DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Part 173

[DOCKET NO. 99F–2907]

Secondary Direct Food Additives Permitted in Food for Human Consumption

AGENCY: Food and Drug Administration, HHS.

ACTION: Final rule.

SUMMARY: The Food and Drug Administration (FDA) is amending the food additive regulations to provide for the safe use of acidified sodium chlorite solutions as an antimicrobial agent on red meat parts and organs. This action is in response to a petition filed by Alcide Corp.

DATES: This rule is effective January 12, 2000; written objections and requests for a hearing by February 11, 2000.

ADDRESSES: Written objections may be sent to the Dockets Management Branch (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852.


SUPPLEMENTARY INFORMATION: In a notice published in the Federal Register of August 30, 1999 (64 FR 47193), FDA announced that a food additive petition (FAP 9A4692) had been filed by Alcide Corp., 8561 154th Ave. NE., Redmond, WA 98052. The petition proposed to amend the food additive regulation in 21 CFR 173.325 (§ 173.325) to provide for the safe use of acidified sodium chlorite solutions as an antimicrobial agent on red meat parts and organs. FDA has evaluated data in the petition and other relevant material. Based on this information, the agency concludes that the proposed use of the additive is safe, that the additive will achieve its intended technical effect, and therefore, that the regulation in § 173.325 should be amended as set forth below.

In accordance with § 171.1(h) (21 CFR 171.1(h)), the petition and the documents that FDA considered and relied upon in reaching its decision to approve the petition are available for inspection at the Center for Food Safety and Applied Nutrition by appointment with the information contact person listed above. As provided in § 171.1(h), the agency will delete from the documents any materials that are not available for public disclosure before making the documents available for inspection.

In the notice of filing, FDA gave interested parties an opportunity to submit comments on the petitioner’s environmental assessment. FDA received no comments in response to that notice.

The agency has carefully considered the potential environmental effects of this action. FDA has concluded that the action will not have a significant impact on the human environment, and that an environmental impact statement is not required. The agency’s finding of no significant impact and the evidence supporting that finding, contained in an environmental assessment, may be seen in the Dockets Management Branch (address above) between 9 a.m. and 4 p.m., Monday through Friday. This final rule contains no collection of information. Therefore, clearance by the Office of Management and Budget under the Paperwork Reduction Act of 1995 is not required.

Any person who will be adversely affected by this regulation may at any time on or before February 11, 2000, file with the Dockets Management Branch (address above) written objections thereto. Each objection shall be separately numbered, and each numbered objection shall specify with particularity the provisions of the regulation to which objection is made and the grounds for the objection. Each numbered objection on which a hearing is requested shall specifically state the failure to request a hearing for any particular objection shall constitute a waiver of the right to a hearing on that objection. Each numbered objection for which a hearing is requested shall include a detailed description and analysis of the specific factual information intended to be presented in support of the objection in the event that a hearing is held. Failure to include such a description and analysis for any particular objection shall constitute a waiver of the right to a hearing on the objection. Three copies of all documents are to be submitted and are to be identified with the docket number found in brackets in the heading of this document. Any objections received in response to the regulation may be seen in the Dockets Management Branch between 9 a.m. and 4 p.m., Monday through Friday.

List of Subjects in 21 CFR Part 173

Food additives.

Therefore, under the Federal Food, Drug, and Cosmetic Act and under authority delegated to the Commissioner of Food and Drugs and redelegated to the Director, Center for Food Safety and Applied Nutrition, 21 CFR part 173 is amended as follows:

PART 173—SECONDARY DIRECT FOOD ADDITIVES PERMITTED IN FOOD FOR HUMAN CONSUMPTION

1. The authority citation for 21 CFR part 173 continues to read as follows:


2. Section 173.325 is amended by revising paragraph (c) to read as follows:

§ 173.325 Acidified sodium chlorite solutions.

