DEPARTMENT OF HEALTH AND HUMAN SERVICES

Office of the Secretary

21 CFR Ch. I

25 CFR Ch. V

42 CFR Chs. I–V

45 CFR Subtitle A; Subtitle B, Chs. II, III, and XIII

Regulatory Agenda

AGENCY: Office of the Secretary, HHS. **ACTION:** Semiannual regulatory agenda.

SUMMARY: The Regulatory Flexibility Act of 1980 and Executive Order 12866 require the Department semiannually to issue an inventory of rulemaking actions under development to provide the public a summary of forthcoming regulatory actions. This information will help the public more effectively participate in the Department's regulatory activity, and the Department welcomes comments on any aspect of this agenda.

FOR FURTHER INFORMATION CONTACT:

Jennifer M. Cannistra, Executive Secretary, Department of Health and Human Services, Washington, DC 20201.

SUPPLEMENTARY INFORMATION: The Department of Health and Human

Services (HHS) is the Federal Government's principal agency for protecting the health of all Americans and providing essential human services, especially for those who are least able to help themselves. HHS enhances the health and well-being of Americans by promoting effective health and human services and by fostering sound, sustained advances in the sciences underlying medicine, public health, and social services. This agenda presents the rulemaking activities that the Department expects to undertake in the foreseeable future to advance this mission. The agenda furthers several Departmental goals, including strengthening health care; advancing scientific knowledge and innovation; advancing the health, safety, and wellbeing of the American people; increasing efficiency, transparency, and accountability of HHS programs; and strengthening the Nation's health and human services infrastructure and workforce.

HHS has an agency-wide effort to support the agenda's purpose of encouraging more effective public participation in the regulatory process. The Department's Public Participation Task Force, which was created as part of the HHS Retrospective Review plan in response to Executive Order 13563 (*Improving Regulation and Regulatory Review*), regularly meets to identify ways to make the rulemaking process

more accessible to the general public. For example, to encourage public participation, HHS regularly updates its main regulatory Web page (http:// www.HHS.gov/regulations/), which includes links to HHS rules currently open for public comment and provides a "regulations toolkit" with background information on regulations, the commenting process, how public comments influence the development of a rule, and how the public can provide effective comments. HHS also actively encourages meaningful public participation in its retrospective review of regulations, including through a comment form on the HHS retrospective review Web page (http://www.HHS.gov/ RetrospectiveReview). In addition, a cross-agency team at HHS is currently considering how to increase efficiency in rulemaking by organizing public comment on proposed rules.

The rulemaking abstracts included in this paper issue of the **Federal Register** only cover, as required by the Regulatory Flexibility Act of 1980, those prospective HHS rulemakings likely to have a significant economic impact on a substantial number of small entities. The Department's complete Regulatory Agenda is accessible online at *http:// www.RegInfo.gov.*

Dated: August 21, 2013.

Jennifer M. Cannistra,

Executive Secretary to the Department.

FOOD AND DRUG ADMINISTRATION—PRERULE STAGE

Sequence No.	Title	Regulation Identifier No.
		0910–AF43 0910–AG14

FOOD AND DRUG ADMINISTRATION—PROPOSED RULE STAGE

Sequence No.	Title	Regulation Identifier No.
276	Food Labeling; Revision of the Nutrition and Supplement Facts Labels (Reg Plan Seq No. 49)	0910-AF22
277	Food Labeling: Serving Sizes of Foods That Can Reasonably Be Consumed at One-Eating Occasion;	0910-AF23
	Dual-Column Labeling; Updating, Modifying, and Establishing Certain RACCs (Reg Plan Seq No. 50).	
278	Over-the-Counter (OTC) Drug Review—Cough/Cold (Antihistamine) Products	0910–AF31
279	Over-the-Counter (OTC) Drug Review—Internal Analgesic Products	0910–AF36
280	Updated Standards for Labeling of Pet Food	0910–AG09
281	Current Good Manufacturing Practice, Hazard Analysis, and Risk-Based Preventive Controls for Food for	0910–AG10
	Animals (Reg Plan Seq No. 51).	
282	Over-the-Counter (OTC) Drug Review—Pediatric Dosing for Cough/Cold Products	0910–AG12
283	Electronic Distribution of Prescribing Information for Human Prescription Drugs Including Biological Prod- ucts.	0910–AG18
284	Produce Safety Regulation	0910–AG35
285	Produce Safety Regulation Hazard Analysis and Risk-Based Preventive Controls	0910-AG36
286	"Tobacco Products" Subject to the Federal Food, Drug, and Cosmetic Act, as Amended by the Family Smoking Prevention and Tobacco Control Act (Reg Plan Seq No. 52).	0910–AG38
287	Requirements for the Testing and Reporting of Tobacco Product Constituents, Ingredients, and Additives	0910–AG59
288	Foreign Supplier Verification Program	0910–AG64
289	Supplemental Applications Proposing Labeling Changes for Approved Drugs and Biological Products (Reg Plan Seq No. 55).	0910–AG94

FOOD AND DRUG ADMINISTRATION—PROPOSED RULE STAGE—Continued

Sequence No.	Title	Regulation Identifier No.
291 292	Veterinary Feed Directive (Reg Plan Seq No. 56) Format and Content of Reports Intended to Demonstrate Substantial Equivalence Radiology Devices; Designation of Special Controls for the Computed Tomography X-Ray System Mammography Quality Standards Act; Regulatory Amendments	

References in boldface appear in The Regulatory Plan in part II of this issue of the Federal Register.

FOOD AND DRUG ADMINISTRATION—FINAL RULE STAGE

Sequence No.	Title	Regulation Identifier No.
294	Content and Format of Labeling for Human Prescription Drugs and Biologics; Requirements for Preg- nancy and Lactation Labeling.	0910–AF11
295	Infant Formula: Current Good Manufacturing Practices; Quality Control Procedures; Notification Require- ments; Records and Reports; and Quality Factors.	0910-AF27
296	Combinations of Bronchodilators With Nasal Decongestants or Expectorants; Cold, Cough, Allergy, Bron- chodilator, and Antiasthmatic Drug Products for Over-the-Counter Human Use.	0910-AF33
297	Laser Products; Proposed Amendment to Performance Standard	0910–AF87
298	Food Labeling: Calorie Labeling of Articles of Food Sold in Vending Machines (Reg Plan Seq No. 57)	0910–AG56
299	Food Labeling: Nutrition Labeling of Standard Menu Items in Restaurants and Similar Retail Food Estab- lishments (Reg Plan Seq No. 58).	0910–AG57
300	Use of Certain Symbols in Labeling	0910–AG74
301	Requirements for the Submission of Data Needed To Calculate User Fees for Manufacturers and Import- ers of Tobacco Products.	0910–AG81

References in boldface appear in The Regulatory Plan in part II of this issue of the Federal Register.

