we are proposing to refine and update the HCAHPS Survey measure by replacing the set of three current Pain Management questions with the “Communication About Pain” composite measure beginning with the FY 2020 payment determination. There is no additional burden associated with the refinement of these questions because we are rewording the existing questions to include language that focuses on communication about pain. In addition, consistent with previous years (81 FR 57261), the burden estimate for the Hospital IQR Program excludes the burden associated with the HCAHPS Survey measure, which is submitted under a separate information collection request and approved under OMB control number 0938–0981.

7. Impact of the Proposed Update to the Hospital 30-Day, All-Cause, Risk-Standardized Mortality Rate Following Acute Ischemic Stroke Measure for the FY 2023 Payment Determination and Subsequent Years

In section IX.A.6.b. of the preamble of this proposed rule, we are proposing to update the Hospital 30-Day, All-Cause, Risk-Standardized Mortality Rate Following Acute Ischemic Stroke measure to include the use of NIH stroke scale claims data for risk adjustment beginning with the FY 2023 payment determination. Because this proposed update would result only in the inclusion of additional claims-based data that are already reported to the Medicare program for payment purposes, we believe no additional burden on hospitals would result from the update to the stroke mortality measure.
8. Impact of Confidential and Potential Future Public Reporting of Readmission Measure Data Stratified by Social Risk Factors

In section IX.A.13. of the preamble of this proposed rule we discuss our intent to provide confidential reports to hospitals that include measure data stratified by dual eligible status for the Hospital 30-day, All-Cause, Risk-Standardized Readmission Rate following Pneumonia Hospitalization and Hospital 30-day, All-Cause, Risk-Standardized Mortality Rate (RSRR) for Pneumonia measures. Because this proposal is related to the way we would display data, and not the methods of data collection implemented by the hospitals, we believe no additional burden on hospitals would result from the confidential reporting of stratified measure data using social risk factor indicators. We note that all measures for which we might consider confidential reporting or public display of stratified measure data would already be included in the Hospital IQR Program, and as claims-based measures, we do not expect any additional burden because these data are already reported to the Medicare program for payment purposes.

9. Impact of Changes to the Hospital IQR Program Extraordinary Circumstances Exceptions (ECE) Policy for the FY 2020 Payment Determination and Subsequent Years

In section IX.A.15.b. of the preamble of this proposed rule we discuss our intent to align the naming of this exception policy and update CFR 412.140 to reflect our current ECE policies. We also are clarifying the timing of CMS response to ECE requests. Because we are not seeking any new or additional information in our ECE proposals, we believe the updates will have no effect on burden for hospitals.

10. Summary of Effects

Historically, 100 hospitals, on average, that participate in the Hospital IQR Program do not receive the full annual percentage increase in any fiscal year due to the requirements of this program. We anticipate that, because of the new requirements for reporting we are proposing for the FY 2020 payment determination, the number of hospitals not receiving the full annual percentage increase may increase due to the changes in policy described above. At this time, information is not available to determine the precise number of hospitals that will not meet the requirements to receive the full annual percentage increase for the FY 2020 payment determination. If the number of hospitals failing to receive the full annual percentage increase does increase because of the new requirements, we anticipate that, over the long run, this number will decline as hospitals gain more experience with these requirements.

In implementing the Hospital IQR Program and other quality reporting programs, we have focused on measures that have high impact and support CMS and HHS priorities for improving the quality of care and value for Medicare beneficiaries.

L. Effects of Proposed Requirements for the PPS-Exempt Cancer Hospital Quality Reporting (PCHQR) Program

In section IX.B. of the preamble of this proposed rule, we discuss our proposed policies for the quality data reporting program for PPS-exempt cancer hospitals (PCHs), which we refer to as the PPS-Exempt Cancer Hospital Quality Reporting Program (PCHQR) Program. The PCHQR Program is authorized under section 1866(k) of the Act, which was added by section 3005 of the Affordable Care Act. There is no financial impact to PCH Medicare reimbursement if a PCH does not submit data.

In section IX.B.4. of the preamble of this proposed rule, we are proposing to adopt four claims-based measures beginning with the FY 2020 program: (1) Proportion of Patients Who Died from Cancer Receiving Chemotherapy in the Last 34 Days of Life (NQF #0210); (2) Proportion of Patients Who Died from Cancer Admitted to the ICU in the Last 30 Days of Life (NQF #0213); (3) Proportion of Patients Who Died from Cancer Not Admitted to the ICU but NQF #0215, and (4) Proportion of Patients Who Died from Cancer Admitted to Hospice for Less Than Three Days (NQF #0216).