* * * * *

(c) The additive is used as an antimicrobial agent in accordance with current industry practice in the processing of red meat, red meat parts, and organs as a component of a spray or in the processing of red meat parts and organs as a component of a dip. Applied as a dip or spray, the additive is used at levels that result in sodium chlorite concentrations between 500 and 1,200 ppm in combination with any GRAS acid at levels sufficient to achieve a solution pH of 2.5 to 2.9.

* * * * *


Janice F. Oliver,
Deputy Director for Operations, Center for Food Safety and Applied Nutrition.

[FR Doc. 00–691 Filed 1–11–00; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Part 314

[DOCKET NO. 94N–0449]

RIN 0910–AA78

New Drug Applications; Drug Master Files

AGENCY: Food and Drug Administration, HHS.

ACTION: Final rule.

SUMMARY: The Food and Drug Administration (FDA) is revising its regulation governing drug master files (DMF’s). FDA is removing the provision for submitting Type I DMF’s and will no longer permit information submitted in a Type I DMF to be incorporated by reference in investigational new drug applications (IND’s), new drug applications (NDA’s), abbreviated new drug applications (ANDA’s), or amendments or supplements to any of
these. This rule is intended to eliminate submissions of information that are not necessary either to conduct inspections of manufacturing facilities or to review the chemistry, manufacturing, and controls sections of IND's, NDA’s, and abbreviated applications.

**EFFECTIVE DATE:** July 10, 2000.

**FOR FURTHER INFORMATION CONTACT:**

**SUPPLEMENTARY INFORMATION:**

I. Background

A DMF is a voluntary submission to FDA that may be used to provide confidential, detailed information about the facilities, processes, or articles used in the manufacturing, processing, packaging, and storing of one or more human drug products. The regulations in 21 CFR 314.420(a) describe five types of DMF’s according to the kind of information to be submitted. Type I submissions include manufacturing site, facilities, operating procedures, and personnel information. Type II submissions include information regarding drug substances, drug substance intermediates, and materials used to prepare them, or drug products. Type III submissions include information about packaging materials. Type IV submissions include information concerning excipients, colorants, flavors, essences, or materials used in their preparation. Type V submissions, detailed in the guidance for industry entitled “Drug Master Files” (September 1, 1989), include FDA-accepted reference information. DMF’s allow regulated industry to submit to FDA information that may be used to support an IND, NDA, ANDA, another DMF, an export application, or amendments or supplements to any of these. DMF information may be incorporated by reference into a drug application or supplement without public disclosure.

FDA intended to use information submitted in a Type I DMF to plan its on-site inspections and travel to foreign drug manufacturing facilities. In December 1992, the Chemistry, Manufacturing, and Controls Coordination Committee (CMCCC) of the Center for Drug Evaluation and Research (CDER) established a DMF task force to review DMF procedures and consider ways of improving the DMF system. One of the task force recommendations was that Type I DMF's be eliminated. The recommendation was based on a number of factors:

1. The information contained in Type I DMF's was often outdated.
2. The Type I DMF was not always easily accessible to FDA investigators.
3. The review divisions in CDER do not review the information in most Type I DMF’s. Although information from Type I DMF’s has often been incorporated by reference into IND’s, NDA’s, and abbreviated applications, the information is not required for review of the chemistry, manufacturing, and controls section of an application. Under 21 CFR 314.50(d)(1)(i) and (d)(1)(ii), a drug product applicant is required to furnish in the application the name and location of facilities used in the manufacture of the drug substance or drug product.
4. Information concerning the facility is maintained onsite where it is available for the investigator.

The CMCCC adopted the recommendation of the DMF Task Force and, subsequently, FDA proposed eliminating Type I DMF’s in the **Federal Register** of July 3, 1995 (60 FR 34486). FDA also proposed to implement a procedure by which DMF holders could request that facility information currently contained in Type I DMF’s be transferred to Types II through V.