FOOD AND DRUG ADMINISTRATION—LONG-TERM ACTIONS

Sequence No.	Title	Regulation Identifier No.
302 303	Over-the-Counter (OTC) Drug Review—Topical Antimicrobial Drug Products Amendment to the Current Good Manufacturing Practice Regulations for Finished Pharmaceuticals—Sec- ond Phase.	0910–AF69 0910–AG20
304 305	Human Subject Protection; Acceptance of Data From Clinical Studies for Medical Devices Amendments to the Current Good Manufacturing Practice Regulations for Finished Pharmaceuticals— Components.	0910–AG48 0910–AG70

FOOD AND DRUG ADMINISTRATION-COMPLETED ACTIONS

Sequence No.	Title	Regulation Identifier No.
306 307		0910–AG31 0910–AG82
308		0910–AG84

CENTERS FOR MEDICARE & MEDICAID SERVICES—PROPOSED RULE STAGE

Sequence No.	Title	Regulation Identifier No.
309	Emergency Preparedness Requirements for Medicare and Medicaid Participating Providers and Suppliers (CMS-3178-P) (Section 610 Review).	0938–AO91
310	Prospective Payment System for Federally Qualified Health Centers; Changes to Contracting Policies for Rural Health Clinics and CLIA Enforcement Actions for Proficiency Testing Referral (CMS–1443–F) (Section 610 Review).	0938–AR62
311	Hospital Inpatient Prospective Payment System for Acute Care Hospitals and the Long-Term Care Hospital Prospective Payment System and Fiscal Year 2015 Rates (CMS-1607-P) (Reg Plan Seq No. 62).	0938–AS11
312	CY 2015 Revisions to Payment Policies under the Physician Fee Schedule and Other Revisions to Medi- care Part B (CMS-1612-P) (Reg Plan Seq No. 63).	0938–AS12
313	CY 2015 Hospital Outpatient Prospective Payment System (PPS) Policy Changes and Payment Rates, and CY 2015 Ambulatory Surgical Center Payment System Policy Changes and Payment Rates (CMS– 1613–P) (Reg Plan Seq No. 64).	0938–AS15

References in boldface appear in The Regulatory Plan in part II of this issue of the Federal Register.

CENTERS FOR MEDICARE & MEDICAID SERVICES—FINAL RULE STAGE

Sequence No.	Title	Regulation Identifier No.
314 315	Covered Outpatient Drugs (CMS–2345–F) (Section 610 Review) CY 2014 Changes to the End-Stage Renal Disease (ESRD) Prospective Payment System, ESRD Quality Incentive Program, and Durable Medical Equipment (CMS–1526–F).	0938–AQ41 0938–AR55
316	Revisions to Payment Policies Under the Physician Fee Schedule and Medicare Part B for CY 2014 (CMS-1600-F).	0938–AR56
317	Adoption of Operating Rules for HIPAA Transactions (CMS-0036-IFC)	0938–AS01

CENTERS FOR MEDICARE & MEDICAID SERVICES—COMPLETED ACTIONS

Sequence No.	Title	Regulation Identifier No.
318	Changes to the Hospital Inpatient and Long-Term Care Prospective Payment System for FY 2014 (CMS- 1599-F).	0938–AR53
319		0938–AR54

DEPARTMENT OF HEALTH AND HUMAN SERVICES (HHS)

Food and Drug Administration (FDA)

Prerule Stage

274. Over-the-Counter (OTC) Drug Review—Sunscreen Products

Legal Authority: 21 U.S.C. 321p; 21 U.S.C. 331; 21 U.S.C. 351 to 353; 21 U.S.C. 355; 21 U.S.C. 360; 21 U.S.C. 371

Abstract: The OTC drug review establishes conditions under which OTC drugs are considered generally recognized as safe and effective and not misbranded. After a final monograph (i.e., final rule) is issued, only OTC drugs meeting the conditions of the monograph, or having an approved new drug application, may be legally marketed. The first of the future actions will address the safety of sunscreen active ingredients.

Timetable:

Action	Date	FR Cite
ANPRM (Sun- screen and In- sect Repellent).	02/22/07	72 FR 7941
ANPRM Comment Period End.	05/23/07	
NPRM (UVA/ UVB).	08/27/07	72 FR 49070
NPRM Comment Period End.	12/26/07	
Final Action (UVA/ UVB).	06/17/11	76 FR 35620
NPRM (Effective- ness).	06/17/11	76 FR 35672
NPRM (Effective- ness) Comment Period End.	09/15/11	
ANPRM (Dosage Forms).	06/17/11	76 FR 35669
ANPRM (Dosage Forms) Com- ment Period End.	09/15/11	

Action	Date	FR Cite
ANPRM (Safety)	06/00/14	

Regulatory Flexibility Analysis

Required: Yes. Agency Contact: David Eng, Regulatory Project Manager, Department of Health and Human Services, Food and Drug Administration, Center for Drug Evaluation and Research, WO 22, Room 5487, 10903 New Hampshire Avenue, Silver Spring, MD 20993, Phone: 301 796–2773, Fax: 301 796– 9899, Email: david.eng@fda.hhs.gov. RIN: 0910–AF43

275. Prescription Drug Marketing Act of 1987; Prescription Drug Amendments of 1992; Policies, Requirements, and Administrative Procedures (Section 610 Review)

Legal Authority: 21 U.S.C. 331; 21 U.S.C. 333; 21 U.S.C. 351 to 353; 21 U.S.C. 360; 21 U.S.C. 371; 21 U.S.C. 374; 21 U.S.C. 381

Abstract: FDA is currently reviewing regulations promulgated under the Prescription Drug Marketing Act (PDMA). FDA is undertaking this review to determine whether the regulations should be changed or rescinded to minimize adverse impacts on a substantial number of small entities. FDA has extended again the completion date by 1 year and will complete the review by November 2013. *Timetable*.

Timetable.		
Action	Date	FR Cite
Begin Review of Current Regula- tion.	11/24/08	
End Review of Current Regula- tion.	11/00/13	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Howard Muller, Office of Regulatory Policy, Department of Health and Human Services, Food and Drug Administration, Center for Drug Evaluation and Research, WO 51, Room 6234, 10903 New Hampshire Avenue, Silver Spring, MD 20993–0002, *Phone:* 301 796–3601, *Fax:* 301 847– 8440, *Email: pdma610(c)review@ fda.hhs.gov.*

RIN: 0910-AG14

DEPARTMENT OF HEALTH AND HUMAN SERVICES (HHS)

Food and Drug Administration (FDA)

Proposed Rule Stage

276. Food Labeling; Revision of the Nutrition and Supplement Facts Labels

Regulatory Plan: This entry is Seq. No. 49 in part II of this issue of the **Federal Register**. *RIN:* 0910–AF22

277. Food Labeling: Serving Sizes of Foods That Can Reasonably Be Consumed at One-Eating Occasion; Dual-Column Labeling; Updating, Modifying, and Establishing Certain RACCs

Regulatory Plan: This entry is Seq. No. 50 in part II of this issue of the **Federal Register**.

RIN: 0910–AF23

278. Over-the-Counter (OTC) Drug Review—Cough/Cold (Antihistamine) Products

Legal Authority: 21 U.S.C. 321p; 21 U.S.C. 331; 21 U.S.C. 351 to 353; 21 U.S.C. 355; 21 U.S.C. 360; 21 U.S.C. 371 *Abstract:* FDA will be proposing a rule to add the common cold indication to certain over-the-counter (OTC) antihistamine active ingredients. This proposed rule is the result of collaboration under the U.S.-Canada Regulatory Cooperation Council (RCC) as part of efforts to reduce unnecessary duplication and differences. This pilot exercise will help determine the feasibility of developing an ongoing mechanism for alignment in review and adoption of OTC drug monograph elements.

