new information to Chile. Finally, the revised guidance notes that FDA considers the information on this list, which is provided voluntarily with the understanding that it will be posted on FDA's Internet site and communicated to, and possibly further disseminated by, Chile, to be information that is not protected from disclosure under 5 U.S.C. 552(b)(4). Under this guidance,

FDA recommends that U.S. firms that want to be placed on the list send the following information to FDA: Name and address of the firm and the manufacturing plant; name, telephone number, and e-mail address (if available) of the contact person; a list of products presently shipped and expected to be shipped in the next 3 years; identities of agencies that inspect

the plant and the date of last inspection; plant number and copy of last inspection notice; and, if other than an FDA inspection, copy of last inspection report. FDA requests that this information be updated every 2 years.

FDA estimates the burden of this collection of information as follows:

TABLE 1.—ESTIMATED ANNUAL REPORTING BURDEN<sup>1</sup>

Activity	No. of Respondents	Annual Frequency per Response	Total Annual Responses	Hours per Response	Total Hours
New written requests to be placed on the list	15	1	15	1.5	22.5
Biannual update	55	1	55	1.0	55.0
Occasional updates	25	1	25	0.5	12.5
Total					90

<sup>&</sup>lt;sup>1</sup>There are no capital costs or operating and maintenance costs associated with this collection of information.

The estimate of the number of firms that will submit new written requests to be placed on the list, biannual updates and occasional updates is based on the FDA's experience maintaining the list over the past 3 years. The estimate of the number of hours that it will take a firm to gather the information needed to be placed on the list or update its information is based on FDA's experience with firms submitting similar requests. FDA believes that the information to be submitted will be readily available to the firms.

To date, over 110 producers have sought to be included on the list. FDA estimates that, each year, approximately 15 new firms will apply to be added to the list. We estimate that a firm will require 1.5 hours to read the guidance, gather the information needed, and to prepare a communication to FDA that contains the information and requests that the firm be placed on the list. Under the revised guidance, every 2 years each producer on the list must provide updated information in order to remain on the list. FDA estimates that each year approximately half of the firms on the list, 55 firms, will resubmit the information to remain on the list. We estimate that a firm already on the list will require 1.0 hours to biannually update and resubmit the information to FDA, including time reviewing the information and corresponding with FDA. In addition, FDA expects that, each year, approximately 25 firms will need to submit an occasional update and each firm will require 0.5 hours to prepare a communication to FDA reporting the change.

Dated: July 21, 2006.

### Jeffrey Shuren,

Assistant Commissioner for Policy.
[FR Doc. E6–12160 Filed 7–28–06; 8:45 am]
BILLING CODE 4160–01–S

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

# **Food and Drug Administration**

[Docket No. 2006D-0246]

# Draft Manufactured Food Regulatory Program Standards; Availability; Correction

**AGENCY:** Food and Drug Administration,

HHS.

**ACTION:** Notice; correction.

SUMMARY: The Food and Drug Administration is correcting a notice that appeared in the Federal Register of July 20, 2006. The document announced the availability of a draft document entitled "Manufactured Food Regulatory Program Standards." The document was published with an incorrect Internet address. This document corrects that error.

### FOR FURTHER INFORMATION CONTACT:

Beverly Kent, Division of Federal-State Relations, Food and Drug Administration, 300 Pearl St., suite 100, Buffalo, NY 14202, 716–541–0331.

SUPPLEMENTARY INFORMATION: In FR Doc. E6–11539, appearing on page 41221 in the Federal Register of Thursday, July 20, 2006, the following correction is made:

1. On page 41222, in the first column, under the "Electronic Access" caption,

the Internet address

"http:www.fda.gov.ohrms/dockets/default.htm" is corrected to read "http://www.fda.gov/ohrms/dockets/98fr/06d-0246-gdl0001.pdf".

Dated: July 25, 2006.

# Jeffrey Shuren,

Assistant Commissioner for Policy.
[FR Doc. E6–12179 Filed 7–28–06; 8:45 am]
BILLING CODE 4160–01–S

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Health Resources and Services Administration

# Cooperative Agreement for Poison Prevention Education

**AGENCY:** Health Resources and Services Administration (HRSA), HHS.

**ACTION:** Notice of Single Source Award.

**SUMMARY:** HRSA will be forming a partnership with the Home Safety Council (HSC) to collaborate on reaching America's low literacy population. Through this project, easy to read and comprehend poison prevention material will be developed and distributed to the public, poison centers, safety and injury prevention professionals, health educators, and first responders.

## FOR FURTHER INFORMATION CONTACT:

Shkeda Johnson, Senior Public Health Analyst, Healthcare Systems Bureau, Division of Healthcare Preparedness, Room 13–103, 5600 Fishers Lane, Rockville, MD 20857. Telephone: 301– 443–1210 Email: sjohnson@hrsa.gov.