the alternate system must be accepted by the appropriate PMI and require the maintenance records to be maintained either indefinitely or until the work is repeated. Records of the piece-part inspections are not required under § 121.380 (a) (2) (vi) of the Federal Aviation Regulations (14 CFR 121.380 (a) (2) (vi)). All other Operators must maintain the records of mandatory inspections required by the applicable regulations governing their operations.

Note 3: The requirements of this AD have been met when the engine manual changes are made and air carriers have modified their continuous airworthiness maintenance plans to reflect the requirements in the engine manuals.

Effective Date

(f) This amendment becomes effective on April 6, 2002.

Issued in Burlington, Massachusetts, on February 21, 2002.

Jay J. Pardee,
Manager, Engine and Propeller Directorate,
Aircraft Certification Service.

[FR Doc. 02–5003 Filed 3–1–02; 8:45 am]

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration


Change in the Removal of the Office of Management and Budget (OMB) Control Numbers; Technical Amendment

AGENCY: Food and Drug Administration, HHS.

ACTION: Final rule; technical amendment.

SUMMARY: The Food and Drug Administration (FDA) is amending its regulations to reflect a change in the removal of OMB control numbers. This action is editorial in nature and is intended to improve the accuracy of the agency’s regulations.

EFFECTIVE DATE: March 4, 2002.

FOR FURTHER INFORMATION CONTACT: Peggy Schlosburg, Office of Information Resources Management (HFA–250), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301–827–1223.

SUPPLEMENTARY INFORMATION: FDA is amending its regulations in 21 CFR parts 56, 58, 60, 101, 107, 179, 310, 312, 314, 510, 514, 606, 610, 640, 660, 680, 720, 814, 1020, and 1040 to reflect a change in the removal of the outdated OMB control numbers. We no longer need to publish OMB control numbers in the CFR, because they are now displayed in a separate Federal Register notice announcing OMB approval for the collection of information.

Publication of this document constitutes final action on these changes under the Administrative Procedure Act (5 U.S.C. 553). Notice and public procedure are unnecessary because FDA is merely correcting nonsubstantive errors.

List of Subjects

21 CFR Part 56

Human research subjects, Reporting and recordkeeping requirements, Safety.

21 CFR Part 58

Laboratories, Reporting and recordkeeping requirements.

21 CFR Part 60

Administrative practice and procedure, Drugs, Food additives, Inventions and patents, Medical devices, Reporting and recordkeeping requirements.

21 CFR Part 101

Food labeling, Nutrition, Reporting and recordkeeping requirements.

21 CFR Part 107

Food labeling, Infants and children, Nutrition, Reporting and recordkeeping requirements, Signs and symbols.

21 CFR Part 179

Food additives, Food labeling, Food packaging, Radiation protection, Reporting and recordkeeping requirements, Signs and symbols.

21 CFR Part 310

Administrative practice and procedure, Drums, Labeling, Medical devices, Reporting and recordkeeping requirements.

21 CFR Part 312

Drugs, Exports, Imports, Investigations, Labeling, Medical research, Reporting and recordkeeping requirements, Safety.

21 CFR Part 314

Administrative practice and procedure, Confidential business information, Drugs, Reporting and recordkeeping requirements.

21 CFR Part 510

Administrative practice and procedure, Animal drugs, Labeling, Reporting and recordkeeping requirements.

21 CFR Part 514

Administrative practice and procedure, Animal drugs, Confidential business information, Reporting and recordkeeping requirements.

21 CFR Part 606

Blood, Labeling, Laboratories, Reporting and recordkeeping requirements.

21 CFR Part 610

Biologics, Labeling, Reporting and recordkeeping requirements.

21 CFR Part 640

Blood, Labeling, Reporting and recordkeeping requirements.

21 CFR Part 660

Biologics, Labeling, Reporting and recordkeeping requirements.

21 CFR Part 680

Biologics, Blood, Reporting and recordkeeping requirements.

21 CFR Part 720

Confidential business information, Cosmetics.

21 CFR Part 814

Administrative practice and procedure, Confidential business information, Medical devices, Medical research, Reporting and recordkeeping requirements.

21 CFR Part 1020

Electronic products, Medical devices, Radiation protection, Reporting and recordkeeping requirements, Television, X-rays.

21 CFR Part 1040

Electronic products, Labeling, Lasers, Medical devices, Radiation protection, Reporting and recordkeeping requirements.

