

changes to reduce the risk from spray drift to non-target terrestrial plants. Quinlorac has not been evaluated under the EDSP. Therefore, the Agency's final registration review decision is dependent upon the result of Section 7 Endangered Species consultation with the Services, and the result of the evaluation of potential endocrine disruptor risk. Pending the outcome of these actions, EPA is planning to issue an interim registration review decision for quinlorac.

Triflumizole (Proposed Interim Decision). The registration review docket for triflumizole (EPA-HQ-OPP-2006-0115) opened in March 2007. Triflumizole is a broad spectrum, imidazole fungicide (group 3) that inhibits ergosterol biosynthesis in fungi, acting as a systemic fungicide. Triflumizole is registered for application to a number of food and non-food crops, including ornamentals in greenhouses/shade houses, interior scapes, and Christmas trees/conifers on nurseries and plantations. It is also used as a pre-plant seed piece treatment on pineapples. EPA conducted a qualitative human health risk assessment and identified occupational handler and post-application exposure risks of concern for several use scenarios. EPA is proposing additional personal protective equipment of a chemical-resistant hat to address occupational handler risks of concern when applying triflumizole with open cab air blast equipment to apple, pear, and cherry. To address post-application risks of concern, EPA is proposing to increase re-entry intervals (REIs) for grapes (table and raisin) to 1-day and hops to 3 days. The ecological risk assessment identified potential risks to listed mammals, birds, herpatofauna, freshwater fish, and aquatic estuarine-marine invertebrates; however, the only non-listed taxa of concern was chronic risk to mammals. To mitigate potential chronic risk to non-listed mammals, the registrant agreed to label changes reducing the number of applications per year for certain crops and increasing the retreatment interval (RTI) to reflect typical usage. The risk assessment for triflumizole did not come to a conclusion of "no effect" to listed species. Therefore, consultation with the Services on the potential risk of triflumizole to listed species will be necessary. Triflumizole has not been evaluated under the EDSP. Therefore, the Agency's final registration review decision is dependent upon the result of Section 7 Endangered Species consultation with the Services and the evaluation of potential endocrine

disruptor risk. Pending the outcome of these actions, EPA is planning to issue an interim registration review decision for triflumizole.

The registration review docket for a pesticide includes earlier documents related to the registration review of the case. For example, the review typically opens with a summary document, containing a Preliminary Work Plan, for public comment. A final Work Plan is placed in the docket following public comment on the initial docket. The documents in the dockets describe EPA's rationales for conducting additional risk assessments, as well as the Agency's subsequent risk findings and consideration of possible risk mitigation measures. A proposed registration review decision will be supported by the rationales included in those documents. Following public comment on a proposed decision, the Agency will issue an interim registration review decision.

The registration review program is being conducted under congressionally mandated time frames, and EPA recognizes the need both to make timely decisions and to involve the public. Section 3(g) of FIFRA (7 U.S.C. 136a(g)) required EPA to establish by regulation procedures for reviewing pesticide registrations, originally with a goal of reviewing each pesticide's registration every 15 years to ensure that a pesticide continues to meet the FIFRA standard for registration. The Agency's final rule to implement this program was issued in August 2006 and became effective in October 2006, and appears at 40 CFR part 155, subpart C. The Pesticide Registration Improvement Act of 2003 (PRIA) was amended and extended in September 2007. FIFRA, as amended by PRIA in 2007, requires EPA to complete registration review decisions by October 1, 2022, for all pesticides registered as of October 1, 2007.

The registration review final rule at 40 CFR 155.58(a) provides for a minimum 60-day public comment period on all proposed registration review decisions. This comment period is intended to provide an opportunity for public input and a mechanism for initiating any necessary amendments to the proposed decision. All comments should be submitted using the methods in **ADDRESSES**, and must be received by EPA on or before the closing date. These comments will become part of the docket for the pesticides included in the table in Unit II.A. Comments received after the close of the comment period will be marked "late." EPA is not required to consider these late comments.

The Agency will carefully consider all comments received by the closing date and will provide a "Response to Comments Memorandum" in the docket as appropriate. The final registration review decision will explain the effect that any comments had on the decision.

