FEDERAL INSECTICIDE, FUNGICIDE, AND RODENTICIDE
ACT

[Ch. 125 of the 80th Congress; Approved June 25, 1947; 61 Stat. 163]

[As Amended Through P.L. 116–8, Enacted March 08, 2019]

Currency: This publication is a compilation of the text of Chapter 125 of the 80th Congress. It was last amended by the public law listed in the As Amended Through note above and below at the bottom of each page of the pdf version and reflects current law through the date of the enactment of the public law listed at https://www.govinfo.gov/app/collection/comps/.

Note: While this publication does not represent an official version of any Federal statute, substantial efforts have been made to ensure the accuracy of its contents. The official version of Federal law is found in the United States Statutes at Large and in the United States Code. The legal effect to be given to the Statutes at Large and the United States Code is established by statute (1 U.S.C. 112, 204).

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AN ACT To regulate the marketing of economic poisons and devices, and for other purposes.

Be it enacted by the Senate and House of Representatives of the United States of America in Congress assembled,

SECTION 1. [7 U.S.C. prec. 121] SHORT TITLE AND TABLE OF CONTENTS.

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For purposes of this Act—

(a) ACTIVE INGREDIENT.—The term “active ingredient” means—

(1) in the case of a pesticide other than a plant regulator, defoliant, desiccant, or nitrogen stabilizer, an ingredient which will prevent, destroy, repel, or mitigate any pest;

(2) in the case of a plant regulator, an ingredient which, through physiological action, will accelerate or retard the rate of growth or rate of maturation or otherwise alter the behavior of ornamental or crop plants or the product thereof;

(3) in the case of a defoliant, an ingredient which will cause the leaves or foliage to drop from a plant;

(4) in the case of a desiccant, an ingredient which will artificially accelerate the drying of plant tissue; and

(5) in the case of a nitrogen stabilizer, an ingredient which will prevent or hinder the process of nitrification, denitrification, ammonia volatilization, or urease production through action affecting soil bacteria.

(b) ADMINISTRATOR.—The term “Administrator” means the Administrator of the Environmental Protection Agency.

(c) ADULTERATED.—The term “adulterated” applies to any pesticide if—

(1) its strength or purity falls below the professed standard of quality as expressed on its labeling under which it is sold;
(2) any substance has been substituted wholly or in part for the pesticide; or
(3) any valuable constituent of the pesticide has been wholly or in part abstracted.

(d) ANIMAL.—The term “animal” means all vertebrate and invertebrate species, including but not limited to man and other mammals, birds, fish, and shellfish.

(e) CERTIFIED APPLICATOR, ETC.—
(1) CERTIFIED APPLICATOR.—The term “certified applicator” means any individual who is certified under section 11 as authorized to use or supervise the use of any pesticide which is classified for restricted use. Any applicator who holds or applies registered pesticides, or uses dilutions of registered pesticides consistent with subsection (ee), only to provide a service of controlling pests without delivering any unapplied pesticide to any person so served is not deemed to be a seller or distributor of pesticides under this Act.
(2) PRIVATE APPLICATOR.—The term “private applicator” means a certified applicator who uses or supervises the use of any pesticide which is classified for restricted use for purposes of producing any agricultural commodity on property owned or rented by the applicator or the applicator’s employer or (if applied without compensation other than trading of personal services between producers of agricultural commodities) on the property of another person.
(3) COMMERCIAL APPLICATOR.—The term “commercial applicator” means an applicator (whether or not the applicator is a private applicator with respect to some uses) who uses or supervises the use of any pesticide which is classified for restricted use for any purpose or on any property other than as provided by paragraph (2).
(4) UNDER THE DIRECT SUPERVISION OF A CERTIFIED APPLICATOR.—Unless otherwise prescribed by its labeling, a pesticide shall be considered to be applied under the direct supervision of a certified applicator if it is applied by a competent person acting under the instructions and control of a certified applicator who is available if and when needed, even though such certified applicator is not physically present at the time and place the pesticide is applied.

(f) DEFOLIANT.—The term “defoliant” means any substance or mixture of substances intended for causing the leaves or foliage to drop from a plant, with or without causing abscission.

(g) DESICCANT.—The term “desiccant” means any substance or mixture of substances intended for artificially accelerating the drying of plant tissue.