Timetable:

Action	Date	FR Cite
Reopening of Ad- ministrative Record.	08/25/00	65 FR 51780
Comment Period End.	11/24/00	
NPRM (Amend- ment) (Common Cold).	12/00/13	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Janice Adams–King, Regulatory Health Project Manager, Department of Health and Human Services, Food and Drug Administration, Center for Drug Evaluation and Research, WO 22, Room 5416, 10903 New Hampshire Avenue, Silver Spring, MD 20993, Phone: 301 796–3713, Fax: 301 796–9899, Email: janice.adams-king@fda.hhs.gov. RIN: 0910–AF31

279. Over-the-Counter (OTC) Drug Review—Internal Analgesic Products

Legal Authority: 21 U.S.C. 321p; 21 U.S.C. 331; 21 U.S.C. 351 to 353; 21 U.S.C. 355; 21 U.S.C. 360; 21 U.S.C. 371; 21 U.S.C. 374; 21 U.S.C. 379e

Abstract: The OTC drug review establishes conditions under which OTC drugs are considered generally recognized as safe and effective and not misbranded. After a final monograph (i.e., final rule) is issued, only OTC drugs meeting the conditions of the monograph, or having an approved new drug application, may be legally marketed. The first action addresses acetaminophen safety. The second action addresses products marketed for children under 2 years old and weightand age-based dosing for children's products.

Timetable:

Action	Date	FR Cite
NPRM (Amend- ment) (Required Warnings and Other Labeling).	12/26/06	71 FR 77314

Action	Date	FR Cite
NPRM Comment Period End.	05/25/07	
Final Action (Re- quired Warn- ings and Other Labeling).	04/29/09	74 FR 19385
Final Action (Cor- rection).	06/30/09	74 FR 31177
Final Action (Technical Amendment).	11/25/09	74 FR 61512
NPRM (Amend- ment) (Pedi- atric).	07/00/14	
NPRM (Amend- ment) (Acetami- nophen).	12/00/14	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Janice Adams–King, Regulatory Health Project Manager, Department of Health and Human Services, Food and Drug Administration, Center for Drug Evaluation and Research, WO 22, Room 5416, 10903 New Hampshire Avenue, Silver Spring, MD 20993, Phone: 301 796–3713, Fax: 301 796–9899, Email: janice.adams-king@fda.hhs.gov.

RIN: 0910-AF36

280. Updated Standards for Labeling of Pet Food

Legal Authority: 21 U.S.C. 343; 21 U.S.C. 371; Pub. L. 110–85, sec 1002(a)(3)

Abstract: FDA is proposing updated standards for the labeling of pet food that include nutritional and ingredient information, as well as style and formatting standards. FDA is taking this action to provide pet owners and animal health professionals more complete and useful information about the nutrient content and ingredient composition of pet food products.

Timetable:

Action	Date	FR Cite
NPRM	06/00/14	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: William Burkholder, Veterinary Medical Officer, Department of Health and Human Services, Food and Drug Administration, Center for Veterinary Medicine, Room 2642 (MPN– 4, HFV–228), 7519 Standish Place, Rockville, MD 20855, *Phone:* 240 453– 6865, *Email: william.burkholder@ fda.hhs.gov.*

RIN: 0910-AG09

281. Current Good Manufacturing Practice, Hazard Analysis, and Risk-Based Preventive Controls for Food for Animals

Regulatory Plan: This entry is Seq. No. 51 in part II of this issue of the **Federal Register**. *RIN:* 0910–AG10

282. Over-the-Counter (OTC) Drug Review—Pediatric Dosing for Cough/ Cold Products

Legal Authority: 21 U.S.C. 331; 21 U.S.C. 351 to 353; 21 U.S.C. 355; 21 U.S.C. 360; 21 U.S.C. 371

Abstract: The OTC drug review establishes conditions under which OTC drugs are considered generally recognized as safe and effective and not misbranded. After a final monograph (i.e., final rule) is issued, only OTC drugs meeting the conditions of the monograph, or having an approved new drug application, may be legally marketed. This action will propose changes to the final monograph to address safety and efficacy issues associated with pediatric cough and cold products.

Timetable:

Action	Date	FR Cite
NPRM	06/00/14	

Regulatory Flexibility Analysis Required: Yes.

Ågency Contact: Janice Adams–King, Regulatory Health Project Manager, Department of Health and Human Services, Food and Drug Administration, Center for Drug Evaluation and Research, WO 22, Room 5416, 10903 New Hampshire Avenue, Silver Spring, MD 20993, *Phone:* 301 796–3713, *Fax:* 301 796–9899, *Email: janice.adams-king@fda.hhs.gov. RIN:* 0910–AG12

283. Electronic Distribution of Prescribing Information for Human Prescription Drugs Including Biological Products

Legal Authority: 21 U.S.C. 321; 21 U.S.C. 331; 21 U.S.C. 351; 21 U.S.C. 352; 21 U.S.C. 353; 21 U.S.C. 355; 21 U.S.C. 358; 21 U.S.C. 360; 21 U.S.C. 360b; 21 U.S.C. 360gg to 360ss; 21 U.S.C. 371; 21 U.S.C. 374; 21 U.S.C. 379e; 42 U.S.C. 216; 42 U.S.C. 241; 42 U.S.C. 262; 42 U.S.C. 264

Abstract: This rule would require electronic package inserts for human drug and biological prescription products with limited exceptions, in lieu of paper, which is currently used. These inserts contain prescribing information intended for healthcare practitioners. This would ensure that the information accompanying the product is the most up-to-date information regarding important safety and efficacy issues about these products.

Timetable:

Action	Date	FR Cite
NPRM	01/00/14	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Megan Velez, Policy Analyst, Department of Health and Human Services, Food and Drug Administration, Office of Policy, WO 32, Room 4249, 10903 New Hampshire Avenue, Silver Spring, MD 20993, Phone: 301 796–9301, Email: megan.velez@fda.hhs.gov.

RIN: 0910-AG18

284. Produce Safety Regulation

Legal Authority: 21 U.S.C. 342; 21 U.S.C. 350h; 21 U.S.C. 371; 42 U.S.C. 264; Pub. L. 111-353 (signed on Jan. 4, 2011)

Abstract: FDA is proposing to establish science-based minimum standards for the safe production and harvesting of those types of fruits and vegetables that are raw agricultural commodities for which the Secretary has determined that such standards minimize the risk of serious adverse health consequences or death. The purpose of the proposed rule is to reduce the risk of illness associated with fresh produce.