Background on the registration review program is provided at: <http://www2.epa.gov/pesticide-reevaluation>. Information regarding earlier documents related to the registration review of these pesticides can be found at: <http://www2.epa.gov/pesticide-reevaluation/individual-pesticides-registration-review>.

B. What is the Agency's authority for taking this action?

Section 3(g) of FIFRA (7 U.S.C. 136a(g)) and 40 CFR part 155, subpart C, provide authority for this action.

Authority: 7 U.S.C. 136 *et seq.*

Dated: September 17, 2014.

Patricia L. Parrott,

Acting Director, Pesticide Re-Evaluation Division, Office of Pesticide Programs.

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ENVIRONMENTAL PROTECTION AGENCY

[EPA-HQ-OPP-2009-1017; FRL-9916-69]

Product Cancellation Order for Certain Pesticide Registrations

AGENCY: Environmental Protection Agency (EPA).

ACTION: Notice.

SUMMARY: This notice announces EPA's order for the cancellations, voluntarily requested by the registrants and accepted by the Agency, of the products listed in Table 1 of Unit II, pursuant to the Federal Insecticide, Fungicide, and Rodenticide Act (FIFRA). This cancellation order follows an August 6, 2014 **Federal Register** Notice of Receipt of Requests from the registrants listed in Table 2 of Unit II, to voluntarily cancel these product registrations. In the August 6, 2014 notice, EPA indicated that it would issue an order implementing the cancellations, unless the Agency received substantive comments within the 30 day comment period that would merit its further review of these requests, or unless the registrants withdrew their requests. The Agency did not receive any comments on the notice. Further, the registrants did not withdraw their requests. Accordingly, EPA hereby issues in this notice a cancellation order granting the

requested cancellations. Any distribution, sale, or use of the products subject to this cancellation order is permitted only in accordance with the terms of this order, including any existing stocks provisions.

DATES: The cancellations are effective September 24, 2014.

FOR FURTHER INFORMATION CONTACT: John W. Pates, Jr., Pesticide Re-Evaluation Division (7508P), Office of Pesticide Programs, Environmental Protection Agency, 1200 Pennsylvania Ave. NW., Washington, DC 20460-0001; telephone number: (703) 308-8195; email address: pates.john@epa.gov.

SUPPLEMENTARY INFORMATION:

I. General Information

A. Does this action apply to me?

This action is directed to the public in general, and may be of interest to a

wide range of stakeholders including environmental, human health, and agricultural advocates; the chemical industry; pesticide users; and members of the public interested in the sale, distribution, or use of pesticides. Since others also may be interested, the Agency has not attempted to describe all the specific entities that may be affected by this action.

B. How can I get copies of this document and other related information?

The docket for this action, identified by docket identification (ID) number EPA-HQ-OPP-2009-1017, is available at <http://www.regulations.gov> or at the Office of Pesticide Programs Regulatory Public Docket (OPP Docket) in the Environmental Protection Agency Docket Center (EPA/DC), West William Jefferson Clinton Bldg., Rm. 3334, 1301

Constitution Ave. NW., Washington, DC 20460-0001. The Public Reading Room is open from 8:30 a.m. to 4:30 p.m., Monday through Friday, excluding legal holidays. The telephone number for the Public Reading Room is (202) 566-1744, and the telephone number for the OPP Docket is (703) 305-5805. Please review the visitor instructions and additional information about the docket available at <http://www.epa.gov/dockets>.

II. What action is the Agency taking?

This notice announces the cancellation, as requested by registrants, of products registered under FIFRA section 3. These registrations are listed in sequence by registration number in Table 1 of this unit.