FDA is finalizing its proposal to eliminate Type I DMF’s. In so doing, the agency will no longer accept Type I DMF’s or correspondence updating existing Type I DMF’s and will no longer permit information previously submitted in a Type I DMF to be incorporated by reference in IND’s, NDA’s, ANDA’s, and supplemental applications for drugs approved under section 505 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 355). The Center for Biologics Evaluation and Research (CBER) has used Type I Master Files in a manner different from that used by CDER. Certain biological products, such as gene therapy products, require review of some facility information to assess their safety for use in clinical trials under IND. CBER will accept facility information for such products in Type V Master Files. CBER intends to issue a guidance on the information that may be submitted in a Type V Master File without previously obtaining permission.

II. Comments on the Proposed Rule

The agency received seven comments on the proposed rule and several of these raised multiple issues. A number of comments expressed general support for the proposal. A summary of the comments and the agency’s responses follows.

1. One firm stated that it will be manufacturing drug products for other U.S. and non-U.S. companies and needs a means to submit confidential, technical information to FDA (e.g., information regarding the firm’s new manufacturing facility, including, but not limited to, air handling systems, milling, blending, and filling technology). The firm emphasized that if Type I DMF’s are eliminated, confidential information regarding the facilities, processes, or articles used in the manufacturing, processing, packaging, and storing of drugs for human use would not be available for referencing by sponsors of IND’s or NDA’s with which the firm will contract. In addition, FDA’s review divisions will not be able to rely on the applications themselves for information typically included in a Type I DMF. The firm noted that without a Type I DMF, a Type II DMF (intermediates, drug substances, and drug products) might be the only alternative for supplying the agency with certain information and that it would be forced to file a Type II DMF for each company for which it does drug product manufacturing. The firm also stated that the submission of multiple Type II DMF’s instead of a single Type I would place an unnecessary paper burden on the agency. The firm further noted that if the agency relies on preapproval inspections, it faces the possibility of multiple inspections in any given year, placing unnecessary burdens on valuable FDA resources (i.e., multiple inspections of the same facility).

One comment noted that it is irrelevant that field investigators do not use Type I DMF’s and that, since Type I submissions are voluntary, the agency should continue to allow firms the convenience of referencing Type I submissions. Another comment suggested that instead of FDA eliminating Type I DMF’s, industry should be required to keep the information current. The comment stated that the privilege of incorporating Type I DMF information by reference should be denied on a case-by-case basis to those firms that do not keep information current.

The agency believes that several of these comments are based on a misunderstanding of the agency’s
reliance on information contained in Type I DMF’s during the drug application review process. Information contained in Type I DMF’s is not reviewed by CDER reviewers, and it plays no role in processing a drug product application.

The Type I DMF was intended to assist FDA in conducting onsite inspections of foreign manufacturing facilities. As noted above, the agency determined that the Type I DMF was not always easily accessible to investigators and that information in the document was often out-of-date. The drug product application is required to provide information on the location of manufacturing facilities and it is this current, product-specific information that is used by CDER review divisions. Continuing to maintain Type I DMF’s when the information is not used by the agency provides no benefit to either regulated industry or the agency.

If a firm is performing different processing steps for a customer, a Type I DMF would not provide the information necessary for adequate review. Moreover, the elimination of Type I DMF’s does not mean that a firm would be required to file a Type II DMF for each company for which it manufactures drug products. Reviewers examine the details of the manufacturing process as they apply to each individual product and procedures used in the manufacture of more than one drug product may be included in the same Type II DMF.

Concerns about a possible strain on FDA resources because of multiple inspections are not relevant to the Type I DMF issue since inspections are conducted in accordance with current agency inspection policy, which applies whether or not a firm has a Type I DMF. The current agency policy on inspections is described in the agency’s Investigations Operations Manual. Prior to the approval of a drug product, the facility that will manufacture the product will generally be inspected by FDA unless there has been a recent inspection for other reasons.

2. One comment stated that the production of “Generic Compounds” (which could conceivably be manufactured in smaller, stand-alone facilities possibly located in remote areas) is generally not adequately described in drug product applications and other written material submitted to FDA. The comment stated that such inadequate descriptions could increase the risk of problems resulting from admixing imported products that may not have been manufactured in a facility for which a DMF has been filed. The comment noted that a full description of a facility enhances FDA’s ability to identify facilities that do not meet FDA criteria.