Timetable:

Action	Date	FR Cite
NPRM	01/16/13	78 FR 3503
NPRM Comment Period End.	05/16/13	
NPRM Comment Period Ex- tended.	04/26/13	78 FR 24692
NPRM Comment Period Ex- tended End.	09/16/13	
NPRM Comment Period Ex- tended.	08/09/13	78 FR 48637
NPRM Comment Period Ex- tended End.	11/15/13	
Notice of Intent To Prepare an Environmental Impact State- ment for the Proposed Rule.	08/19/13	78 FR 50358

Action	Date	FR Cite	Action	Date
Notice of Intent To Prepare En- vironmental Im-	11/15/13		NPRM Comment Period Ex- tended.	08/09/13
pact Statement for the Pro- posed Rule			NPRM Comment Period Ex- tended End.	11/15/13
Comment Pe- riod End.	11/00/110		NPRM Comment Period Ex-	11/20/13
NPRM Comment Period Ex- tended.	11/20/13	78 FR 69605	tended. NPRM Comment Period Ex-	11/22/13
NPRM Comment Period Ex- tended End.	11/22/13		tended End.	vihility And
Environmental Im- pact Statement for the Pro- posed Rule; Comment Pe- riod Extended.	11/18/13	78 FR 69006	Required: Yes. Agency Contac Advisor, Departn Human Services, Administration, (t: Jenny Sc nent of Hea Food and I
Environmental Im- pact Statement for the Pro- posed Rule;	03/14/14		5100 Paint Branc Park, MD 20740, Email: jenny.scot RIN: 0910–AG	h Parkway, <i>Phone:</i> 240 t@fda.hhs.g
Comment Pe- riod Extended End.			286. "Tobacco Pı Federal Food, Dr	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Samir Assar, Supervisory Consumer Safety Officer, Department of Health and Human Services, Food and Drug Administration, Center for Food Safety and Applied Nutrition, Office of Food Safety, 5100 Paint Branch Parkway, College Park, MD 20740, Phone: 240 402-1636, Email: samir.assar@ fda.hhs.gov.

RIN: 0910-AG35

285. Hazard Analysis and Risk-Based **Preventive Controls**

Legal Authority: 21 U.S.C. 342; 21 U.S.C. 371; 42 U.S.C. 264; Pub. L. 111-353 (signed on Jan. 4, 2011)

Abstract: This proposed rule would require a food facility to have and implement preventive controls to significantly minimize or prevent the occurrence of hazards that could affect food manufactured, processed, packed, or held by the facility. This action is intended to prevent or, at a minimum, quickly identify foodborne pathogens before they get into the food supply. Timetable:

Action	Date	FR Cite
NPRM NPRM Comment Period End.	01/16/13 05/16/13	78 FR 3646
NPRM Comment Period Ex- tended.	04/26/13	78 FR 24691
NPRM Comment Period Ex- tended End.	09/16/13	

bility Analysis Jenny Scott, Senior nt of Health and

ood and Drug fice of Food Safety, Parkway, College hone: 240 402–1488, fda.hhs.gov.

FR Cite

78 FR 48636

78 FR 69604

ducts" Subject to the , and Cosmetic Act, as Amended by the Family Smoking Prevention and Tobacco Control Act

Regulatory Plan: This entry is Seq. No. 52 in part II of this issue of the Federal Register. RIN: 0910-AG38

287. Requirements for the Testing and **Reporting of Tobacco Product** Constituents, Ingredients, and Additives

Legal Authority: 21 U.S.C. 301 et seq.; 21 U.S.C. 387; The Family Smoking Prevention and Tobacco Control Act

Abstract: The Federal Food, Drug, and Cosmetic Act, as amended by the Family Smoking Prevention and Tobacco Control Act, requires the Food and Drug Administration to promulgate regulations that require the testing and reporting of tobacco product constituents, ingredients, and additives, including smoke constituents, that the agency determines should be tested to protect the public health.

Timetable:

Action	Date	FR Cite
NPRM	12/00/13	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Carol Drew, Regulatory Counsel, Department of Health and Human Services, Food and Drug Administration, Center for Tobacco Products, 9200 Corporate Boulevard, Room 240 H, Rockville, MD 20850, Phone: 877 287-1373, Fax: 240 276-3904, Email: carol.drew@ fda.hhs.gov.

RIN: 0910-AG59

288. Foreign Supplier Verification Program

Legal Authority: 21 U.S.C. 384a; title III, sec 301 of FDA Food Safety Modernization Act, Pub. L. 111-353, establishing sec 805 of the Federal Food, Drug, and Cosmetic Act (FD&C Act)

Abstract: FDA is proposing regulations that describe what a food importer must do to verify that its foreign suppliers produce food that is as safe as food produced in the United States. FDA is taking this action to improve the safety of food that is imported into the United States.

Timetable:

Action	Date	FR Cite
NPRM NPRM Comment Period End.	07/29/13 11/26/13	78 FR 45729
NPRM Comment Period Ex- tended.	11/20/13	78 FR 69602
NPRM Comment Period Ex- tended End.	01/27/14	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Brian L. Pendleton, Senior Policy Advisor, Department of Health and Human Services, Food and Drug Administration, Office of Policy, WO 32, Room 4245, 10903 New Hampshire Avenue, Silver Spring, MD 20993-0002, Phone: 301 796-4614, Fax: 301 847-8616, Email: brian.pendleton@ fda.hhs.gov.

RIN: 0910-AG64

289. Supplemental Applications Proposing Labeling Changes for Approved Drugs and Biological Products

Regulatory Plan: This entry is Seq. No. 55 in part II of this issue of the Federal Register.

RIN: 0910-AG94

290. Veterinary Feed Directive

Regulatory Plan: This entry is Seq. No. 56 in part II of this issue of the Federal Register. RIN: 0910-AG95

291. Format and Content of Reports **Intended To Demonstrate Substantial** Equivalence

Legal Authority: 21 U.S.C. 387e(j); 21 U.S.C. 387j(a); secs 905(j) and 910(a) of the Federal Food, Drug, and Cosmetic Act

Abstract: This regulation would establish the format and content of reports intended to demonstrate substantial equivalence and compliance

with the FD&C Act (sections 905(j) and 910(a) of the FD&C Act). This regulation also would provide information as to how the Agency will review and act on these submissions.

Timetable:

Action	Date	FR Cite
NPRM	03/00/14	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Gerie Voss, Regulatory Counsel, Department of Health and Human Services, Food and Drug Administration, Center for Tobacco Products, 9200 Corporate Boulevard, Rockville, MD 20850, Phone: 877 287-1373, Fax: 240 276-4193, Email: gerie.voss@fda.hhs.gov. RIN: 0910-AG96

292. Radiology Devices; Designation of **Special Controls for the Computed Tomography X-Ray System**

Legal Authority: 21 U.S.C. 360 Abstract: The proposed rule would establish special controls for the computed tomography (CT) X-ray system, a class II device as defined in 21 CFR 892.1750. A CT X-ray system is a diagnostic X-ray imaging system intended to produce cross-sectional images of the body through use of a computer to reconstruct an image from the same axial plane taken at different angles. High doses of ionizing radiation can cause acute (deterministic) effects such as burns, reddening of the skin, cataracts, hair loss, sterility, or, in extremely high doses, radiation poisoning. Therefore, the design of a CT X-ray system needs to balance the benefits of the device (i.e., the ability of the device to produce a diagnostic quality image) with the known risks (e.g., exposure to ionizing radiation). FDA is establishing special controls, combined with the general controls, to provide reasonable assurance of the safety and effectiveness of a class II CT X-ray system.