TABLE 1—PRODUCT CANCELLATIONS

EPA Registration No.	Product name	Chemical name
000100-00729	Primo® Liquid	Trinexapac-ethyl.
000100-00752	Primo® WSB	Trinexapac-ethyl.
000279-09556	Intruder Residual Cylinder with Cyfluthrin	Piperonyl butoxide, pyrethrins (No inert use), and cyfluthrin.
003546-00041	Shoo-fly Flying Insect Killer	Permethrin, piperonyl butoxide, and pyrethrins (No inert use).
010807-00127	Misty Insect Repellent II	MGK 264, MGK326, and diethyl toluamide.
046386-00002	Prometrex Technical	Prometryn.
053883-00241	CSI Wipe & Spray Insecticide	Stabilene, piperonyl butoxide, and pyrethrins (No inert use).
053883-00295	CSI Folpet Technical	Folpet.
053883-00301	CSI Folpet MUP	Folpet.
062719-00601	Acetochlor Technical	Acetochlor.
071711-00022	AC 801,757 Miticide-Insecticide	Tebufenpyrad.
071711-00023	AC 801,757 3EC Miticide-Insecticide	Tebufenpyrad.
ME-080001	Nexter	Pyridaben.
PR-130002	IMI 1% G Insecticide	Imidacloprid.
PR-140001	Quali-pro Imidacloprid 1G Nursery & Greenhouse Insecticide.	Imidacloprid.
WA-860025	Drexel Dimethoate 2.67 EC	Dimethoate.

Table 2 of this unit includes the names and addresses of record for all registrants of the products in Table 1 of

this unit, in sequence by EPA company number. This number corresponds to the first part of the EPA registration

numbers of the products listed in Table 1 of this unit.

TABLE 2—REGISTRANTS OF CANCELLED PRODUCTS

EPA Company No.	Company name and address
100	Syngenta Crop Protection, LLC, 410 Swing Rd., P.O. Box 18300, Greensboro, NC 27419-8300.
279	FMC Corp. Agricultural Products Group, 1735 Market St., Rm. 1978, Philadelphia, PA 19103.
3546	Lynwood Labs, Inc., 945 Great Plain Ave., Needham, MA 02492-3004.
10807	Amrep, Inc., Agent: Zep, Inc. c/o Compliance Services, 1259 Seaboard Industrial Blvd. NW., Atlanta, GA 30318.
46386	Verolit Chemical Manufacturers, LTD, c/o/Makhteshim-Agan of North America, Inc., Agent: Makhteshim-Agan of North America, Inc., 3120 Highwoods Blvd., Suite 100, Raleigh, NC 27604.
53883, PR-130002	Control Solutions, Inc., 5903 Genoa Red Bluff Rd., Pasadena, TX 77507-1041.
62719	Dow Agrosciences, LLC, 9339 Zionsville Rd., 308/2E, Indianapolis, IN 46268-1054.
PR-140001	Makhteshim Agan of North America, Inc., d/b/a ADAMA, 3120 Highwoods Blvd., Suite 100, Raleigh, NC 27604.
71711	Nichino America, Inc., Agent: Exponent, Inc., 1150 Connecticut Ave., NW., Suite 1100, Washington, DC 20036.
ME-080001	Gowan Company, P.O. Box 5569, Yuma, AZ 85366.
WA-860025	Drexel Chemical Company, P.O. Box 13327, Memphis, TN 38113-0327.

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

During the public comment period provided, EPA received no comments in response to the August 6, 2014 **Federal Register** notice (79 FR 45803) (FRL-9914-36) announcing the Agency's receipt of the requests for voluntary cancellations of products listed in Table 1 of Unit II.

IV. Cancellation Order

Pursuant to FIFRA section 6(f), EPA hereby approves the requested cancellations of the registrations identified in Table 1 of Unit II. Accordingly, the Agency hereby orders that the product registrations identified in Table 1 of Unit II. are canceled. The effective date of the cancellations that are the subject of this notice is September 24, 2014. Any distribution, sale, or use of existing stocks of the products identified in Table 1 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** of August 6, 2014. The comment period closed on September 5, 2014.

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 products listed in Table 1 of Unit II. until September 24, 2015, which is 1 year after the publication of the Cancellation Order in the **Federal Register**. Thereafter, the registrants are prohibited from selling or distributing products listed in Table 1 of Unit II.,

except for export in accordance with FIFRA section 17, or proper disposal. Persons other than the registrants may sell, distribute, or use existing stocks of products listed in Table 1 of Unit II. until existing stocks are exhausted, provided that such sale, distribution, or use is consistent with the terms of the previously approved labeling on, or that accompanied, the canceled products.