In conjunction with our proposal in section IX.B.4. of the preamble of this proposed rule to remove the existing chart-abstracted measures beginning with the FY 2020 program—(1) Adjuvant Chemotherapy is Considered or Administered Within 4 Months (120 Days) of Diagnosis to Patients Under the Age of 80 with AJCC III (Lymph Node Positive) Colon Cancer (PCH–01/NQF #0259); (2) Adjuvant Chemotherapy is Considered or Administered Within 4 Months (120 Days) of Diagnosis to Patients Under the Age of 70 with AJCC T4c, or Stage II or III Hormone Receptor Negative Breast Cancer (PCH–02/NQF #0250); (3) Adjuvant Chemotherapy is Considered or Administered Within 4 Months (120 Days) of Diagnosis to Patients Under the Age of 70 with AJCC T4c, or Stage II or III Hormone Receptor Negative Breast Cancer (PCH–02/NQF #0250); (4) Proportion of Patients Who Died from Cancer Receiving Chemotherapy in the Last 34 Days of Life (NQF #0210); (5) Proportion of Patients Who Died from Cancer Admitted to the ICU in the Last 30 Days of Life (NQF #0213); (6) Proportion of Patients Who Died from Cancer Not Admitted to the ICU but...
In accordance with section 1866(s)(4)(A)(i) of the Act, we will implement a 2 percentage point reduction in the FY 2020 market basket update for IPFs that have failed to comply with the IPFQR Program requirements for the FY 2020 payment determination. In section IX.D. of the preamble of this proposed rule, we discuss how the 2 percentage point reduction will be applied. For the FY 2017 payment determination (that is, data collected during CY 2015 and submitted in CY 2016) of the 1,647 IPFs eligible for the IPFQR Program, we did not consider the full market basket update due to reasons specific to the IPFQR Program; 22 of these IPFs chose not to participate and 27 did not meet the requirements of the Program. We anticipate that even fewer IPFs would receive the reduction for FY 2018 as IPFs become more familiar with the requirements. Thus, we estimate that the IPFQR Program will have a negligible impact on overall IPF payments for FY 2018.

We intend to closely monitor the effects of this quality reporting program on IPFs and help facilitate reducing outcomes through ongoing stakeholder education, national trainings, and a technical help desk. We are proposing provisions that impact the FY 2018 procedural requirements and subsequent years, and the FY 2020 payment determinations and subsequent years. We refer readers to section XIII.B.10. of the preamble of this proposed rule for details discussing information collection requirements for the IPFQR Program.

O. Effects of Proposed Requirements Regarding the Electronic Health Record (EHR) Incentive Programs and Meaningful Use

In section IX.E. of the preamble of this proposed rule, we discuss proposed policies for eligible hospitals and CAHs reporting CQMs electronically under the Medicare and Medicaid EHR Incentive Programs in 2017.

As outlined in this proposed rule, we are proposing the following modifications to the CY 2017 final CQM policies: (1) Revise the CY 2017 reporting requirements for eligible hospitals and CAHs reporting CQMs electronically under the Medicare and Medicaid EHR Incentive Programs; (2) revise the number of CQMs eligible hospitals and CAHs are required to report electronically for CY 2017 to 6 (self-selected) available CQMs, and (3) require eligible hospitals and CAHs to report CQMs for reporting period requires the same number of quarters of data; and (2) revise the number of CQMs eligible hospitals and CAHs are required to report electronically for CY 2017 to 6 (self-selected) available CQMs. In addition, we are proposing the following CQM reporting requirements for CY 2018: (1) Eligible hospitals and CAHs reporting CQMs electronically that demonstrate meaningful use for the first time in 2018 or that have demonstrated meaningful use in any year prior to 2018, the reporting period would be the first 3 quarters of CY 2018 with a submission period (Medicare EHR Incentive Program only) consisting of the 2 months following the close of the calendar year, ending on February 28, 2019; (2) eligible hospitals and CAHs reporting CQMs electronically would be required to report at least 6 (self-selected) of the available CQMs; (3) eligible hospitals and CAHs that report CQMs by attestation under the Medicare EHR Incentive Program because electronic reporting is not feasible, and eligible hospitals and CAHs that report CQMs by attestation under their State’s Medicaid EHR Incentive Program, would be required to report on all 16 available CQMs; and (4) eligible hospitals and CAHs reporting CQMs by attestation under the Medicare EHR Incentive Program would have a submission period that would be the 2 months following the close of the CY 2018 CQM reporting period, ending February 28, 2019.

Because the proposed reporting requirements for data collection regarding the reporting of CQMs electronically under the Medicare and Medicaid EHR Incentive Programs would align with the reporting requirements under the Hospital IQR Program, we do not believe that there is any additional burden for the collection of such information. We are not proposing modifications for the CQMs reporting requirements by attestation. Therefore, there would be no change in burden associated with attestation of CQMs.

In section IX.F. of the preamble of this proposed rule, we discuss proposed policies regarding clinical quality measures for EPs participating in the Medicare EHR Incentive Program. We note that there may be costs incurred by States associated with systems development as a result of the proposed policies. State attestation systems would likely require minor updates, which may be eligible for support through enhanced Federal funding, subject to CMS prior approval, if outlined in an updated Implementation Advance Planning Document (IAPD). We anticipate that eligible professionals (EPs) may also face minor burden increases and increased costs for updating clinical quality measures and reporting capabilities in the EHR. However, we intend to reduce EP burden and simplify the program through these proposals, which are intended to better align CQM alignment at the State attestation system and EP levels would both reduce burden associated with reporting on multiple CMS programs and enhance State and CMS operational efficiency.

In section IX.G.1. of the preamble of this proposed rule, we discuss our proposals to change the EHR reporting period in 2018 from the full CY 2018 to any continuous 90-day period within CY 2018 for all returning EPs, eligible hospitals and CAHs in the Medicare and Medicaid EHR Incentive Programs. We do not believe that modifying the EHR reporting period would cause an increase in cost as the reporting requirements for a 90 day reporting period are virtually the same for a full calendar year reporting period as the requirements for a full calendar year reporting period and 90 day EHR reporting period reporting requires the same number of objectives and measures to be met.