Timetable:

Action	Date	FR Cite
NPRM	05/00/14	

Regulatory Flexibility Analysis Required: Yes.

Âgency Contact: Erica Blake, Regulatory Counsel, Department of Health and Human Services, Food and Drug Administration, Center for Devices and Radiological Health, WO 66, Room 4426, 10903 New Hampshire Avenue, Silver Spring, MD 20993, Phone: 301 796-6248, Fax: 301 847-8145, Email: erica.blake@fda.hhs.gov.

RIN: 0910-AH03

293. Mammography Quality Standards Act; Regulatory Amendments

Legal Authority: 21 U.S.C. 360i; 21 U.S.C. 360nn; 21 U.S.C. 374(e); 42 U.S.C. 263b

Abstract: FDA is proposing to amend its regulations governing mammography. The amendments would update the regulations issued under the Mammography Quality Standards Act of 1992 (MQSA). FDA is taking this action to address changes in mammography technology and mammography processes, such as breast density reporting, that have occurred since the regulations were published in 1997. *Timetable:*

Action	Date	FR Cite
NPRM	12/00/13	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Nancy Pirt, Regulatory Counsel, Department of Health and Human Services, Food and Drug Administration, Center for Devices and Radiological Health, WO 66, Room 4438, 10903 New Hampshire Avenue, Silver Spring, MD 20993, Phone: 301 796-6248, Fax: 301 847-8145, Email: nancy.pirt@fda.hhs.gov.

RIN: 0910-AH04

DEPARTMENT OF HEALTH AND **HUMAN SERVICES (HHS)**

Food and Drug Administration (FDA)

Final Rule Stage

294. Content and Format of Labeling for **Human Prescription Drugs and Biologics; Requirements for Pregnancy** and Lactation Labeling

Legal Authority: 21 U.S.C. 321; 21 U.S.C. 331; 21 U.S.C. 351 to 353; 21 U.S.C. 355; 21 U.S.C. 358; 21 U.S.C. 360; 21 U.S.C. 360b; 21 U.S.C. 360gg to 360ss; 21 U.S.C. 371; 21 U.S.C. 374; 21 U.S.C. 379e; 42 U.S.C. 216; 42 U.S.C. 241; 42 U.S.C. 262; 42 U.S.C. 264

Abstract: This final rule will amend the content and format of the "Pregnancy," "Labor and delivery," and "Nursing mothers" subsections of the "Use in Specific Populations" section of regulations regarding the labeling for human prescription drug and biological products (21 CFR 201.56 and 201.57) to better communicate risks.

Timetable:

Action	Date	FR Cite
NPRM	05/29/08	73 FR 30831

Action	Date	FR Cite
NPRM Comment Period End.	08/27/08	
Final Action	05/00/14	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Molly Flannery, Regulatory Counsel, Department of Health and Human Services, Food and Drug Administration, Center for Drug Evaluation and Research, WO 51, Room 6246, 10903 New Hampshire Avenue, Silver Spring, MD 20993, Phone: 301 796-3543, Email: molly.flannery@ fda.hhs.gov.

RIN: 0910-AF11

295. Infant Formula: Current Good Manufacturing Practices; Quality **Control Procedures; Notification Requirements; Records and Reports;** and Quality Factors

Legal Authority: 21 U.S.C. 321; 21 U.S.C. 342; 21 U.S.C. 350a; 21 U.S.C. 371

Abstract: The Food and Drug Administration (FDA) is revising its infant formula regulations in 21 CFR parts 106 and 107 to establish requirements for current good manufacturing practices (CGMP), including audits; to establish requirements for quality factors; and to amend FDA's quality control procedures, notification, and record and reporting requirements for infant formula. FDA is taking this action to improve the protection of infants who consume infant formula products.

Timetable:

Action	Date	FR Cite
NPRM	07/09/96	61 FR 36154
NPRM Comment Period End.	12/06/96	
NPRM Comment	04/28/03	68 FR 22341
Period Re- opened.		
NPRM Comment	06/27/03	68 FR 38247
Period Ex- tended.		
NPRM Comment	08/26/03	
Period End. NPRM Comment	08/01/06	71 FR 43392
Period Re-		
opened. NPRM Comment	09/15/06	
Period End.		
Final Rule	11/00/13	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Benson Silverman, Staff Director, Infant Formula and Medical Foods, Department of Health and Human Services, Food and Drug Administration, Center for Food Safety and Applied Nutrition (HFS-850), 5100 Paint Branch Parkway, College Park, MD 20740, Phone: 240 402-1459, Email: benson.silverman@fda.hhs.gov RIN: 0910-AF27

296. Combinations of Bronchodilators With Nasal Decongestants or Expectorants; Cold, Cough, Allergy, **Bronchodilator, and Antiasthmatic Drug Products for Over-the-Counter** Human Use

Legal Authority: 21 U.S.C. 321p; 21 U.S.C. 331; 21 U.S.C. 351 to 353; 21 U.S.C. 355; 21 U.S.C. 360; 21 U.S.C. 371

Abstract: The OTC drug review establishes conditions under which OTC drugs are considered generally recognized as safe and effective and not misbranded. After a final monograph (i.e., final rule) is issued, only OTC drugs meeting the conditions of the monograph, or having an approved new drug application, may be legally marketed. This action addresses cough/ cold drug products containing an oral bronchodilator (ephedrine and its salts) in combination with any expectorant or any oral nasal decongestant. Timetable:

Action	Date	FR Cite
NPRM (Amend- ment).	07/13/05	70 FR 40232
NPRM Comment Period End.	11/10/05	
Final Action (Technical Amendment).	03/19/07	72 FR 12730
Final Action	06/00/14	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Janice Adams-King, Regulatory Health Project Manager, Department of Health and Human Services, Food and Drug Administration, Center for Drug Evaluation and Research, WO 22, Room 5416, 10903 New Hampshire Avenue, Silver Spring, MD 20993, Phone: 301 796-3713, Fax: 301 796-9899, Email: janice.adams-king@fda.hhs.gov. RIN: 0910-AF33

297. Laser Products; Proposed Amendment to Performance Standard

Legal Authority: 21 U.S.C. 360hh to 360ss; 21 U.S.C. 371; 21 U.S.C. 393 Abstract: FDA is proposing to amend

the performance standard for laser products to achieve closer harmonization between the current standard and the International Electrotechnical Commission (IEC) standard for laser products and medical laser products. The proposed amendment is intended to update FDA's performance standard to reflect advancements in technology.

Timetable:

Action	Date	FR Cite
NPRM NPRM Comment Period End.	06/24/13 09/23/13	78 FR 37723
Final Action	06/00/14	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Nancy Pirt, Regulatory Counsel, Department of Health and Human Services, Food and Drug Administration, Center for Devices and Radiological Health, WO 66, Room 4438, 10903 New Hampshire Avenue, Silver Spring, MD 20993, Phone: 301 796-6248, Fax: 301 847-8145, Email: nancy.pirt@fda.hhs.gov.