Authority: 7 U.S.C. 136 *et seq.*

Dated: September 12, 2014.

Richard P. Keigwin, Jr.,

*Director, Pesticide Re-Evaluation Division,
Office of Pesticide Programs.*

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ENVIRONMENTAL PROTECTION AGENCY

[EPA-HQ-OPP-2009-1017; FRL-9916-41]

Product Cancellation Order for Certain Pesticide Registrations; Correction

AGENCY: Environmental Protection Agency (EPA).

ACTION: Notice; correction.

SUMMARY: EPA issued a notice in the **Federal Register** of August 13, 2014, and June 4, 2014, concerning receipt of requests to voluntarily cancel certain pesticide registrations and its follow-up cancellation order, respectively. In both notices, EPA inadvertently listed the incorrect existing stocks language for products Ronilan Manufacturer's Concentrate (EPA Reg. No. 007969-00057), Ronilan EG Fungicide (EPA Reg. No. 007969-00085), and Curalan EG Fungicide (EPA Reg. No. 007969-00224). This document corrects the existing stocks language listed in both the August 13, 2014, and June 4, 2014, **Federal Register** notices.

FOR FURTHER INFORMATION CONTACT: John W. Pates, Jr., Pesticide Re-Evaluation Division (7508P), Office of Pesticide Programs, Environmental Protection Agency, 1200 Pennsylvania Ave. NW., Washington, DC 20460-0001; telephone number: (703) 308-8195; email address: pates.john@epa.gov.

SUPPLEMENTARY INFORMATION:

I. General Information

A. Does this action apply to me?

The Agency included in the **Federal Register** notices of August 13, 2014 (79 FR 47454) (FRL 9914-00) and June 4, 2014 (79 FR 32288) (FRL 9910-97) a list of those who may be potentially affected by this action.

B. How can I get copies of this document and other related information?

The docket for this action, identified by docket identification (ID) number EPA-HQ-OPP-2009-1017, is available at <http://www.regulations.gov> or at the Office of Pesticide Programs Regulatory Public Docket (OPP Docket) in the Environmental Protection Agency Docket Center (EPA/DC), West William Jefferson Clinton Bldg., Rm. 3334, 1301 Constitution Ave. NW., Washington, DC 20460-0001. The Public Reading Room is open from 8:30 a.m. to 4:30 p.m., Monday through Friday, excluding legal holidays. The telephone number for the Public Reading Room is (202) 566-1744, and the telephone number for the OPP Docket is (703) 305-5805. Please review the visitor instructions and additional information about the docket available at <http://www.epa.gov/dockets>.

II. What does this correction do?

FR Docs. 2014-18961 and 2014-12922 published in the **Federal Register** of August 13, 2014 (79 FR 47454) (FRL 9914-00) and June 4, 2014 (79 FR 32288) (FRL 9910-97), respectively, are corrected as follows:

1. On pages 47456 and 32290, respectively, second column, under the heading B. For Products (007969-00057, 007969-00085, and 007969-00224), paragraph 1, sentences 3 and 4, correct "Thereafter, registrants, and persons other than registrants, are prohibited from selling or distributing existing stocks of products containing vinclozolin identified in Table 1 of Unit II., except for export consistent with FIFRA section 17 or for proper disposal. Existing stocks of products containing vinclozolin already in the hands of users can be used legally until such stocks are exhausted, provided that the use is consistent with the terms of the previously approved labeling on, or that accompanied, the cancelled products" to read "Thereafter, registrants are prohibited from selling and distributing existing stocks of products containing vinclozolin identified in Table 1 of Unit II., except for export consistent with FIFRA section 17 or for proper disposal. Persons other than registrants may sell, distribute, or use existing stocks of products listed in Table 1 of Unit II. until such stocks are exhausted, provided that such sale, distribution, or use is consistent with the terms of the previously approved labeling on, or that accompanied, the canceled products."

Authority: 7 U.S.C. 136 *et seq.*