In section IX.G.2. of the preamble of this proposed rule, as required by the 21st Century Cures Act (Pub. L. 114–255), we are proposing an exemption from the payment adjustments under sections 1848(a)(7)(A), 1886(b)(3)(B)(ix)(I), and 1814(l)(4) of the Act for EPs, eligible hospitals and CAHs, respectively, that demonstrate through an application process that compliance with the
requirement for being a meaningful EHR user is not possible because their certified EHR technology has been decertified under ONC’s Health IT Certification Program. The application process involves participants completing an application form for an exception. While the form is standardized, we believe it is exempt from the PRA. The form is structured as an attestation. Therefore, we believe it is exempt under 5 CFR 1320.3(h)(1) of the implementing regulations of the PRA. The form is an attestation that imposes no burden beyond what is required to provide identifying information and to attest to the applicable information.

In section IX.G.3. of the preamble of this proposed rule, as required by the 21st Century Cures Act, we are proposing to exempt ambulatory surgical center-based EPs from the 2017 and 2018 payment adjustments under section 1848(a)(7)(A) of the Act if they furnish substantially all of their covered professional services in an ambulatory surgical center. We do not believe this requirement would cause an increase in burden as CMS would identify the EPs who might meet this requirement.

For the information collection requirements relating to the above proposals, we refer readers to section XIII.B.11. of the preamble of this proposed rule.

P. Effects of Proposed Electronic Signature and Electronic Submission of the Certification and Settlement Summary Page of Medicare Cost Reports

In section X.A. of the preamble of this proposed rule, we discuss our proposal to allow providers to use an electronic signature on the certification statement of the Certification and Settlement Summary page of the Medicare cost report and submit it electronically. This proposal would result in savings to providers.

Using the most current data from Medicare’s System for Tracking Audit and Reimbursement, approximately 51,000 providers file a Medicare cost report and, therefore, must currently mail the Certification and Settlement Summary page. Because most providers mail the Certification and Settlement Summary page with an electronic signature, this proposal would collectively save these providers approximately $362,000 in postage costs. This is an underestimate as it does not include mailing costs when providers choose to mail the Certification and Settlement Summary page to their contractors via overnight mail at a significantly higher expense.

Q. Effects of Proposed Changes Relating to Survey and Certification Requirements

In section XI.A. of the preamble of this proposed rule, we discuss our proposals to revise the application and reapplication procedures for national accrediting organizations (AOs) to require them to post final survey results and acceptable plans of corrections (PoCs) to the Web sites. The AOs programs consist of 10 provider-supplier AO programs and 4 Advanced Diagnostic Imaging (ADI) AO programs. All of these AO programs would be affected by the proposal.

As of the end of FY 2016, there were a total of 12,454 deemed providers and suppliers divided among 4 CMS provider/supplier-approved AO programs. Accreditation surveys for deemed provider and suppliers are conducted on a triennial basis, with a varying number of surveys conducted annually by the AO based on the provider’s or supplier’s entry into the AO program. It is estimated that approximately 5,492 survey reports and corresponding PoCs would need to be posted annually across the 10 provider/supplier AO programs. In addition to the provider/supplier-approved AO programs, there were 16,873 ADI suppliers divided among 4 CMS-approved ADI AOs. It is estimated that approximately 2,128 survey reports and corresponding PoCs would need to be posted annually across the 4 ADI AOs. We are not able to estimate the cost associated with the proposed requirement for posting of the surveys reports and corresponding PoCs at this time. We are seeking public comments, particularly from AOs, regarding the potential initial cost of modifications to the AOs’ existing public Web sites and the ongoing cost associated with uploading survey reports and PoCs. We recommend that AOs provide public comments in response to this proposed rule on their estimated costs for posting survey reports and corresponding PoCs. We will consider any public comments received and address them in the final rule.

There is no financial impact of the proposal on deemed facilities as the survey reports and associated PoCs would not be posted by the facilities, but would be posted by the AOs affiliated with the providers or suppliers or ADIs. The overall impact would be determined based on the total costs for posting of the survey reports for the 10 provider-supplier AO programs and the 4 ADI AOs.

In section XI.B. of the preamble of this proposed rule, we discuss our proposals to eliminate the term “newspaper” from the requirement to publish public notice upon a provider’s involuntary termination for RHCs, FQHCs, ASCs, and OPOs. Eliminating the term “newspaper” would allow greater flexibility for the CMS Regional Offices in publishing public notices and would also reduce burden on the CMS Regional Offices. The print newspaper advertisements for an involuntary termination are required to be purchased by the CMS Regional Office assigned to that provider or supplier. The advertisement is placed under the legal advertisement section of the local newspaper outlet. A single CMS Regional Office may incur an average annual cost of approximately $3,000 to $5,000 for the publication of involuntary termination notices for the providers or suppliers assigned to its region. For example, from 2014 to 2016, the Dallas Regional Office spent $14,331.89 on the publication of termination notices in local newspapers, with costs of $3,949.45 in 2014, costs of $5,386.67 in 2015, and costs of $4,908.77 in 2016. In same timeframe of 2014 to 2016, the Philadelphia Regional Office spent a total of $7,114.75 and the Kansas Regional Office spent a total of $11,121.40. The table below depicts the actual FY 2016 costs for all 10 CMS Regional Offices.

<table>
<thead>
<tr>
<th>Regional Office</th>
<th>2016 Costs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Boston</td>
<td>$4,766</td>
</tr>
<tr>
<td>New York</td>
<td>$645</td>
</tr>
<tr>
<td>Philadelphia</td>
<td>$3,570</td>
</tr>
<tr>
<td>Atlanta</td>
<td>$6,712</td>
</tr>
<tr>
<td>Chicago</td>
<td>$10,853</td>
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<tr>
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</tr>
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<td>Denver</td>
<td>$910</td>
</tr>
<tr>
<td>San Francisco</td>
<td>$1,507</td>
</tr>
<tr>
<td>Seattle</td>
<td>$707</td>
</tr>
</tbody>
</table>

Total Cost ................................ 37,020.00

If one CMS Regional Office spends approximately $5,000 annually, and there are 10 CMS Regional Offices, the overall cost nationwide per annum for termination notices could be as high as $50,000.