RIN: 0910-AF87

298. Food Labeling: Calorie Labeling of **Articles of Food Sold in Vending Machines**

Regulatory Plan: This entry is Seq. No. 57 in part II of this issue of the Federal Register.

RIN: 0910-AG56

299. Food Labeling: Nutrition Labeling of Standard Menu Items in Restaurants and Similar Retail Food Establishments

Regulatory Plan: This entry is Seq. No. 58 in part II of this issue of the Federal Register.

RIN: 0910-AG57

300. Use of Certain Sysmbols in Labeling

Legal Authority: sec 502(c) of the Food Drug and Cosmetic Act (FD&C Act), 21 U.S.C. 352(c); sec 514(c) of FD&C Act, 21 U.S.C. 360d(c), enacted by the Food and Drug Modernization Act of 1997 (FDAMA)

Abstract: The purpose of this rule is to allow for the inclusion of certain stand-alone symbols contained in a standard that FDA recognizes, provided that such symbols are explained in a symbols glossary that contemporaneously accompanies the medical device.

Timetable:

Action	Date	FR Cite
NPRM NPRM Comment Period End	04/19/13 06/18/13	78 FR 23508
Final Action	04/00/14	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Mary Follette Story, Human Factors and Accessible Medical Technology Specialist, Department of Health and Human Services, Food and Drug Administration, Center for Devices and Radiological Health, Room 2553, 10903 New Hampshire Avenue, Silver Spring, MD 20993, *Phone:* 301 796– 1456, *Email: molly.story@fda.hhs.gov. RIN:* 0910–AG74

301. Requirements for the Submission of Data Needed To Calculate User Fees for Manufacturers and Importers of Tobacco Products

Legal Authority: 21 U.S.C. 371; 21 U.S.C. 387s; Pub. L. 111–31

Abstract: FDA is proposing to require manufacturers and importers of tobacco products to submit certain market share data to FDA. USDA currently collects such data, but its program sunsets at the end of September 2014 and USDA will cease collection of this information. FDA is taking this action so that it may continue to calculate market share percentages needed to compute user fees.

Timetable:

Action	Date	FR Cite
NPRM NPRM Comment Period End.	05/31/13 08/14/13	78 FR 32581
Final Action	06/00/14	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Annette L. Marthaler, Regulatory Counsel, Department of Health and Human Services, Food and Drug Administration, Center for Tobacco Products, Room 340K, 9200 Corporate Boulevard, Rockville, MD 20850, Phone: 877 287–1373, Fax: 240 276–3904, Email: annette.marthaler@ fda.hhs.gov.

RIN: 0910-AG81

DEPARTMENT OF HEALTH AND HUMAN SERVICES (HHS)

Food and Drug Administration (FDA)

Long-Term Actions

302. Over-the-Counter (OTC) Drug Review—Topical Antimicrobial Drug Products

Legal Authority: 21 U.S.C. 321p; 21 U.S.C. 331; 21 U.S.C. 351 to 353; 21 U.S.C. 355; 21 U.S.C. 360; 21 U.S.C. 371

Abstract: The OTC drug review establishes conditions under which OTC drugs are considered generally recognized as safe and effective and not misbranded. After a final monograph (i.e., final rule) is issued, only OTC drugs meeting the conditions of the monograph, or having an approved new drug application, may be legally marketed. This action addresses antimicrobial agents in consumer hand wash products.

Timetable:

Action	Date	FR Cite
NPRM (Healthcare).	06/17/94	59 FR 31402
Comment Period End.	12/15/95	
NPRM (Consumer Hand Wash Products).	12/00/14	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: David Eng, Regulatory Project Manager, Department of Health and Human Services, Food and Drug Administration, Center for Drug Evaluation and Research, WO 22, Room 5487, 10903 New Hampshire Avenue, Silver Spring, MD 20993, *Phone:* 301 796–2773, *Fax:* 301 796– 9899, *Email: david.eng@fda.hhs.gov.*

RIN: 0910-AF69

303. Amendment to the Current Good Manufacturing Practice Regulations for Pharmaceuticals—Second Phase

Legal Authority: 21 U.S.C. 321; 21 U.S.C. 351; 21 U.S.C. 352; 21 U.S.C. 355; 21 U.S.C. 360b; 21 U.S.C. 371; 21 U.S.C. 374; 42 U.S.C. 262; 42 U.S.C. 264

Abstract: FDA will revise regulations for "current good manufacturing practice" for oversight and controls over the manufacture of drugs to ensure quality, including managing the risk of and establishing the safety of raw materials, materials used in the manufacturing of drugs, and finished drug products. This revision will update and harmonize requirements and improve detection and response to emerging product safety and quality signals.

Timetable:

Action	Date	FR Cite
NPRM	11/00/14	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Paula Katz, Regulatory Counsel, Office of Compliance, Department of Health and Human Services, Food and Drug Administration, Center for Drug Evaluation and Research, WO 51, Room 4314, 10903 New Hampshire Avenue, Silver Spring, MD 20993, *Phone:* 301 796–6972, *Fax:* 301 847–8742, *Email: paula.katz@fda.hhs.gov.*

RIN: 0910-AG20

304. Human Subject Protection; Acceptance of Data From Clinical Studies for Medical Devices

Legal Authority: 21 U.S.C. 321; 21 U.S.C. 331; 21 U.S.C. 351; 21 U.S.C. 352; 21 U.S.C. 360; 21 U.S.C. 360c; 21 U.S.C. 360e; 21 U.S.C. 360i; 21 U.S.C. 360j; 21 U.S.C. 371; 21 U.S.C. 374; 21 U.S.C. 381; 21 U.S.C. 393; 42 U.S.C. 264; 42 U.S.C. 271; * * *

Abstract: This rule will amend FDA's regulations on acceptance of data from clinical studies conducted in support of a premarket approval application, humanitarian device exemption application, an investigational device exemption application, or a premarket notification submission for a medical device.

Timetable:

Action	Date	FR Cite
NPRM NPRM Comment Period End.	02/25/13 05/28/13	78 FR 12664
Final Action	12/00/14	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Sheila Anne Brown, Policy Analyst, Investigational Device Exemptions Staff, Department of Health and Human Services, Food and Drug Administration, WO 66, Room 1651, 10903 New Hampshire Avenue, Silver Spring, MD 20993, *Phone:* 301 796– 6563, *Fax:* 301 847–8120, *Email: sheila.brown@fda.hhs.gov. BIN:* 0010 ACA8

RIN: 0910–AG48

305. Amendments to the Current Good Manufacturing Practice Regulations for Finished Pharmaceuticals— Components

Legal Authority: 21 U.S.C. 321; 21 U.S.C. 351; 21 U.S.C. 352; 21 U.S.C. 355; 21 U.S.C. 360b; 21 U.S.C. 360bbb–7; 21 U.S.C. 371; 21 U.S.C. 374; 42 U.S.C. 262; 42 U.S.C. 264

Abstract: FDA will revise regulations for "current good manufacturing practice" with regard to the control over components used in manufacturing finished pharmaceuticals.