Therefore, under the Federal Food, Drug, and Cosmetic Act and under authority delegated to the Commissioner of Food and Drugs, 21 CFR parts 56, 58, 60, 101, 107, 179, 310, 312, 314, 510, 514, 606, 610, 640, 660, 680, 720, 814, 1020, and 1040 are amended as follows:

PART 56—INSTITUTIONAL REVIEW BOARDS

1. The authority citation for 21 CFR part 56 continues to read as follows:

PART 58—GOOD LABORATORY PRACTICE FOR NONCLINICAL LABORATORY STUDIES

4. The authority citation for 21 CFR part 58 continues to read as follows:

PART 107—INVESTIGATIONAL NEW DRUG APPLICATION

27. The authority citation for 21 CFR part 312 continues to read as follows:

PART 312—INVESTIGATIONAL NEW DRUG APPLICATION

28. In §312.7 Promotion and charging for investigational drugs, remove the parenthetical phrase at the end of the section.

PART 310—NEW DRUGS

25. The authority citation for 21 CFR part 310 continues to read as follows:
§ 312.38 [Amended]
37. In § 312.38 Withdrawal of an IND, remove the parenthetical phrase at the end of the section.

§ 312.41 [Amended]
38. In § 312.41 Comment and advice on an IND, remove the parenthetical phrase at the end of the section.

§ 312.44 [Amended]
39. In § 312.44 Termination, remove the parenthetical phrase at the end of the section.

§ 312.45 [Amended]
40. In § 312.45 Inactive status, remove the parenthetical phrase at the end of the section.

§ 312.47 [Amended]
41. In § 312.47 Meetings, remove the parenthetical phrase at the end of the section.

§ 312.53 [Amended]
42. In § 312.53 Selecting investigators and monitors, remove the parenthetical phrase at the end of the section.

§ 312.55 [Amended]
43. In § 312.55 Informing investigators, remove the parenthetical phrase at the end of the section.

§ 312.56 [Amended]
44. In § 312.56 Review of ongoing investigations, remove the parenthetical phrase at the end of the section.

§ 312.57 [Amended]
45. In § 312.57 Recordkeeping and record retention, remove the parenthetical phrase at the end of the section.

§ 312.59 [Amended]
46. In § 312.59 Disposition of unused supply of investigational drug, remove the parenthetical phrase at the end of the section.

§ 312.62 [Amended]
47. In § 312.62 Investigator recordkeeping and record retention, remove the parenthetical phrase at the end of the section.

§ 312.64 [Amended]
48. In § 312.64 Investigator reports, remove the parenthetical phrase at the end of the section.

§ 312.66 [Amended]
49. In § 312.66 Assurance of IRB review, remove the parenthetical phrase at the end of the section.

§ 312.70 [Amended]
50. In § 312.70 Disqualification of a clinical investigator, remove the parenthetical phrase at the end of the section.

§ 312.110 [Amended]
51. In § 312.110 Import and export requirements, remove the parenthetical phrase at the end of the section.

§ 312.120 [Amended]
52. In § 312.120 Foreign clinical studies not conducted under an IND, remove the parenthetical phrase at the end of the section.

§ 312.140 [Amended]
53. In § 312.140 Address for correspondence, remove the parenthetical phrase at the end of the section.

§ 312.160 [Amended]
54. In § 312.160 Drugs for investigational use in laboratory research animals or in vitro tests, remove the parenthetical phrase at the end of the section.

§ 312.170 [Amended]
55. In § 312.170 Changes in ownership, remove the parenthetical phrase at the end of the section.

PART 314—APPLICATIONS FOR FDA APPROVAL TO MARKET A NEW DRUG


§ 314.50 [Amended]
56. In § 314.50 Content and format of an application, remove the parenthetical phrase at the end of the section.

§ 314.70 [Amended]
57. In § 314.70 Supplements and other changes to an approved application, remove the parenthetical phrase at the end of the section.

§ 314.71 [Amended]
58. In § 314.71 Procedures for submission of a supplement to an approved application, remove the parenthetical phrase at the end of the section.

§ 314.72 [Amended]
59. In § 314.72 Changes in ownership of an application, remove the parenthetical phrase at the end of the section.

§ 314.80 [Amended]
60. In § 314.80 Postmarketing reporting of adverse drug experiences, remove the parenthetical phrase at the end of the section.

§ 314.90 [Amended]
61. In § 314.90 Waivers, remove the parenthetical phrase at the end of the section.

PART 510—NEW ANIMAL DRUGS


§ 510.455 [Amended]
66. In § 510.455 New animal drug requirements regarding free-choice administration in feeds, remove the parenthetical phrase at the end of the section.

PART 514—NEW ANIMAL DRUG APPLICATIONS

67. The authority citation for 21 CFR part 514 continues to read as follows: Authority: 21 U.S.C. 351, 352, 360b, 371, 379e, 381.

§ 514.1 [Amended]
68. In § 514.1 Applications, remove the parenthetical phrase at the end of the section.