The cost associated with the involuntary termination notice is assessed only to the CMS Regional Offices. The provider or supplier is not required to post a notice for an involuntary termination. Therefore, there would be no associated costs for the provider or supplier.

All CMS Regional Offices have Web sites available to the public, which are regularly maintained and updated. Creation of a subsite to reflect termination notices for providers would be at no cost to CMS. In addition, the use of Regional Press Officers to convey termination of a provider would be a minimal cost to CMS and absorbed through the Survey & Certification budget.

R. Effects of Clarification of Limitations on the Valuation of Depreciable Assets Disposed of on or After December 1, 1997

In section X.B. of the preamble of this proposed rule, we discuss our proposal to revise the Medicare provider reimbursement regulations to clarify our longstanding policy pertaining to allowable costs and the limits on the valuation of a depreciable asset that may be recognized in establishing an appropriate allowance for depreciation for assets disposed of on or after December 1, 1997. Specifically, we are clarifying that the elimination of the gain or loss for depreciable assets applies to assets a provider disposes of by sale or scrapping on or after December 1, 1997, regardless of whether the asset is scrapped, sold as an individual asset of a Medicare participating provider, or sold incident to a provider change of ownership.

Because we are not proposing any change in policy, but rather are restating longstanding Medicare policy, there is no economic impact on providers resulting from this policy clarification.

S. Alternatives Considered

This proposed rule contains a range of proposed policies. It also provides descriptions of the statutory provisions that are addressed, identifies the proposed policies, and presents rationales for our decisions and, where relevant, alternatives that were considered.
As discussed in section III.H. of the preamble of this proposed rule, we are not proposing to extend the imputed floor policy for developing the hospital wage index. We note that if the imputed floor policy were not to expire at the end of FY 2017, we would estimate that IPPS payments would increase by approximately $19 million in New Jersey, $19 million in Rhode Island, and $9 million in Delaware. Because the imputed floor policy is budget neutral nationally, these additional IPPS payments as a result of the imputed floor policy not expiring would reduce payments to all IPPS hospitals by approximately $47 million.

T. Reducing Regulation and Controlling Regulatory Costs

Executive Order 13771, titled “Reducing Regulation and Controlling Regulatory Costs,” was issued on January 30, 2017. Section 2(a) of Executive Order 13771 requires an agency, unless prohibited by law, to identify at least two existing regulations to be repealed when the agency publicly proposes for notice and comment, or otherwise promulgates, a new regulation. In furtherance of this requirement, section 2(c) of Executive Order 13771 requires that the new incremental costs associated with new regulations shall, to the extent permitted by law, be offset by the elimination of existing costs associated with at least two prior regulations. OMB’s implementation guidance, issued on April 5, 2017, explains that “Federal spending regulatory actions that cause only income transfers between taxpayers and program beneficiaries (for example, regulations associated with Medicare spending) are considered ‘transfer rules’ and are not covered by EO 13771. . . . However . . . such regulatory actions may impose requirements apart from transfers. . . . In those cases, the actions would need to be offset to the extent they impose more than de minimis costs.

Examples of ancillary requirements that may require offsets include new reporting or recordkeeping requirements. . . . Analogously, if an action reduces the stringency of requirements or conditions . . . the action may qualify as an EO 13771 deregulatory action.” Table I of section I.G., Table III of section I.I., and Table IV of section I.J. of this Appendix show the IPPS operating and capital costs and LTCH PPS costs, respectively, on affected entities. The implications of the rule’s costs and cost savings will be further considered in the context of our compliance with Executive Order 13771.

U. Overall Conclusion

1. Acute Care Hospitals

Table I of section I.G. of this Appendix demonstrates the estimated distributional impact of the IPPS budget neutrality requirements for the proposed MS–DRG and wage index changes, and for the wage index reclassifications under the MCGRR. Table I also shows a projected overall increase of 1.7 percent in operating payments before accounting for the proposed changes in Medicare DSH payments and uncompensated care payments. When combined with the impact of those proposed changes, consistent with our policy discussed in section V.G. of the preamble of this proposed rule, we estimate that operating payments would increase by approximately 2.9 percent in FY 2018, or approximately $3.2 billion. We also currently estimate that the proposed changes in new technology add-on payments for FY 2018 would decrease spending by approximately $2 million and the proposed changes to the volume decrease adjustment would increase in spending by approximately $15 million. In addition, we estimate the change in low-volume hospital payments, including the statutory expiration of the temporary increase in the low-volume hospital payment adjustment in FY 2018 would decrease spending by approximately $311 million in FY 2018. These estimates, combined with our estimated increase in FY 2018 operating payment of $3.2 billion, would result in an estimated increase of approximately $2.8 billion for FY 2018. We estimate that hospitals would experience a 2.4 percent increase in capital payments per case, as shown in Table III of section I.I. of this Appendix. We project that there would be a $212 million increase in capital payments in FY 2018 compared to FY 2017. The cumulative operating and capital payments would result in a net increase of approximately $3.1 billion to IPPS providers. The discussions presented in the previous pages, in combination with the rest of this proposed rule, constitute a regulatory impact analysis.