Timetable:

Action	Date	FR Cite
NPRM	11/00/14	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Brian Hasselbalch, Consumer Safety Officer, Department of Health and Human Services, Food and Drug Administration, Center for Drug Evaluation and Research, WO 51, Room 4364, 10903 New Hampshire Avenue, Silver Spring, MD 20993, *Phone:* 301 796–3279, *Email: brian.hasselbalch@ fda.hhs.gov.*

Paula Katz, Consumer Safety Officer, Department of Health and Human Services, Food and Drug Administration, Center for Drug Evaluation and Research, WO 51, Room 1320, 10903 New Hampshire Avenue, Silver Spring, MD 20993, *Phone:* 301 796–6972, *Email: paula.katz@ fda.hhs.gov.*

RIN: 0910–AG70

DEPARTMENT OF HEALTH AND HUMAN SERVICES (HHS)

Food and Drug Administration (FDA)

Completed Actions

306. Unique Device Identification

Legal Authority: 21 U.S.C. 351; 21 U.S.C. 352; 21 U.S.C. 360; 21 U.S.C. 360h; 21 U.S.C. 360i; 21 U.S.C. 360j; 21 U.S.C. 360l; 21 U.S.C. 371

Abstract: FDA is issuing a final rule establishing a unique device identification system for medical devices. A unique device identification system would allow healthcare professionals and others to rapidly and precisely identify a device and obtain important information concerning the device and would reduce medical errors.

Timetable:

Action	Date	FR Cite
NPRM NPRM Comment Period End.	07/10/12 11/07/12	77 FR 40735
Second NPRM Second NPRM Comment Pe- riod End.	11/19/12 12/19/13	77 FR 69393
Final Action	09/24/13	78 FR 58786

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: John J. Crowley, Senior Advisor for Patient Safety, Department of Health and Human Services, Food and Drug Administration, Center for Devices and Radiological Health, WO 66, Room 2315, 10903 New Hampshire Avenue, Silver Spring, MD 20993, *Phone:* 301 980–1936, *Email: jay.crowley@ fda.hhs.gov.*

RIN: 0910-AG31

307. Food Labeling: Serving Sizes; Reference Amount and Serving Size Declaration for Hard Candies and Breath Mints

Legal Authority: 21 U.S.C. 321; 21 U.S.C. 343; 21 U.S.C. 371

Abstract: FDA is proposing to change the nutrition label serving size for breath mints to one unit. FDA is taking this action in response to a citizen petition that requested a serving size for breath mints that more accurately reflects the amount customarily consumed per eating occasion and comments received on an advance notice of proposed rulemaking published in 2005.

Timetable:

Action	Date	FR Cite
NPRM NPRM Comment Period End.	12/30/97 03/16/98	62 FR 67775
ANPRM ANPRM Comment Period End. Withdrawn and	04/05/05 06/20/05 08/14/13	70 FR 17010
Merged with 0910–AF23.		

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Mark Kantor, Nutritionist, Department of Health and Human Services, Food and Drug Administration, HFS–830, 5100 Paint Branch Parkway, College Park, MD 20740, *Phone:* 240 402–1450, *Fax:* 301 436–1191, *Email: mark.kantor@ fda.hhs.gov.*

RIN: 0910-AG82

308. Food Labeling; Gluten-Free Labeling of Foods

Legal Authority: Title II of Pub. L. 108–282; 21 U.S.C. 321; 21 U.S.C. 331; 21 U.S.C. 342; 21 U.S.C. 343; 21 U.S.C. 348; 21 U.S.C. 371

Abstract: FDA is amending its regulations to define the term "glutenfree" for voluntary use in the labeling of foods. FDA is taking this action to assist persons who have celiac disease to more easily identify foods that they can eat while following a "gluten-free" diet. *Timetable:*

Action	Date	FR Cite
NPRM	01/23/07	72 FR 2795
NPRM Comment	04/23/07	
Period End.		
NPRM Comment	08/03/11	76 FR 46671
Period Re-		
opened.		
NPRM Comment	10/03/11	
Period Re-		
opened End.		
Final Action	08/05/13	78 FR 47154

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Felicia Billingslea, Director, Food Labeling and Standard Staff, Department of Health and Human Services, Food and Drug Administration, Room 4D045, HFS 820, 5100 Paint Branch Parkway, College Park, MD 20740, *Phone:* 240 402–1803, *Fax:* 301 436–2636, *Email: felicia.billingslea@fda.hhs.gov. RIN:* 0910–AG84

DEPARTMENT OF HEALTH AND HUMAN SERVICES (HHS)

Centers for Medicare & Medicaid Services (CMS)

Proposed Rule Stage

309. Emergency Preparedness Requirements for Medicare and Medicaid Participating Providers and Suppliers (CMS-3178-P) (Section 610 Review)

Legal Authority: 42 U.S.C. 1821; 42 U.S.C. 1861 (ff) (3)(B)(i)(ii); 42 U.S.C. 1913 (c)(1) et al

Abstract: This rule proposes emergency preparedness requirements for Medicare and Medicaid participating providers and suppliers to ensure that they adequately plan for both natural and man-made disasters and coordinate with Federal, State, tribal, regional, and local emergency preparedness systems. This rule would ensure providers and suppliers are adequately prepared to meet the needs of patients, residents, clients, and participants during disasters and emergency situations.

Timetable:

Action	Date	FR Cite
NPRM	11/00/13	

Regulatory Flexibility Analysis Required: Yes.

Âgency Contact: Janice Graham, Health Insurance Specialist, Clincal Standards Group, Department of Health and Human Services, Centers for Medicare & Medicaid Services, Center for Clincial Standards and Quality, Mail Stop S3–02–01, 7500 Security Boulevard, Baltimore, MD 21244–1850, *Phone:* 410 786–8020, *Email: janice.graham@cms.hhs.gov. RIN:* 0938–AO91

310. Prospective Payment System for Federally Qualified Health Centers; Changes to Contracting Policies for Rural Health Clinics and CLIA Enforcement Actions for Proficiency Testing Referral (CMS–1443–F) (Section 610 Review)

Legal Authority: Pub. L. 111–148, sec 10501

Abstract: This final rule establishes methodology and payment rates for a prospective payment system (PPS) for federally qualified health center (FQHC) services under Medicare Part B beginning on October 1, 2014, in compliance with the statutory requirement of the Affordable Care Act. This rule also establishes a policy which would allow rural health clinics (RHCs) to contract with nonphysician practitioners when statutory requirements for employment of nurse practitioners and physician assistants are met, and makes other technical and conforming changes to the RHC and FQHC regulations. Finally, this rule makes changes to the Clinical Laboratory Improvement Amendments (CLIA) regulations regarding enforcement actions for proficiency testing referral.

