non-experimental/investigational (Category B) device if all other coverage requirements are met.

(c) Other considerations. Medicare contractors must consider whether any restrictions concerning site of service, indications for use, or any other list of conditions for coverage have been placed on the device's use.

EFFECTIVE DATE NOTE: At 78 FR 74809, Dec. 10, 2013, §405.211 was revised, effective Jan. 1, 2015. For the convenience of the user, the revised text is set forth as follows:

# \$405.211 Coverage of items and services in FDA-approved IDE studies.

(a) Coverage of routine care items and services for Category A (Experimental) devices. Medicare covers routine care items and services furnished in an FDA-approved Category A (Experimental) IDE study if CMS (or its designated entity) determines that the Medicare coverage IDE study criteria in 405.212 are met.

(b) Coverage of Category B (Nonexperimental/ investigational) IDE devices and routine care items and services. Medicare may make payment for a Category B (Nonexperimental/investigational) IDE device and routine care items and services furnished in an FDA-approved Category B (Nonexperimental/investigational) IDE study if CMS (or its designated entity) determines prior to the submission of the first related claim that the Medicare coverage IDE study criteria in §405.212 are met.

(c) CMS (or its designated entity) must review the following to determine if the Medicare coverage IDE study criteria in §405.212 are met for purposes of coverage of items and services described in paragraphs (a) and (b) of this section:

(1) FDA approval letter of the IDE.

(2) IDE study protocol.

(3) IRB approval letter.

(4) NCT number.

(5) Supporting materials, as needed.

(d) Notification. A listing of all CMS-approved Category A (Experimental) IDE studies and Category B (Nonexperimental/investigational) IDE studies shall be posted on the CMS Web site and published in the FEDERAL REGISTER.

# § 405.212 Medicare Coverage IDE study criteria.

(a) For Medicare coverage of items and services described in §405.211, a Category A (Experimental) or Category B (Nonexperimental/investigational) IDE study must meet all of the following criteria:

(1) The principal purpose of the study is to test whether the device improves 42 CFR Ch. IV (10–1–14 Edition)

health outcomes of appropriately selected patients.

(2) The rationale for the study is well supported by available scientific and medical information, or it is intended to clarify or establish the health outcomes of interventions already in common clinical use.

(3) The study results are not anticipated to unjustifiably duplicate existing knowledge.

(4) The study design is methodologically appropriate and the anticipated number of enrolled subjects is adequate to confidently answer the research question(s) being asked in the study.

(5) The study is sponsored by an organization or individual capable of successfully completing the study.

(6) The study is in compliance with all applicable Federal regulations concerning the protection of human subjects found at 21 CFR parts 50, 56, and 812 and 45 CFR part 46.

(7) Where appropriate, the study is not designed to exclusively test toxicity or disease pathophysiology in healthy individuals. Studies of all medical technologies measuring therapeutic outcomes as one of the objectives may be exempt from this criterion only if the disease or condition being studied is life threatening and the patient has no other viable treatment options.

(8) The study is registered with the National Institutes of Health's National Library of Medicine's ClinicalTrials.gov.

(9) The study protocol describes the method and timing of release of results on all pre-specified outcomes, including release of negative outcomes and that the release should be hastened if the study is terminated early.

(10) The study protocol must describe how Medicare beneficiaries may be affected by the device under investigation, and how the study results are or are not expected to be generalizable to the Medicare beneficiary population. Generalizability to populations eligible for Medicare due to age, disability, or other eligibility status must be explicitly described.

(b) [Reserved]

[79 FR 74809, Dec. 10, 2014]

### Centers for Medicare & Medicaid Services, HHS

EFFECTIVE DATE NOTE: At 78 FR 74809, Dec. 10, 2013, §405.212 was added, effective Jan. 1, 2015

#### §405.213 Re-evaluation of a device categorization.

(a) General rules. (1) Any sponsor that does not agree with an FDA decision that categorizes its device as experimental/investigational (Category A) may request re-evaluation of the categorization decision.

(2) A sponsor may request review by CMS only after the requirements of paragraph (b) of this section are met.

(3) No reviews other than those described in paragraphs (b) and (c) of this section are available to the sponsor.

(4) Neither the FDA original categorization or re-evaluation (described in paragraph (b) of this section) nor CMS's review (described in paragraph (c) of this section) constitute an initial determination for purposes of the Medicare appeals processes under part 405, subpart G or subpart H, or parts 417, 473, or 498 of this chapter.

(b) Request to FDA. A sponsor that does not agree with the FDA's categorization of its device may submit a written request to the FDA at any time requesting re-evaluation of its original categorization decision, together with any information and rationale that it believes support recategorization. The FDA notifies both CMS and the sponsor of its decision.

(c) Request to CMS. If the FDA does not agree to recategorize the device, the sponsor may seek review from CMS. A device sponsor must submit its request in writing to CMS. CMS obtains copies of relevant portions of the application, the original categorization decision, and supplementary materials. CMS reviews all material submitted by the sponsor and the FDA's recommendation. CMS reviews only information in the FDA record to determine whether to change the categorization of the device. CMS issues a written decision and notifies the sponsor of the IDE and the FDA.

EFFECTIVE DATE NOTE: At 78 FR 74810, Dec. 10. 2013. §405.213 was amended by revising paragraph (a)(1), effective Jan. 1, 2015, For the convenience of the user, the revised text is set forth as follows:

#### §405.213 Re-evaluation of a device categorization.

(a) \* \* \*

(1) Any sponsor that does not agree with an FDA decision that categorizes its device as Category A (experimental) may request reevaluation of the categorization decision.

#### §405.215 Confidential commercial and trade secret information.

To the extent that CMS relies on confidential commercial or trade secret information in any judicial proceeding, CMS will maintain confidentiality of the information in accordance with Federal law.

### Subpart C—Suspension of Payment, Recovery of Overpayments, and Repayment of Scholarships and Loans

AUTHORITY: Secs. 1102, 1815, 1833, 1842, 1862, 1866, 1870, 1871, 1879 and 1892 of the Social Security Act (42 U.S.C. 1302, 1395g, 1395l, 1395u, 1395y, 1395cc, 1395gg, 1395hh, 1395pp and 1395ccc) and 31 U.S.C. 3711.

SOURCE: 31 FR 13534, Oct. 20, 1966, unless otherwise noted. Redesignated at 42 FR 52826, Sept. 30, 1977.

EDITORIAL NOTE: Nomenclature changes to subpart C of part 4405 appear at 76 FR 5961, Feb. 2. 2011.

#### GENERAL PROVISIONS

#### §405.301 Scope of subpart.

This subpart sets forth the policies and procedures for handling of incorrect payments and recovery of overpayments.

[54 FR 41733, Oct. 11, 1989]

LIABILITY FOR PAYMENTS TO PROVIDERS OR SUPPLIERS AND HANDLING OF IN-CORRECT PAYMENTS

#### §405.350 Individual's liability for payments made to providers and other persons for items and services furnished the individual.

Any payment made under title XVIII of the Act to any provider of services or other person with respect to any item or service furnished an individual shall be regarded as a payment to the individual, and adjustment shall be

§405.350