2. LTCHs

Overall, LTCHs are projected to experience a decrease in estimated payments per discharge in FY 2018. In the impact analysis, we are using the proposed rates, factors, and policies presented in this proposed rule based on the best available claims and CCR data to estimate the change in payments under the LTCH PPS for FY 2018. Accordingly, based on the best available data for the 415 LTCHs in our database, we estimate that FY 2017 LTCH PPS payments would decrease approximately $173 million relative to FY 2017 as a result of the proposed payment rates and factors presented in this proposed rule.

V. Regulatory Review Costs

If regulations impose administrative costs on private entities, such as the time needed to read and interpret this proposed rule, we should estimate the cost associated with regulatory review. Due to the uncertainty involved with accurately quantifying the number of entities that will review this proposed rule, we assume that the total number of commenters on last year’s proposed rule will be the number of reviewers of this proposed rule. We acknowledge that this assumption may understate or overstate the costs of reviewing this rule. It is possible that not all commenters reviewed last year’s rule in detail, and it is also possible that some reviewers chose not to comment on the proposed rule. For these reasons, we believe that the number of past commenters would be a fair estimate of the number of reviewers of this proposed rule. We welcome any public comments on the approach in estimating the number of entities that will review this proposed rule.

We also recognize that different types of entities are in many cases affected by mutually exclusive sections of this proposed rule. Therefore, for the purposes of our estimate, we assume that each reviewer reads approximately 50 percent of the proposed rule. We are seek public comments on this assumption.

Using the wage information from the BLS for medical and health service managers (Code 11–9111), we estimate that the cost of reviewing this proposed rule is $90.16 per hour, including overhead and fringe benefits (https://www.bls.gov/oes/2015/may/oaus4_631000.htm). Assuming an average reading speed, we estimate that it would take approximately 16 hours for the staff to review half of this proposed rule. For each IPPS hospital or LTCH that reviews this proposed rule, the estimated cost is $1,442.56 (16 hours × $90.16). Therefore, we estimate that the total cost of reviewing this proposed rule is $2,071.516 ($1,442.56 × 1,436 reviewers).

II. Accounting Statements and Tables

A. Acute Care Hospitals

As required by OMB Circular A–4 (available at http://www.whitehouse.gov/omb/circulars/a004/a-4.pdf), in the following Table V, we have prepared an accounting statement showing the classification of the expenditures associated with the provisions of this proposed rule as they relate to acute care hospitals. This table provides our best estimate of the change in Medicare payments to providers as a result of the proposed changes to the IPPS presented in this proposed rule. All expenditures are classified as transfers to Medicare providers.

The costs to the Federal Government associated with the proposed policies in this proposed rule are estimated at $3.1 billion.

<table>
<thead>
<tr>
<th>Category</th>
<th>Transfers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Annualized Monetized Transfers</td>
<td>$3.1 billion.</td>
</tr>
<tr>
<td>From Whom to Whom</td>
<td>Federal Government to IPPS Medicare Providers.</td>
</tr>
</tbody>
</table>

### Table V—Accounting Statement: Classification of Estimated Expenditures Under the IPPS from FY 2017 to FY 2018

- **Annualized Monetized Transfers**: $3.1 billion
- **From Whom to Whom**: Federal Government to IPPS Medicare Providers
B. LTCHs

As discussed in section I.J. of this Appendix, the impact analysis of the proposed payment rates and factors presented in this proposed rule under the LTCH PPS is projected to result in a decrease in estimated aggregate LTCH PPS payments in FY 2018 relative to FY 2017 of approximately $173 million based on the data for 415 LTCHs in our database that are subject to payment under the LTCH PPS. Therefore, as required by OMB Circular A–4 (available at http://www.whitehouse.gov/omb/circulars/a004/a-4.pdf), in Table VI, we have prepared an accounting statement showing the classification of the expenditures associated with the provisions of this proposed rule as they relate to savings associated with the changes to the LTCH PPS. Table VI provides our best estimate of the estimated change in Medicare payments under the LTCH PPS as a result of the proposed payment rates and factors and other provisions presented in this proposed rule based on the data for the 415 LTCHs in our database. All expenditures are classified as transfers to Medicare providers (that is, LTCHs).

The savings to the Federal Government associated with the policies for LTCHs in this proposed rule are estimated at $173 million.

<table>
<thead>
<tr>
<th>TABLE VI—ACCOUNTING STATEMENT: CLASSIFICATION OF ESTIMATED EXPENDITURES FROM THE FY 2017 LTCH PPS TO THE FY 2018 LTCH PPS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Category</td>
</tr>
<tr>
<td>Annualized Monetized Transfers</td>
</tr>
<tr>
<td>From Whom to Whom</td>
</tr>
</tbody>
</table>

III. Regulatory Flexibility Act (RFA)

Analysis

The RFA requires agencies to analyze options for regulatory relief of small entities. For purposes of the RFA, small entities include small businesses, nonprofit organizations, and small government jurisdictions. We estimate that most hospitals and most other providers and suppliers are small entities as that term is used in the RFA. The great majority of hospitals and most other health care providers and suppliers are small entities, either by being nonprofit organizations or by meeting the SBA definition of a small business (having revenues of less than $7.5 million to $38.5 million in any 1 year). (For details on the latest definitions of health care providers, we refer readers to page 36 of the Table of Small Business Size Standards for NAIC 622 found on the SBA Web site at: http://www.sba.gov/sites/default/files/files/Size_Standards_Table.pdf.) For purposes of the RFA, all hospitals and other providers and suppliers are considered to be small entities. Individuals and States are not included in the definition of a small entity. We believe that the provisions of this proposed rule relating to acute care hospitals will have a significant impact on small entities as explained in this Appendix. For example, we refer readers to “Table I—Impact Analysis of Proposed Changes to the IPPS for Operating Costs for FY 2018.” Because we lack data on individual hospital receipts, we cannot determine the number of small proprietary LTCHs. Therefore, we are assuming that all LTCHs are considered small entities for the purpose of the analysis in section I.J. of this Appendix. MACs are not considered to be small entities. Because we acknowledge that many of the affected entities are small entities, the analysis discussed throughout the preamble of this proposed rule constitutes our regulatory flexibility analysis. This proposed rule contains a range of proposed policies. It provides statutory provisions that are addressed, identifies the proposed policies, and presents rationales for our decisions and, where relevant, alternatives that were considered.