PART 606—CURRENT GOOD MANUFACTURING PRACTICE FOR BLOOD AND BLOOD COMPONENTS


§ 606.170 [Amended]
70. In § 606.170 Adverse reaction file, remove the parenthetical phrase at the end of the section.

PART 610—GENERAL BIOLOGICAL PRODUCTS

§ 660.25 [Amended]  
82. In § 660.25 Potency tests without reference preparations, remove the parenthetical phrase at the end of the section.

§ 660.26 [Amended]  
83. In § 660.26 Specificity tests and avidity tests, remove the parenthetical phrase at the end of the section.

§ 660.28 [Amended]  
84. In § 660.28 Labeling, remove the parenthetical phrase at the end of the section.

§ 660.34 [Amended]  
85. In § 660.34 Processing, remove the parenthetical phrase at the end of the section.

§ 660.35 [Amended]  
86. In § 660.35 Labeling, remove the parenthetical phrase at the end of the section.

§ 660.36 [Amended]  
87. In § 660.36 Samples and protocols, remove the parenthetical phrase at the end of the section.

§ 660.51 [Amended]  
88. In § 660.51 Processing, remove the parenthetical phrase at the end of the section.

§ 660.52 [Amended]  
89. In § 660.52 Reference preparations, remove the parenthetical phrase at the end of the section.

§ 660.53 [Amended]  
90. In § 660.53 Controls for serological procedures, remove the parenthetical phrase at the end of the section.

§ 660.54 [Amended]  
91. In § 660.54 Potency tests, specificity tests, tests for heterospecific antibodies, and additional tests for nonspecific properties, remove the parenthetical phrase at the end of the section.

§ 660.55 [Amended]  
92. In § 660.55 Labeling, remove the parenthetical phrase at the end of the section.

PART 680—ADDITIONAL STANDARDS FOR MISCELLANEOUS PRODUCTS

93. The authority citation for 21 CFR part 680 continues to read as follows:  

§ 680.1 [Amended]  
94. In § 680.1 Allergenic products, remove the parenthetical phrase at the end of the section.

§ 680.2 [Amended]  
95. In § 680.2 Manufacture of allergenic products, remove the parenthetical phrase in paragraph (f) of this section.

§ 680.3 [Amended]  
96. In § 680.3 Tests, remove the parenthetical phrase at the end of the section.

PART 720—VOLUNTARY FILING OF COSMETIC PRODUCT INGREDIENT COMPOSITION STATEMENTS

97. The authority citation for 21 CFR part 720 continues to read as follows:  

§ 720.6 [Amended]  
98. In § 720.6 Amendments to statement, remove the parenthetical phrase at the end of the section.

PART 814—PREMARKET APPROVAL OF MEDICAL DEVICES

99. The authority citation for 21 CFR part 814 continues to read as follows:  

§ 814.20 [Amended]  
100. In § 814.20 Application, remove the parenthetical phrase at the end of the section.

§ 814.39 [Amended]  
101. In § 814.39 PMA supplements, remove the parenthetical phrase at the end of the section.

§ 814.84 [Amended]  
102. In § 814.84 Reports, remove the parenthetical phrase at the end of the section.

PART 1020—PERFORMANCE STANDARDS FOR LIGHT-EMITTING PRODUCTS

103. The authority citation for 21 CFR part 1020 continues to read as follows:  

§ 1020.33 [Amended]  
104. In § 1020.33 Computed tomography (CT) equipment, remove the parenthetical phrase at the end of the section.

PART 1040—PERFORMANCE STANDARDS FOR LIGHT-EMITTING PRODUCTS

105. The authority citation for 21 CFR part 1040 continues to read as follows:  

§ 1040.20 [Amended]  
106. In § 1040.20 Sunlamp products and ultraviolet lamps intended for use in sunlamp products, remove the
DEPARTMENT OF TRANSPORTATION

Coast Guard

33 CFR Part 165

COTP Pittsburgh-02–001

RIN 2115–AA97

Security Zone; Ohio River Mile 119.0 to 119.8, Natrium, West Virginia

AGENCY: Coast Guard, DOT.

ACTION: Temporary final rule.

SUMMARY: The Coast Guard is establishing a temporary security zone encompassing all water extending 200 feet from the shoreline of the left descending bank on the Ohio River, beginning from mile marker 119.0 and ending at mile marker 119.8. This security zone is necessary to protect the PPG Plant in Natrium, West Virginia from any and all subversive actions from any groups or individuals whose objective it is to cause disruption to the daily operations of the PPG Plant. Entry of vessels into this security zone is prohibited unless authorized by the Coast Guard Captain of the Port Pittsburgh or his designated representative.