CDER believes that a current, accurate facility description at the manufacturing site and an inspection of the facility are the best sources of information for assessing a facility’s ability to meet FDA standards. Current, accurate information is particularly important when a facility is remote.

3. One comment noted that agency investigators of foreign manufacturers had stated that the Type I DMF was of immense value because of the information provided. The comment noted that “having more information was preferable to having none,” and that the Type I format was superior in providing that information.

The agency agrees that accurate manufacturing information is important in evaluating drug product applications and preparing for inspections. FDA does not agree, for reasons explained above and in the proposed rule, that the Type I DMF is the most effective method of providing this information.

4. One comment stated that the proposed rule should be reconsidered because it is not globally oriented. The comment stated that, at the present time, several foreign governments link approval and acceptance of U.S. products to the data listed in Type I DMF’s.

It is not clear from the comment how foreign governments link approval and acceptance of U.S. products to the data listed in Type I DMF’s since these data are not reviewed in the approval process for U.S. products. Foreign governments that have previously relied on the information in a Type I DMF can request that the firm provide a description of the manufacturing facility to them.

5. One comment asserted that switching information from one type of DMF to another would result in a reduction in paperwork, because there would be no basic change in the system. The comment suggested that a proposal to eliminate Type I Master Files might be more appropriate. The comment observed that there would be a reduction in paperwork if the amount of information incorporated in a Type I DMF were limited to that specified in the proposed rule as appropriate for transfer to a Type V DMF. Another comment observed that the elimination of Type I DMF’s will increase the paperwork burden for industry if information about facilities, processes, and other data included in the manufacturing, processing, packaging, and storing of human drugs can no longer be reported in a Type I DMF and incorporated by reference.

Because FDA investigators and CDER review divisions do not rely on information in a Type I DMF document for inspection or approval purposes, the agency finds that the mere withdrawal of inactive Type I DMF’s would not address the agency’s concern that the Type I DMF is an inadequate vehicle for information. To address this concern, the agency is eliminating the production and maintenance of all Type I DMF documents. Therefore, based on FDA’s experience, the agency concludes that it is reasonable to anticipate a reduction in the paperwork burden by eliminating the requirement that industry produce and maintain the Type I DMF document.

6. One comment asserted that the proposal would require a rewrite of the current guidance to provide industry with information regarding the format and content of the Type V DMF’s. The agency notes that the guidance for the industry on DMF’s is currently undergoing revision and any changes regarding Type V DMF’s will require no significant additional resources. The agency advises that the only Type I DMF’s that may be converted to Type V’s are those covering sterile processing facilities and other special cases. As detailed in the discussion on implementation below, these will be examined on a case-by-case basis to decide if transferring them is justified. The agency does not anticipate that substantial agency resources will be required to evaluate the transfer of information currently included in Type I DMF’s to Types II, III, IV, or V DMF’s.

III. Implementation of the Rule

7. One comment suggested that the proposed implementation date of 60 days after publication should be reconsidered because this timeframe does not permit adequate time to revise operating procedures. One comment suggested that the proposed rule should be implemented in conjunction with an educational effort, including a workshop on DMF’s and publicity to prepare those affected by the new requirements. One comment suggested that the transfer of information from a Type I DMF to another type would require a review of written requests by the DMF staff and that this could result in a significant economic impact on the agency. One comment asserted that the proposed rule did not address those current applications which reference Type I DMF’s.

Based on comments and FDA’s own evaluation, the agency has concluded...
that the proposed implementation period is inadequate, particularly for foreign firms seeking approval where Type I DMF’s were referenced. Some firms will need time to develop alternative procedures. The agency has determined that the effective date will be 180 days after the date of publication of the final rule in the Federal Register.

After the effective date of the rule, the agency will no longer accept new Type I DMF’s or correspondence updating existing Type I DMF’s. Type I DMF’s will be transferred to the Federal Records Center and the information in Type I DMF’s currently on file may no longer be incorporated by reference into new applications, amendments, or supplements. These changes will supersede all information regarding Type I DMF’s detailed in the current guidance for industry on DMF’s.