In this proposed rule, we are soliciting public comments on our estimates and analysis of the impact of our proposals on those small entities. Any public comments that we receive and our responses will be presented in the final rule.

IV. Impact on Small Rural Hospitals

Section 1102(b) of the Social Security Act requires us to prepare a regulatory impact analysis for any proposed or final rule that may have a significant impact on the operations of a substantial number of small rural hospitals. This analysis must conform to the provisions of section 603 of the RFA. With the exception of hospitals located in certain New England counties, for purposes of section 1102(b) of the Act, we define a small rural hospital as a hospital that is located outside of an urban area and has fewer than 100 beds. Section 601(g) of the Social Security Amendments of 1983 (Pub. L. 98–21) designated hospitals in certain New England counties as belonging to the adjacent urban area. Thus, for purposes of the IPPS and the LTCH PPS, we continue to classify these hospitals as urban hospitals. (We refer readers to Table I in section I.G. of this Appendix for the quantitative effects of the proposed policy changes under the IPPS for operating costs.)

V. Unfunded Mandates Reform Act Analysis

Section 202 of the Unfunded Mandates Reform Act (Pub. L. 104–4) also requires that agencies assess anticipated costs and benefits before issuing any rule whose mandates require spending in any 1 year of $100 million in 1995 dollars, updated annually for inflation. In 2017, that threshold level is approximately $146 million. This proposed rule would not mandate any requirements for State, local, or tribal governments, nor would it affect private sector costs.

VI. Executive Order 13175

Executive Order 13175 directs agencies to consult with Tribal officials prior to the formal promulgation of regulations having tribal implications. This proposed rule contains provisions applicable to hospitals and facilities operated by the Indian Health Service or Tribes or Tribal organizations under the Indian Self-Determination and Education Assistance Act and, thus, has tribal implications. Therefore, in accordance with Executive Order 13175 and the CMS Tribal Consultation Policy (December 2015), CMS will consult with Tribal officials on these Indian-specific provisions of the proposed rule prior to the formal promulgation of this rule.

VII. Executive Order 12866

In accordance with the provisions of Executive Order 12866, the Executive Office of Management and Budget reviewed this proposed rule.

Appendix B: Recommendation of Update Factors for Operating Cost Rates of Payment for Inpatient Hospital Services

I. Background

Section 1886(e)(4)(A) of the Act requires that the Secretary, taking into consideration the recommendations of MedPAC, recommend update factors for inpatient hospital services for each fiscal year that take into account the amounts necessary for the efficient and effective delivery of medically appropriate and necessary care of high quality. Under section 1886(e)(5) of the Act, we are required to publish update factors recommended by the Secretary in the proposed and final IPPS rules, respectively. Accordingly, this Appendix provides the recommendations for the update factors for the IPPS national standardized amount, the hospital-specific rate for SICs, and the rate-of-increase limits for certain hospitals excluded from the IPPS, as well as LTCHs. In prior years, we have made a recommendation in the IPPS proposed rule and final rule for the update factors for the payment rates for IRFs and IPFs. However, for FY 2018, consistent with approach for FY 2017, we are including the Secretary’s recommendation for the update factors for IRFs and IPFs in separate Federal Register documents at the time that we announce the annual updates for IRFs and IPFs. We also discuss our response to MedPAC’s recommended update factors for inpatient hospital services.

II. Inpatient Hospital Update for FY 2018

A. Proposed FY 2018 Inpatient Hospital Update

As discussed in section V.B. of the preamble to this proposed rule, consistent with section 1886(b)(3)(B) of the Act, as amended by sections 3401(a) and 10319(a) of the Affordable Care Act, we are setting the
applicable percentage increase by applying the following adjustments in the following sequence. Specifically, the applicable percentage increase under the IPPS is equal to the rate-of-increase in the hospital market basket for IPPS hospitals in all areas, subject to a reduction of one-quarter of the applicable percentage increase (prior to the application of other statutory adjustments; also referred to as the market basket update or rate-of-increase (with no adjustments)) for hospitals that fail to submit quality information under rules established by the Secretary in accordance with section 1886(b)(3)(B)(viii) of the Act and a reduction of three-quarters of the applicable percentage increase (prior to the application of other statutory adjustments; also referred to as the market basket update or rate-of-increase (with no adjustments)) for hospitals not considered to be meaningful electronic health record (EHR) users in accordance with section 1866(b)(3)(B)(ix) of the Act, and then subject to an adjustment based on changes in economy-wide productivity (the multifactor productivity (MFP) adjustment), and an additional reduction of 0.75 percentage point as required by section 1886(b)(3)(B)(xii) of the Act. Sections 1886(b)(3)(B)(xi) and (b)(3)(B)(xii) of the Act, as added by section 3401(a) of the Affordable Care Act, state that application of the MFP adjustment and the additional FY 2018 adjustment of 0.75 percentage point may result in the applicable percentage increase being less than zero. We note that, in compliance with section 404 of the MMA, in this proposed rule, we are proposing to replace the FY 2010-based IPPS operating and capital market baskets with the revised and rebased 2014-based IPPS operating and capital market baskets for FY 2018.