(h) DEVICE.—The term “device” means any instrument or contrivance (other than a firearm) which is intended for trapping, destroying, repelling, or mitigating any pest or any other form of plant or animal life (other than man and other than bacteria, virus, or other microorganism on or in living man or other living animals); but not including equipment used for the application of pesticides when sold separately therefrom.
(i) DISTRICT COURT.—The term “district court” means a United States district court, the District Court of Guam, the District Court of the Virgin Islands, and the highest court of American Samoa.

(j) ENVIRONMENT.—The term “environment” includes water, air, land, and all plants and man and other animals living therein, and the interrelationships which exist among these.

(k) FUNGUS.—The term “fungus” means any non-chlorophyll-bearing thallophyte (that is, any non-chlorophyll-bearing plant of a lower order than mosses and liverworts), as for example, rust, smut, mildew, mold, yeast, and bacteria, except those on or in living man or other animals and those on or in processed food, beverages, or pharmaceuticals.

(l) IMMINENT HAZARD.—The term “imminent hazard” means a situation which exists when the continued use of a pesticide during the time required for cancellation proceeding would be likely to result in unreasonable adverse effects on the environment or will involve unreasonable hazard to the survival of a species declared endangered by the Secretary of the Interior under Public Law 91–135.

(m) INERT INGREDIENT.—The term “inert ingredient” means an ingredient which is not active.

(n) INGREDIENT STATEMENT.—The term “ingredient statement” means a statement which contains—

1. the name and percentage of each active ingredient, and the total percentage of all inert ingredients, in the pesticide; and
2. if the pesticide contains arsenic in any form, a statement of the percentages of total and water soluble arsenic, calculated as elementary arsenic.

(o) INSECT.—The term “insect” means any of the numerous small invertebrate animals generally having the body more or less obviously segmented, for the most part belonging to the class insecta, comprising six-legged, usually winged forms, as for example, beetles, bugs, bees, flies, and to other allied classes of arthropods whose members are wingless and usually have more than six legs, as for example, spiders, mites, ticks, centipedes, and woodlice.

(p) LABEL AND LABELING.—

1. LABEL.—The term “label” means the written, printed, or graphic matter on, or attached to, the pesticide or device or any of its containers or wrappers.
2. LABELING.—The term “labeling” means all labels and all other written, printed, or graphic matter—

A. accompanying the pesticide or device at any time; or
B. to which reference is made on the label or in literature accompanying the pesticide or device, except to current official publications of the Environmental Protection Agency, the United States Departments of Agriculture and Interior, the Department of Health and Human Services, State experiment stations, State agricultural colleges, and other similar Federal or State institutions or agencies authorized by law to conduct research in the field of pesticides.
(q) **Misbranded.—**

(1) A pesticide is misbranded if—

(A) its labeling bears any statement, design, or graphic representation relative thereto or to its ingredients which is false or misleading in any particular;

(B) it is contained in a package or other container or wrapping which does not conform to the standards established by the Administrator pursuant to section 25(c)(3);

(C) it is an imitation of, or is offered for sale under the name of, another pesticide;

(D) its label does not bear the registration number assigned under section 7 to each establishment in which it was produced;

(E) any word, statement, or other information required by or under authority of this Act to appear on the label or labeling is not prominently placed thereon with such conspicuousness (as compared with other words, statements, designs, or graphic matter in the labeling) and in such terms as to render it likely to be read and understood by the ordinary individual under customary conditions of purchase and use;

(F) the labeling accompanying it does not contain directions for use which are necessary for effecting the purpose for which the product is intended and if complied with, together with any requirements imposed under section 3(d) of this Act, are adequate to protect health and the environment;

(G) the label does not contain a warning or caution statement which may be necessary and if complied with, together with any requirements imposed under section 3(d) of this Act, is adequate to protect health and the environment; or

(H) in the case of a pesticide not registered in accordance with section 3 of this Act and intended for export, the label does not contain, in words prominently placed thereon with such conspicuousness (as compared with other words, statements, designs, or graphic matter in the labeling) as to render it likely to be noted by the ordinary individual under customary conditions of purchase and use, the following: “Not Registered for Use in the United States of America”.

