Engineer, Airframe Branch, ANM–120S, FAA, Seattle Aircraft Certification Office, 1601 Lind Avenue, SW., Renton, Washington 98057–3356; phone: (425) 917–6428; fax: (425) 917–6590; e-mail: nathan.p.weigand@faa.gov.

(k) For service information identified in this AD, contact Boeing Commercial Airplanes, Attention: Data & Services Management, P.O. Box 3707, MC 2H–65, Seattle, WA 98124–2207; telephone 206–544–5000, extension 1; fax 206–766–5680; e-mail me.boeing@boeing.com; Internet https://www.myboeingfleet.com. You may review copies of the referenced service information at the FAA, Transport Airplane Directorate, 1601 Lind Avenue, SW., Renton, Washington. For information on the availability of this material at the FAA, call 425–227–1221.

Issued in Renton, Washington, on August 25, 2011.

Ali Bahrami,
Manager, Transport Airplane Directorate, Aircraft Certification Service.

[FR Doc. 2011–22371 Filed 8–31–11; 8:45 am]
BILLING CODE 4910–13–P

DEPARTMENT OF THE INTERIOR

National Indian Gaming Commission

25 CFR Chapter III

Regulatory Review Schedule

AGENCY: National Indian Gaming Commission.

ACTION: Notice of cancellation of consultation meeting.

SUMMARY: The purpose of this document is to cancel ten tribal consultations scheduled during November 2011, December 2011, January 2012, and February 2012 and to modify the dates for six tribal consultations scheduled during September 2011, October 2011 and November 2011.

DATES: See SUPPLEMENTARY INFORMATION below for dates and locations of cancelled consultations.

FOR FURTHER INFORMATION CONTACT: Lael Echo-Hawk, National Indian Gaming Commission, 1441 L Street, NW., Suite 9100, Washington, DC 20005. Telephone: 202–632–7003; e-mail: reg.review@nigc.gov.

SUPPLEMENTARY INFORMATION: On November 18, 2010, the National Indian Gaming Commission (NICG) issued a Notice of Inquiry and Notice of Consultation advising the public that it was conducting a review of its regulations promulgated to implement 25 U.S.C. 2701–2721 of the Indian Gaming Regulatory Act (IGRA) and requesting public comment on the process for conducting the regulatory review. On April 4, 2011, after holding eight consultations and reviewing all comments, NICG published a Notice of Regulatory Review Schedule in the Federal Register setting out consultation schedules and review processes. (76 FR 18457, April 4, 2011).

The Commission’s regulatory review process established a tribal consultation schedule with a description of the regulation groups to be covered during consultation. Group 1 included a review of:

(a) A Buy Indian Act regulation;

(b) Part 523—Review and Approval of Existing Ordinances or Resolutions;

(c) Part 514—Fees;

(d) Part 559—Facility License Notifications, Renewals, and Submissions; and

(e) Part 542—Minimum Internal Control Standards.

Group 2 included a review of:

(a) Part 573—Enforcement; and

(b) Regulations concerning proceedings before the Commission, including: Parts 519—Service, Part 524—Appeals, Part 539—Appeals, and

will be received through October 26, 2011.

ADDRESSES: You may submit comments, identified by docket ID number HHS–OPHS–2011–0005, by one of the following methods:

• Federal eRulemaking Portal: http://www.regulations.gov. Enter the above docket ID number in the “Enter Keyword or ID” field and click on “Search.” On the next web page, click on “Submit a Comment” action and follow the instructions.

• Mail/Hand delivery/Courier [For paper, disk, or CD–ROM submissions]

  to: Jerry Menikoff, M.D., J.D., OHRP, 1101 Wootton Parkway, Suite 200, Rockville, MD 20852.

  Comments received, including any personal information, will be posted without change to http://www.regulations.gov.

FOR FURTHER INFORMATION CONTACT: Jerry Menikoff, M.D., J.D., Office for Human Research Protections (OHRP), Department of Health and Human Services, 1101 Wootton Parkway, Suite 200, Rockville, MD 20852.

SUPPLEMENTARY INFORMATION: The ANPRM was published in the Federal Register on July 26, 2011 (Volume 76, Number 143, page 44512) with a deadline for comments of September 26, 2011. The ANPRM requests comments on how current regulations for protecting human subjects who participate in research might be modernized and revised to be more effective and how to better protect human subjects who are involved in research, while facilitating valuable research and reducing burden, delay, and ambiguity for investigators. Since the ANPRM was published the Department has received requests to extend the comment period to allow sufficient time for a full review of the ANPRM. HHS and OSTP are committed to affording the public a meaningful opportunity to comment on the ANPRM and welcome comments.

Dated: August 26, 2011.

Kathleen Sebelius,
Secretary.

[FR Doc. 2011–22341 Filed 8–31–11; 8:45 am]
BILLING CODE 4150–28–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Office of the Secretary

45 CFR Parts 46, 160, and 164

Food and Drug Administration

21 CFR Parts 50 and 56

Human Subjects Research Protections: Enhancing Protections for Research Subjects and Reducing Burden, Delay, and Ambiguity for Investigators; Extension of Comment Period

AGENCIES: The Office of the Secretary, HHS, and the Food and Drug Administration, HHS.

ACTION: Advance notice of proposed rulemaking; extension of comment period.

SUMMARY: The Office of the Secretary of the Department of Health and Human Services (HHS) in coordination with the Office of Science and Technology Policy (OSTP) is extending the comment period for an advance notice of proposed rulemaking (ANPRM) requesting comment on how current regulations for protecting human subjects who participate in research might be modernized and revised to be more effective. That ANPRM was published in the Federal Register on July 26, 2011.

DATES: The comment period for the proposed rule published July 26, 2011, at 76 FR 44512 is extended. Comments received, including any personal information, will be posted without change to http://www.regulations.gov.

FOR FURTHER INFORMATION CONTACT: Jerry Menikoff, M.D., J.D., Office for Human Research Protections (OHRP), Department of Health and Human Services, 1101 Wootton Parkway, Suite 200, Rockville, MD 20852.

SUPPLEMENTARY INFORMATION: The ANPRM was published in the Federal Register on July 26, 2011 (Volume 76, Number 143, page 44512) with a deadline for comments of September 26, 2011. The ANPRM requests comments on how current regulations for protecting human subjects who participate in research might be modernized and revised to be more effective and how to better protect human subjects who are involved in research, while facilitating valuable research and reducing burden, delay, and ambiguity for investigators. Since the ANPRM was published the Department has received requests to extend the comment period to allow sufficient time for a full review of the ANPRM. HHS and OSTP are committed to affording the public a meaningful opportunity to comment on the ANPRM and welcome comments.

Dated: August 26, 2011.

Kathleen Sebelius, Secretary.