For this FY 2018 IPPS/LTCH PPS proposed rule, in accordance with section 1886(b)(3)(B) of the Act, we are proposing to base the proposed FY 2018 market basket update used to determine the applicable percentage increase for the IPPS on the IHS Global Insight, Inc.’s (IGI’s) fourth quarter 2016 forecast of the proposed 2014-based IPPS market basket rate-of-increase with historical data through third quarter 2016, which is estimated to be 2.9 percent. In accordance with section 1886(b)(3)(B) of the Act, as amended by section 3401(a) of the Affordable Care Act, in section V.B. of the preamble of this proposed rule, we are proposing an MFP adjustment of 0.4 percent for FY 2018. Therefore, based on IGI’s fourth quarter 2016 forecast of the proposed 2014-based IPPS market basket, depending on whether a hospital submits quality data under the rules established in accordance with section 1886(b)(3)(B)(viii) of the Act (hereafter referred to as a hospital that submits quality data) and is a meaningful EHR user under section 1886(b)(3)(B)(ix) of the Act (hereafter referred to as a hospital that is a meaningful EHR user), there are four possible applicable percentage increases that can be applied to the standardized amount. Below we provide a table summarizing the four proposed applicable percentage increases.

<table>
<thead>
<tr>
<th>FY 2018</th>
<th>Hospital submitted quality data and is a meaningful EHR user</th>
<th>Hospital submitted quality data and is NOT a meaningful EHR user</th>
<th>Hospital did NOT submit quality data and is a meaningful EHR user</th>
<th>Hospital did NOT submit quality data and is NOT a meaningful EHR user</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proposed Market Basket Rate-of-Increase</td>
<td>2.9</td>
<td>2.9</td>
<td>2.9</td>
<td>2.9</td>
</tr>
<tr>
<td>Proposed Adjustment for Failure to Submit Quality Data under Section 1886(b)(3)(B)(viii) of the Act</td>
<td>0.0</td>
<td>0.0</td>
<td>-0.725</td>
<td>-0.725</td>
</tr>
<tr>
<td>Proposed Adjustment for Failure to be a Meaningful EHR User under Section 1886(b)(3)(B)(x) of the Act</td>
<td>0.0</td>
<td>-2.175</td>
<td>0.0</td>
<td>-2.175</td>
</tr>
<tr>
<td>Proposed MFP Adjustment under Section 1886(b)(3)(B)(xi) of the Act</td>
<td>-0.75</td>
<td>-0.75</td>
<td>-0.75</td>
<td>-0.75</td>
</tr>
<tr>
<td>Proposed Applicable Percentage Increase Applied to Standardized Amount</td>
<td>1.75</td>
<td>-0.425</td>
<td>1.025</td>
<td>-1.15</td>
</tr>
</tbody>
</table>

B. Proposed Update for SCHs for FY 2018

Section 1886(b)(3)(B)(iv) of the Act provides that the FY 2018 applicable percentage increase in the hospital-specific rate for SCHs equals the applicable percentage increase set forth in section 1886(b)(3)(B)(i) of the Act (that is, the same update factor as for all other hospitals subject to the IPPS).

(We note that, as discussed in section V.H. of the preamble of this proposed rule, section 205 of the Medicare Access and CHIP Reauthorization Act of 2015 (MACRA) (Pub. L. 114–10, enacted on April 16, 2015) extended the MDH program (which, under previous law, was to be in effect for discharges occurring on or before March 31, 2015 only) for discharges occurring on or after April 1, 2015, through FY 2017 (that is, for discharges occurring on or before September 30, 2017). Therefore, under current law, the MDH program will expire at the end of FY 2017. However, as discussed in section V.H. of the preamble of this proposed rule, MDHs have the opportunity to apply for SCH status in advance of the expiration of the MDH program and be paid as such under certain conditions, as specified in the regulations at 42 CFR 412.92(b)(2)(i) and (b)(2)(v).)

As previously mentioned, the update to the hospital specific rate for SCHs is subject to section 1886(b)(3)(B)(i) of the Act, as amended by sections 3401(a) and 10319(a) of the Affordable Care Act. Accordingly, depending on whether a hospital submits quality data and is a meaningful EHR user, we are proposing the same four possible applicable percentage increases in the table above for the hospital-specific rate applicable to SCHs.

C. Proposed FY 2018 Puerto Rico Hospital Update

As discussed in the FY 2017 IPPS/LTCH PPS final rule (81 FR 56939), prior to January 1, 2016, Puerto Rico hospitals were paid based on 75 percent of the national standardized amount and 25 percent of the Puerto Rico-specific standardized amount. Section 601 of Public Law 114–113 amended section 1886(d)(9)(E) of the Act to specify that the payment calculation with respect to operating costs of inpatient hospital services of a subsection (d) Puerto Rico hospital for inpatient hospital discharges on or after January 1, 2016, shall use 100 percent of the national standardized amount. Because Puerto Rico hospitals are no longer paid with a Puerto Rico-specific standardized amount under the amendments to section 1886(d)(9)(E) of the Act, there is no longer a need for us to propose an update to the Puerto Rico standardized amount. Hospitals in Puerto Rico are now paid 100 percent of the national standardized amount and, therefore, are subject to the same update to the national standardized amount discussed under section V.B.1. of the preamble of this proposed rule. Accordingly, for FY 2018, we are proposing an applicable percentage increase of 1.75 percent to the standardized amount for hospitals located in Puerto Rico.