(2) A pesticide is misbranded if—

(A) the label does not bear an ingredient statement on that part of the immediate container (and on the outside container or wrapper of the retail package, if there be one, through which the ingredient statement on the immediate container cannot be clearly read) which is presented or displayed under customary conditions of purchase, except that a pesticide is not misbranded under this subparagraph if—

(i) the size or form of the immediate container, or the outside container or wrapper of the retail package, makes it impracticable to place the ingredient state-
ment on the part which is presented or displayed under customary conditions of purchase; and

(ii) the ingredient statement appears prominently on another part of the immediate container, or outside container or wrapper, permitted by the Administrator;

(B) the labeling does not contain a statement of the use classification under which the product is registered;

(C) there is not affixed to its container, and to the outside container or wrapper of the retail package, if there be one, through which the required information on the immediate container cannot be clearly read, a label bearing—

(i) the name and address of the producer, registrant, or person for whom produced;

(ii) the name, brand, or trademark under which the pesticide is sold;

(iii) the net weight or measure of the content, except that the Administrator may permit reasonable variations; and

(iv) when required by regulation of the Administrator to effectuate the purposes of this Act, the registration number assigned to the pesticide under this Act, and the use classification; and

(D) the pesticide contains any substance or substances in quantities highly toxic to man, unless the label shall bear, in addition to any other matter required by this Act—

(i) the skull and crossbones;

(ii) the word “poison” prominently in red on a background of distinctly contrasting color; and

(iii) a statement of a practical treatment (first aid or otherwise) in case of poisoning by the pesticide.

(r) NEMATODE.—The term “nematode” means invertebrate animals of the phylum nemathelminthes and class nematoda, that is, unsegmented round worms with elongated, fusiform, or saclike bodies covered with cuticle, and inhabiting soil, water, plants, or plant parts; may also be called nemas or eelworms.

(s) PERSON.—The term “person” means any individual, partnership, association, corporation, or any organized group of persons whether incorporated or not.

(t) PEST.—The term “pest” means (1) any insect, rodent, nematode, fungus, weed, or (2) any other form of terrestrial or aquatic plant or animal life or virus, bacteria, or other micro-organism (except viruses, bacteria, or other micro-organisms on or in living man or other living animals) which the Administrator declares to be a pest under section 25(c)(1).

(u) PESTICIDE.—The term “pesticide” means (1) any substance or mixture of substances intended for preventing, destroying, repelling, or mitigating any pest, (2) any substance or mixture of substances intended for use as a plant regulator, defoliant, or desiccant, and (3) any nitrogen stabilizer, except that the term “pesticide” shall not include any article that is a “new animal drug” within the meaning of section 201(w) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 321(w)), that has been determined by the Secretary of Health and Human Services not to be a new ani-
mal drug by a regulation establishing conditions of use for the article, or that is an animal feed within the meaning of section 201(x) of such Act (21 U.S.C. 321(x)) bearing or containing a new animal drug. The term “pesticide” does not include liquid chemical sterilant products (including any sterilant or subordinate disinfectant claims on such products) for use on a critical or semi-critical device, as defined in section 201 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 321). For purposes of the preceding sentence, the term “critical device” includes any device which is introduced directly into the human body, either into or in contact with the bloodstream or normally sterile areas of the body and the term “semi-critical device” includes any device which contacts intact mucous membranes but which does not ordinarily penetrate the blood barrier or otherwise enter normally sterile areas of the body.

(v) PLANT REGULATOR.—The term “plant regulator” means any substance or mixture of substances intended, through physiological action, for accelerating or retarding the rate of growth or rate of maturation, or for otherwise altering the behavior of plants or the produce thereof, but shall not include substances to the extent that they are intended as plant nutrients, trace elements, nutritional chemicals, plant inoculants, and soil amendments. Also, the term “plant regulator” shall not be required to include any of such of those nutrient mixtures or soil amendments as are commonly known as vitamin-hormone horticultural products, intended for improvement, maintenance, survival, health, and propagation of plants, and as are not for pest destruction and are nontoxic, non-poisonous in the undiluted packaged concentration.

(w) PRODUCER AND PRODUCE.—The term “producer” means the person who manufacturers, prepares, compounds, propagates, or processes any pesticide or device or active ingredient used in producing a pesticide. The term “produce” means to manufacture, prepare, compound, propagate, or process any pesticide or device or active ingredient used in producing a pesticide. The dilution by individuals of formulated pesticides for their own use and according to the directions on registered labels shall not of itself result in such individuals being included in the definition of “producer” for the purposes of this Act.