DATES: This rule is effective from 8 a.m. on February 8, 2002 through 8 a.m. on June 15, 2002.

ADDRESS: Documents indicated in this preamble as being available in the docket, are part of docket COTP Pittsburgh-02–001 and are available for inspection or copying at Marine Safety Office Pittsburgh, Suite 1150 Kossman Bldg., 100 Forbes Ave. Pittsburgh, PA between 8 a.m. and 4 p.m., Monday through Friday, except federal holidays.

FOR FURTHER INFORMATION CONTACT: Chief Petty Officer, Brian Smith, Marine Safety Office Pittsburgh at (412) 644–5808.

SUPPLEMENTARY INFORMATION:

Regulatory Information

We did not publish a notice of proposed rulemaking (NPRM) for this regulation. Under 5 U.S.C. 553(b)(B), the Coast Guard finds that good cause exists for not publishing an NPRM, and, under 5 U.S.C. 553(d)(3), good cause exists for making this rule effective less than 30 days after publication in the Federal Register. The catastrophic nature of, and resulting devastation from, the September 11, 2001 attacks on the World Trade Center towers in New York City and the Pentagon in Washington D.C., makes this rulemaking necessary for the protection of national security interests. National security and intelligence officials warn that future terrorist attacks against United States interests are likely. Any delay in making this regulation effective would be contrary to the public interest because immediate action is necessary to protect against the possible loss of life, injury, or damage to property.

Background and Purpose

On September 11, 2001, both towers of the World Trade Center and the Pentagon were attacked by terrorists. In response to these terrorist acts, heightened awareness and security of our ports and harbors is necessary. To enhance that security the Captain of the Port, Pittsburgh is establishing a temporary security zone.

This security zone includes all water extending 200 feet from the shoreline of the left descending bank on the Ohio River beginning from mile marker 119.0 and ending at mile marker 119.8. This security zone is necessary to protect the public, facilities, and surrounding area from possible acts of terrorism at the PPG Plant. All vessels and persons are prohibited from entering the zone without the permission of the Captain of the Port Pittsburgh or his designated representative.

Regulatory Evaluation

This rule is not a “significant regulatory action” under section 3(f) of Executive Order 12866 and does not require an assessment of potential costs and benefits under section 6(a)(3) of that Order. The Office of Management and Budget has not reviewed it under that Order. It is not “significant” under the regulatory policies and procedures of the Department of Transportation (DOT)(44 FR 11040, February 26, 1979).

The Coast Guard expects the economic impact of this proposal to be so minimal that a full Regulatory Evaluation under paragraph 10 (e) of the regulatory policies and procedures of DOT is unnecessary. This rule will not obstruct the regular flow of vessel traffic and will allow vessel traffic to pass safely around the security zone and vessels may be permitted to enter the security zone on a case-by-case basis.

Small Entities

Under the Regulatory Flexibility Act (5 U.S.C. 601–612), we considered whether this rule would have a significant economic impact on a substantial number of small entities. The term “small entities” comprises small businesses, not-for-profit organizations that are independently owned and operated and are not dominant in their fields, and governmental jurisdictions with populations of less than 50,000.

The Coast Guard certifies under 5 U.S.C. 605(b) that this rule will not have a significant economic impact on a substantial number of small entities. This security zone will not have an impact on a substantial number of small entities because this rule will not obstruct the regular flow of vessel traffic and will allow vessel traffic to pass safely around the security zone.

If you are a small business entity and are significantly affected by this regulation please contact Chief Petty Officer Brian Smith, U.S. Coast Guard Marine Safety Pittsburgh, Suite 1150 Kossman Bldg., 100 Forbes Ave. Pittsburgh, PA at (412) 644–5808.

Assistance for Small Entities

Under section 213(a) of the Small Business Regulatory Enforcement Fairness Act of 1996 (Pub. L. 104–121), we offer to assist small entities in understanding the rule so that they can better evaluate its effects on them and participate in the rulemaking process. Small businesses may send comments on the actions of Federal employees who enforce, or otherwise determine compliance with, Federal regulations to the Small Business and Agriculture Regulatory Enforcement Ombudsman and the Regional Small Business Regulatory Fairness Boards. The Ombudsman evaluates these actions annually and rates each agency’s responsiveness to small business. If you wish to comment on actions by employees of the Coast Guard, call 1–888–REG–FAIR (1–888–734–3247).

Collection of Information

This rule calls for no new collection of information under the Paperwork Reduction Act of 1995 (44 U.S.C. 3501–3520).

Federalism

A rule has implications for federalism under Executive Order 13132. Federalism, if it has a substantial direct effect on State or local governments and would either preempt State law or impose a substantial direct cost of compliance on them. We have analyzed this rule under that Order and have