To accommodate firms that have submitted information under a Type I DMF that should have been filed under DMF Types II through V, a list of all CDER Type I DMF’s is available for public review in the Dockets Management Branch (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852, under the docket number found in brackets in the heading of this document. The list is also available on the CDER Internet site at http://www.fda.gov/cder/dmf/index.htm. If a DMF holder believes that its Type I DMF should be recategorized or transferred to another type of DMF, the DMF holder may contact the Drug Master File Staff within 180 days of publication of this rule in the Federal Register. The agency agrees with the suggestion that the final rule should be implemented in conjunction with an educational effort and will work with the press and trade associations to publicize the obligations and options provided by the regulation.

IV) under which manufacturing information for the specific item is filed. For instance, DMF’s concerned with sterilization procedures for rubber stoppers would be reclassified as Type III DMF’s (packaging materials). Contract manufacturers of sterile drug substances and sterile finished drug products including biotechnology products filed as DMF’s, contract sterilization firms (e.g., ethylene oxide, gamma radiation, and electron beam radiation), and manufacturers of sterile finished drug products that are the subject of a drug product application may request a transfer from Type I to Type V DMF of nonproduct-specific information and procedures that are submitted to support a claim of sterility. Where applicable, the content and format of such transferred information should follow FDA’s guidance for industry entitled “Submission of Documentation for Sterilization Process Validation Applications for Human and Veterinary Drug Products” (November 1, 1994).

CBER intends to administratively recategorize current Type I Master Files that are still needed to other Master File Types as appropriate. CBER will make a list of those Type I Master Files that have not been recategorized available for public review in the Dockets Management Branch (address above), under the docket number found in brackets in the heading of this document, no later than 30 days after date of publication of this document in the Federal Register. The list will also be available on the CBER Internet site at www.fda.gov/CBER. If a holder of a Type I Master File believes that the Master File should be recategorized, the holder may contact the Division of Manufacturing and Product Quality (DMPQ) (HFM–207), Office of Compliance and Biologics Quality, CBER, 1401 Rockville Pike, Rockville, MD 20852–1448. DMPQ may also be reached at 301–827–3031. The agency advises that applicants who have current approved applications that reference Type I DMF’s transferred to Type V DMF’s may notify the agency of this change in an annual report as provided in 21 CFR 314.70.

V. Environmental Impact

The agency has determined under 21 CFR 25.30(h) that this action is of a type that does not individually or cumulatively have a significant effect on the human environment. Therefore, neither an environmental assessment nor an environmental impact statement is required.

VI. Analysis of Impacts

FDA has examined the impacts of the final rule under Executive Order 12866 and the Regulatory Flexibility Act (5 U.S.C. 601–612). Executive Order 12866 directs agencies to assess all costs and benefits of available regulatory alternatives and, when regulation is necessary, to select regulatory approaches that maximize net benefits (including potential economic, environmental, public health and safety, and other advantages; distributive impacts; and equity). The agency believes that this final rule is consistent with the regulatory philosophy and principles identified in the Executive Order. In addition, the final rule is not a significant regulatory action as defined by the Executive Order and so is not subject to review under the Executive Order.

The Regulatory Flexibility Act requires agencies to analyze regulatory options that would minimize any significant impact of a rule on small entities. Because the final rule will lessen paperwork and recordkeeping burdens and impose no significant new burdens, the agency certifies that the regulation will not have a significant economic impact on a substantial number of small entities. Therefore, under the Regulatory Flexibility Act, no further analysis is required.

The Unfunded Mandates Reform Act (Public Law 104–4) requires that agencies prepare a written statement including an assessment of anticipated costs and benefits before proposing any rule that may result in an annual expenditure by State, local, and tribal governments, in the aggregate, or by the private sector, of $100 million or more. This final rule does not impose any mandates on State, local, or tribal
DEPARTMENT OF EDUCATION

34 CFR Part 611

RIN 1840–AC65

Teacher Quality Enhancement Grants Program

AGENCY: Office of Postsecondary Education, Department of Education.