D. Proposed Update for Hospitals Excluded From the IPPS for FY 2018

Section 1886(b)(3)(B)(ii) of the Act is used for purposes of determining the percentage increase in the rate-of-increase limits for children’s hospitals, cancer hospitals, and hospitals located outside the 50 States, the District of Columbia, and Puerto Rico (that is, short-term acute care hospitals located in the U.S. Virgin Islands, Guam, the Northern Marianas Islands, and America Samoa). Section 1886(b)(3)(B)(iii) of the Act sets the percentage increase in the rate-of-increase limits equal to the market basket percentage increase. In accordance with § 403.752(a) of the regulations, RNHCHs are paid under the provisions of § 413.40, which also use section 1886(b)(3)(B)(ii) of the Act to update the percentage increase in the rate-of-increase limits.

Currently, children's hospitals, PPS-excluded cancer hospitals, RNHCHs, and short-term acute care hospitals located in the U.S. Virgin Islands, Guam, the Northern Marianas Islands, and American Samoa are among the remaining types of hospitals still paid under the reasonable cost methodology, subject to the rate-of-increase limits. As
discussed in section VII. of the preamble of this proposed rule, we are proposing to use the percentage increase in the 2014-based IPPS operating market basket to update the target amounts for children’s hospitals, PPS-excluded cancer hospitals, RNHCIs, and short-term acute care hospitals located in the U.S. Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa for FY 2018 and subsequent fiscal years. Accordingly, for FY 2018, the rate-of-increase percentage to be applied to the target amount for these children’s hospitals, cancer hospitals, RNHCIs, and short-term acute care hospitals located in the U.S. Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa would be the FY 2018 percentage increase in the 2014-based IPPS operating market basket. For this proposed rule, the current estimate of the IPPS operating market basket percentage increase for FY 2018 is 2.9 percent.

E. Proposed Update for LTCHs for FY 2018

Section 123 of Public Law 106–113, as amended by section 307(b) of Public Law 106–554 (and codified at section 1886(m)(1) of the Act), provides the statutory authority for updating payment rates under the LTCH PPS. As discussed in section V.A. of the Addendum to this proposed rule, we are proposing to update the LTCH PPS standard Federal payment rate by 1.0 percent for FY 2018, consistent with the amendments to section 1886(m)(3) of the Act provided by section 411 of MACRA. In accordance with the LTCHQR Program under section 1886(m)(5) of the Act, we are proposing to reduce the annual update to the LTCH PPS standard Federal rate by 2.0 percentage points for failure of a LTCH to submit the required quality data. Accordingly, we are proposing to apply an update factor of 0.99 percent in determining the LTCH PPS standard Federal rate for FY 2018. For LTCHs that fail to submit quality data for FY 2018, we are proposing to apply an annual update to the LTCH PPS standard Federal rate of −1.0 percent (that is, the proposed annual update for FY 2018 of 1.0 percent less 2.0 percentage points for failure to submit the required quality data in accordance with section 1886(m)(5)(C) of the Act and our rules) by applying a proposed update factor of 0.99 percent in determining the LTCH PPS standard Federal rate for FY 2018.

III. Secretary’s Recommendations

MedPAC is recommending an inpatient hospital update in the amount specified in current law for FY 2018. MedPAC’s rationale for this update recommendation is described in more detail below. As mentioned above, section 1886(e)(4)(A) of the Act requires that the Secretary, taking into consideration the recommendations of MedPAC, recommend update factors for inpatient hospital services for each fiscal year that take into account the amounts necessary for the efficient and effective delivery of medically appropriate and necessary care of high quality. Consistent with current law, depending on whether a hospital submits quality data and is a meaningful EHR user, we are recommending the four applicable percentage increases to the standardized amount listed in the table under section II. of this Appendix B. We are recommending that the same applicable percentage increases apply to SCHs.

In addition to making a recommendation for IPPS hospitals, in accordance with section 1886(e)(4)(A) of the Act, we are recommending update factors for certain other types of hospitals excluded from the IPPS. Consistent with our policies for these facilities, we are recommending an update to the target amounts for children’s hospitals, cancer hospitals, RNHCIs, and short-term acute care hospitals located in the U.S. Virgin Islands, Guam, the Northern Mariana Islands, and American Samoa of 2.9 percent.

For FY 2018, consistent with the amendments to section 1886(m)(3) of the Act provided by section 411 of MACRA, for LTCHs that submit quality data, we are recommending an update of 1.0 percent to the LTCH PPS standard Federal rate. For LTCHs that fail to submit quality data for FY 2018, we are recommending an annual update to the LTCH PPS standard Federal rate of −1.0 percent.

IV. MedPAC Recommendation for Assessing Payment Adequacy and Updating Payments in Traditional Medicare

In its March 2017 Report to Congress, MedPAC assessed the adequacy of current payments and costs, and the relationship between payments and an appropriate cost base. MedPAC recommended an update to the hospital inpatient rates in the amount specified in current law. We refer the reader to the March 2017 MedPAC report, which is available for download at www.medpac.gov for a complete discussion on this recommendation. MedPAC expects Medicare margins to decline from 2015 to 2017.

Response: We agree with MedPAC, and consistent with current law, we are proposing an applicable percentage increase for FY 2018 of 1.75 percent, provided the hospital submits quality data and is a meaningful EHR user, consistent with statutory requirements.

We note that, because the operating and capital prospective payment systems remain separate, we are continuing to use separate updates for operating and capital payments. The update to the capital rate is discussed in section III. of the Addendum to this proposed rule.

[FR Doc. 2017–07800 Filed 4–14–17; 4:15 pm]