Timetable:

Action	Date	FR Cite
NPRM NPRM Comment Period End.	09/23/13 11/18/13	78 FR 58386
Final Action	08/00/14	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Sarah Harding, Health Insurance Specialist, Department of Health and Human Services, Centers for Medicare & Medicaid Services, Mail Stop C4-01-26, 7500 Security Boulevard, Windsor Mill, MD 21244, Phone: 410 786-4001, Email: sarah.harding@cms.hhs.gov. RIN: 0938-AR62

311. • Hospital Inpatient Prospective **Payment System for Acute Care** Hospitals and the Long-Term Care **Hospital Prospective Payment System** and Fiscal Year 2015 Rates (CMS-1607-P)

Regulatory Plan: This entry is Seq. No. 62 in part II of this issue of the Federal Register. RIN: 0938-AS11

312. • CY 2015 Revisions to Payment **Policies Under the Physician Fee** Schedule and Other Revisions to Medicare Part B (CMS-1612-P)

Regulatory Plan: This entry is Seq. No. 63 in part II of this issue of the Federal Register.

RIN: 0938-AS12

313. • CY 2015 Hospital Outpatient Prospective Payment System (PPS) Policy Changes and Payment Rates, and CY 2015 Ambulatory Surgical Center **Payment System Policy Changes and** Payment Rates (CMS-1613-P)

Regulatory Plan: This entry is Seq. No. 64 in part II of this issue of the Federal Register.

RIN: 0938-AS15

DEPARTMENT OF HEALTH AND HUMAN SERVICES (HHS)

Centers for Medicare & Medicaid Services (CMS)

Final Rule Stage

314. Covered Outpatient Drugs (CMS-2345–F) (Section 610 Review)

Legal Authority: Pub. L. 111–48, secs 2501, 2503, 3301(d)(2); Pub. L. 111-152, sec 1206; Pub. L. 111-8, sec 221

Abstract: This final rule revises requirements pertaining to Medicaid reimbursement for covered outpatient drugs to implement provisions of the Affordable Care Act. This rule also revises other requirements related to covered outpatient drugs, including key aspects of Medicaid coverage, payment, and the drug rebate program.

Timetable:

Action	Date	FR Cite
NPRM NPRM Comment Period End.	02/02/12 04/02/12	77 FR 5318
Final Action	05/00/14	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Wendy Tuttle, Health Insurance Specialist, Department of Health and Human Services, Centers for Medicare & Medicaid Services, Center for Medicaid and State Operations, Mail Stop S2-14-26, 7500 Security Boulevard, Baltimore, MD 21244, Phone: 410 786-8690, Email: wendy.tuttle@cms.hhs.gov. RIN: 0938-AQ41

315. CY 2014 Changes to the End-Stage **Renal Disease (ESRD) Prospective Payment System, ESRD Quality** Incentive Program, and Durable Medical Equipment (CMS-1526-F)

Legal Authority: MIPPA sec 153(b); Pub. L. 111-148 sec 3401(h); ATRA sec 632(a)

Abstract: This final rule updates the bundled payment system for End Stage Renal Disease (ESRD) facilities by 1/1/ 13. The rule also updates the Quality Incentives in the ESRD Program. In addition, this rule clarifies the grandfathering provision related to the 3-year minimum lifetime requirement for Durable Medical Equipment (DME). It also provides clarification of the definition of routinely purchased DME. Timetable:

Action	Date	FR Cite
NPRM NPRM Comment Period End. Final Action	07/08/13 08/30/13 11/00/13	78 FR 40835

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Michelle Cruse, Health Insurance Specialist, Department of Health and Human Services, Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Mailstop C5–05–27, Baltimore, MD 21244, Phone: 410 786-7540, Email: michelle.cruse@ cms.hhs.gov.

RIN: 0938–AR55

316. Revisions to Payment Policies Under the Physician Fee Schedule and Medicare Part B for CY 2014 (CMS-1600-F)

Legal Authority: Social Security Act secs 1102, 1871, 1848

Abstract: This final rule revises payment polices under the Medicare physician fee schedule, and make other policy changes to payment under Medicare Part B. These changes are applicable to services furnished on or after January 1 annually.

Timetable:

Action	Date	FR Cite
NPRM NPRM Comment Period End.	07/19/13 09/06/13	78 FR 43282
Final Action	11/00/13	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Kathy Bryant, Deputy Director, Division of Practitioner Services, Department of Health and Human Services, Centers for Medicare & Medicaid Services, Mail Stop C4-01-27, 7500 Security Boulevard, Baltimore, MD 21244, Phone: 410 786-3448, Email: kathy.bryant@cms.hhs.gov.

RIN: 0938-AR56

317. • Adoption of Operating Rules for HIPAA Transactions (CMS-0036-IFC)

Legal Authority: Pub. L. 104–191, sec 1104

Abstract: Under the Affordable Care Act, this interim final rule adopts operating rules for HIPAA transactions for health care claims or equivalent encounter information, enrollment and disenrollment of a health plan, health plan premium payments, and referral certification and authorization.

Timetable:

Action	Date	FR Cite
Interim Final Rule	06/00/14	

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Christine Stahlecker, Acting Director, Administrative Simplification Group, Office of E-Health Standards and Services, Department of

Health and Human Services, Centers for Medicare & Medicaid Services, Mail Stop S2–26–17, 7500 Security Boulevard, Baltimore, MD 21244, *Phone:* 410 786–6405, *Email: christine.stahlecker@cms.hhs.gov. RIN:* 0938–AS01

DEPARTMENT OF HEALTH AND HUMAN SERVICES (HHS)

Centers for Medicare & Medicaid Services (CMS)

Completed Actions

318. Changes to the Hospital Inpatient and Long-Term Care Prospective Payment System for FY 2014 (CMS– 1599–F)

Legal Authority: sec 1886(d) of the Social Security Act

Abstract: This annual rule revises the Medicare hospital inpatient and longterm care hospital prospective payment systems for operating and capital-related costs. This rule implements changes arising from our continuing experience with these systems.

Timetable:

Action	Date	FR Cite
NPRM NPRM Comment Period End.	05/10/13 06/25/13	78 FR 27485
Final Action	08/19/13	78 FR 50419

Regulatory Flexibility Analysis Required: Yes.

Ågency Contact: Roechel Kujawa, Health Insurance Specialist, Department of Health and Human Services, Centers for Medicare & Medicaid Services, Mail Stop C4–07–07, 7500 Security Boulevard, Baltimore, MD 21244, Phone: 410 786–9111, Email: roechel.kujawa@cms.hhs.gov. RIN: 0938–AR53

319. Changes to the Hospital Outpatient Prospective Payment System and Ambulatory Surgical Center Payment System for CY 2014 (CMS-1601-F)

Legal Authority: sec 1833 of the Social Security Act

Abstract: This final rule revises the Medicare hospital outpatient prospective payment system to implement applicable statutory requirements and changes arising from our continuing experience with this system. The rule also describes changes to the amounts and factors used to determine payment rates for services. In addition, the rule finalizes changes to the Ambulatory Surgical Center Payment System list of services and rates.

Timetable:

Action	Date	FR Cite
NPRM NPRM Comment	07/19/13 09/06/13	78 FR 43534
Period End. Final Action	09/06/13	78 FR 54842

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: Marjorie Baldo, Health Insurance Specialist, Department of Health and Human Services, Centers for Medicare & Medicaid Services, Center for Medicare Management, Mail Stop C4–03–06, 7500 Security Boulevard, Baltimore, MD 21244, Phone: 410 786–4617, Email: marjorie.baldo@cms.hhs.gov. RIN: 0938–AR54

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Department of Homeland Security

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