ACTION: Final regulations.

SUMMARY: The Assistant Secretary for Postsecondary Education issues regulations to implement a requirement of section 204(e) of the Higher Education Act (HEA), as amended by the Higher Education Amendments of 1998. Section 204(e) requires that students in teacher preparation programs funded under the Teacher Recruitment Program must repay scholarships provided with program funds if they do not teach in high-need local educational agencies for the period of time for which they receive scholarship assistance. These regulations also would apply to any scholarships awarded to students in teacher preparation programs funded under the State and Partnership Programs authorized in sections 202 and 203 of the HEA.

DATES: These regulations are effective January 12, 2000.

FOR FURTHER INFORMATION CONTACT: Dr. Louis Venuto, Higher Education Programs, Office of Postsecondary Education, Office of Policy, Planning, and Innovation, 1990 K Street, NW., Washington, DC 20006–8525; Telephone: (202) 502–7763. Inquiries also may be sent by e-mail to: Louis_Venuto@ed.gov or by FAX to: (202) 502–7699. If you use a telecommunications device for the deaf (TDD), you may call the Federal Information Relay Service (FIRS) at 1–800–877–8339. Individuals with disabilities may obtain this document in an alternate format (e.g., Braille, large print, audiotape, or computer diskette) on request to the contact person listed in the preceding paragraph.

SUPPLEMENTARY INFORMATION:

Background

On October 8, 1998, the President signed into law the Higher Education Amendments of 1998 (Pub. L. 105–244). Title II of this law addresses the Nation’s need to ensure that new teachers enter the classroom prepared to teach all students to high standards by authorizing, as Title II of the Higher Education Act (HEA), Teacher Quality Enhancement Grants for States and Partnerships.

The new Teacher Quality Enhancement Grants Program consists of three different competitive grant programs. Together, the State Grants Program, the Partnership Grants for Improving Teacher Preparation Program, and the Teacher Recruitment Program, these programs are designed to increase student achievement by supporting comprehensive approaches to improving teacher quality.

One key aspect of the Teacher Recruitment Grants Program is the availability of scholarships to students who are enrolled in teacher preparation programs at the grantee institutions of higher education (IHEs) (or at IHEs working with State Teacher Recruitment Program grantees), and who agree to teach in high-need school districts. As provided in section 204(e) of the HEA, in exchange for scholarship support recipients must agree to incur a contractual obligation, under terms the Department establishes, to teach in high-need LEAs for a period equivalent to the period for which they receive the scholarship.

On November 5, 1999, the Secretary published a notice of proposed rulemaking (NPRM) for this part in the Federal Register (64 FR 60632). In the preamble to the NPRM, the Secretary discussed on pages 60632 through 60638 the proposed terms and conditions of this contractual agreement. The major issues addressed by the NPRM included—

• Whether all with Teacher Recruitment Program scholarship recipients should have to meet their service obligations by teaching in high-need schools of high-need LEAs;

• The definition of a “high-need LEA” and a “high-need school” in which scholarship recipients would need to teach in order to avoid responsibility for repaying their scholarships;

• How, in order to retain the financial assistance as a scholarship, the Department will calculate the period of time in which the scholarship recipient must teach in a high-need school of a high-need LEA;

• Conditions under which the Department may defer a scholarship recipient’s service obligation;

• The amount of the scholarship recipient’s indebtedness to the Federal government for failure to meet the service obligation, terms of repayment, and any limited circumstances under which the Department would discharge this indebtedness;

• The content of the scholarship agreement that the scholarship recipient would execute;

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• Conditions under which the Department may defer a scholarship recipient’s service obligation;

• The amount of the scholarship recipient’s indebtedness to the Federal government for failure to meet the service obligation, terms of repayment, and any limited circumstances under which the Department would discharge this indebtedness;

• The content of the scholarship agreement that the scholarship recipient would execute;