TABLE 3—REGISTRANTS OF CANCELLED PRODUCTS—Continued

<table>
<thead>
<tr>
<th>EPA company No.</th>
<th>Company name and address</th>
</tr>
</thead>
<tbody>
<tr>
<td>46515</td>
<td>Celex, Division of United Industries Corp., P.O. Box 142642, St. Louis, MO 63114.</td>
</tr>
<tr>
<td>63376</td>
<td>Family Products SDN BHD, Agent: Regulatory West Co., LLC, 8203 West 20th St., Suite A, Greeley, CO 80634.</td>
</tr>
<tr>
<td>82539</td>
<td>Kerslig Candle Light, 5807 Church Hill Way, Medina, OH 44256.</td>
</tr>
<tr>
<td>83467</td>
<td>Multinational Resources, Inc., 9380 SW. 72 St., Suite B211, Miami, FL 33173.</td>
</tr>
</tbody>
</table>

III. Summary of Public Comments Received and Agency Response to Comments

The Agency received one comment indicating that all pesticides should be disallowed because of their effect on the environment. This did not merit further review of the requests.

IV. Cancellation Order

Pursuant to FIFRA section 6(f), EPA hereby approves the requested cancellation of the registrations identified in Tables 1 and 2 of Unit II. Accordingly, the Agency orders that the product registrations identified in Tables 1 and 2 of Unit II, are canceled. The effective date of the cancellations that are the subject of this notice is as follows: The cancellation for the manufacturing products, listed in Table 1 of Unit II., will be effective September 30, 2015, no use of listed manufacturing products to formulate any end use products will be permitted after December 31, 2015, and the cancellation for the end use products listed in Table 2 of Unit II., will be effective December 31, 2016. Any distribution, sale, or use of existing stocks of the products identified in Tables 1 and 2 of Unit II., in a manner inconsistent with any of the provisions for disposition of existing stocks set forth in Unit VI., will be a violation of FIFRA.

V. What is the Agency’s authority for taking this action?

Section 6(f)(1) of FIFRA provides that a registrant of a pesticide product may at any time request that any of its pesticide registrations be canceled or amended to terminate one or more uses. FIFRA further provides that, before acting on the request, EPA must publish a notice of receipt of any such request in the Federal Register. Thereafter, following the public comment period, the EPA Administrator may approve such a request. The notice of receipt for this action was published for comment in the Federal Register issue of May 29, 2013 (78 FR 32248) (FRL–9386–4). The comment period closed on June 28, 2013.

VI. Provisions for Disposition of Existing Stocks

Existing stocks are those stocks of registered pesticide products which are currently in the United States and which were packaged, labeled, and released for shipment prior to the effective date of the cancellation action. The existing stocks provisions for the products subject to this order are as follows.

The registrants may continue to sell and distribute existing stocks of manufacturing use products listed in Table 1 of Unit II., up to and including September 30, 2015, and end use products listed in Table 2 of Unit II., up to and including December 31, 2016. The following terms and conditions are applicable to existing stocks:

- No sale or distribution of listed manufacturing products by any person, other than for purposes of disposal or export, will be permitted after September 30, 2015.
- No use of listed manufacturing products to formulate end use products will be permitted after December 31, 2015.
- As of January 1, 2017, only persons other than registrants will be allowed to sell, distribute, or use existing stocks of cancelled end use products until such stocks are exhausted. Use of existing stocks will be permitted only to the extent that the use is consistent with the terms of the previously approved labeling accompanying the product used.

List of Subjects

Environmental protection, Pesticides and pests, Allethrin.

Dated: September 12, 2013.

Richard P. Keigwin, Jr., Director, Pesticide Re-Evaluation Division, Office of Pesticide Programs.

[FR Doc. 2013–22718 Filed 9–17–13; 8:45 am]
DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Disease, Disability, and Injury Prevention and Control Special Emphasis Panel (SEP): Initial Review

The meeting announced below concerns Capacity Building Assistance for High Impact HIV Prevention, Funding Opportunity Announcement (FOA) PS14–1403, Initial Review.

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92–463), the Centers for Disease Control and Prevention (CDC) announces the following meeting of the aforementioned committee:

Times and Dates: 8:00 a.m.–5:00 p.m. EST, October 31, 2013. 8:00 a.m.–1:00 p.m. EST, November 1, 2013.

Place: Centers for Disease Control and Prevention, 4770 Buford Highway, Chamblee Building 107, Rooms 1A and 1B, Atlanta, Georgia 30341

Limited teleconference access is also available. Login information is as follows:

For Public:

TOLL-FREE PHONE #: 888–899–8135
Participant passcode: BREASTCANCER
Net Conference URL: https://www.mymeetings.com/cdc/join/
Conference number: PW7128790
Audience passcode: BREASTCANCER

Public can join the event directly:

TOLL-FREE PHONE #: 888–899–8135
There is also a toll free number for anyone outside of the USA: TOLL # 1–203–827–7034
Participant passcode: BREASTCANCER

Matters To Be Discussed: The meeting will include the initial review, discussion, and evaluation of applications received in response to “Capacity Building Assistance for High Impact HIV Prevention”, FOA PS14–1403.

Contact Person for More Information:

Harriette A. Lynch, Public Health Analyst, CDC, 1600 Clifton Road NE., Mailstop E07, Atlanta, Georgia 30333, Telephone: (404) 718–8837.

Elaine L. Baker,
Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Advisory Committee on Breast Cancer in Young Women (ACBCYW)

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92–463), the Centers for Disease Control and Prevention (CDC), announces the following meeting of the aforementioned committee:

Times and Dates: 8:00 a.m.–5:00 p.m. EST, October 31, 2013.

Place: Centers for Disease Control and Prevention, 4770 Buford Highway, Chamblee Building 107, Rooms 1A and 1B, Atlanta, Georgia 30341

Limited teleconference access is also available. Login information is as follows:

For Public:

TOLL-FREE PHONE #: 888–899–8135
Participant passcode: BREASTCANCER
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Elaine L. Baker,
Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

Agency Information Collection Activities; Proposed Collection; Comment Request; Food Canning Establishment Registration, Process Filing, and Recordkeeping for Acidified Foods and Thermally Processed Low-Acid Foods in Hermetically Sealed Containers

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA or we) is announcing an opportunity for public comment on the proposed collection of certain information by the Agency. Under the Paperwork Reduction Act of 1995 (the PRA), Federal Agencies are required to publish notice in the Federal Register concerning each proposed collection of information, including each proposed extension of an existing collection of information, and to allow 60 days for public comment in response to the notice. This notice invites comments on the information collection provisions of reporting and recordkeeping requirements for firms that process acidified foods and thermally processed low-acid foods in hermetically sealed containers, and provides notice of and invites comments on our proposed revisions to the