(x) PROTECT HEALTH AND THE ENVIRONMENT.—The terms “protect health and the environment” and “protection of health and the environment” mean protection against any unreasonable adverse effects on the environment.

(y) REGISTRANT.—The term “registrant” means a person who has registered any pesticide pursuant to the provisions of this Act.

(z) REGISTRATION.—The term “registration” includes reregistration.

(aa) STATE.—The term “State” means a State, the District of Columbia, the Commonwealth of Puerto Rico, the Virgin Islands, Guam, the Trust Territory of the Pacific Islands, and American Samoa.

(bb) UNREASONABLE ADVERSE EFFECTS ON THE ENVIRONMENT.—The term “unreasonable adverse effects on the environment” means (1) any unreasonable risk to man or the environment, taking into account the economic, social, and environmental costs and benefits of the use of any pesticide, or (2) a human dietary risk
from residues that result from a use of a pesticide in or on any food inconsistent with the standard under section 408 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 346a). The Administrator shall consider the risks and benefits of public health pesticides separate from the risks and benefits of other pesticides. In weighing any regulatory action concerning a public health pesticide under this Act, the Administrator shall weigh any risks of the pesticide against the health risks such as the diseases transmitted by the vector to be controlled by the pesticide.

(cc) **WEED.—**The term “weed” means any plant which grows where not wanted.

(dd) **ESTABLISHMENT.—**The term “establishment” means any place where a pesticide or device or active ingredient used in producing a pesticide is produced, or held, for distribution or sale.

(ee) **TO USE ANY REGISTERED PESTICIDE IN A MANNER INCONSISTENT WITH ITS LABELING.—**The term “to use any registered pesticide in a manner inconsistent with its labeling” means to use any registered pesticide in a manner not permitted by the labeling, except that the term shall not include (1) applying a pesticide at any dosage, concentration, or frequency less than that specified on the labeling unless the term shall not include any target pest not specified on the labeling if the application is to the crop, animal, or site specified on the labeling, unless the Administrator has required that the labeling specifically state that the pesticide may be used only for the pests specified on the labeling after the Administrator has determined that the use of the pesticide against other pests would cause an unreasonable adverse effect on the environment, (3) employing any method of application not prohibited by the labeling unless the labeling specifically states that the product may be applied only by the methods specified on the labeling, (4) mixing a pesticide or pesticides with a fertilizer when such mixture is not prohibited by the labeling, (5) any use of a pesticide in conformance with section 5, 18, or 24 of this Act, or (6) any use of a pesticide in a manner that the Administrator determines to be consistent with the purposes of this Act. After March 31, 1979, the term shall not include the use of a pesticide for agricultural or forestry purposes at a dilution less than label dosage unless before or after that date the Administrator issues a regulation or advisory opinion consistent with the study provided for in section 27(b) of the Federal Pesticide Act of 1978, which regulation or advisory opinion specifically requires the use of definite amounts of dilution.

(ff) **OUTSTANDING DATA REQUIREMENT.—**

(1) **IN GENERAL.—**The term “outstanding data requirement” means a requirement for any study, information, or data that is necessary to make a determination under section 3(c)(5) and which study, information, or data—

(A) has not been submitted to the Administrator; or

(B) if submitted to the Administrator, the Administrator has determined must be resubmitted because it is
not valid, complete, or adequate to make a determination under section 3(c)(5) and the regulations and guidelines issued under such section.

(2) FACTORS.—In making a determination under paragraph (1)(B) respecting a study, the Administrator shall examine, at a minimum, relevant protocols, documentation of the conduct and analysis of the study, and the results of the study to determine whether the study and the results of the study fulfill the data requirement for which the study was submitted to the Administrator.

(gg) TO DISTRIBUTE OR SELL.—The term “to distribute or sell” means to distribute, sell, offer for sale, hold for distribution, hold for sale, hold for shipment, ship, deliver for shipment, release for shipment, or receive and (having so received) deliver or offer to deliver. The term does not include the holding or application of registered pesticides or use dilutions thereof by any applicator who provides a service of controlling pests without delivering any unapplied pesticide to any person so served.

(hh) NITROGEN STABILIZER.—The term “nitrogen stabilizer” means any substance or mixture of substances intended for preventing or hindering the process of nitrification, denitrification, ammonia volatilization, or urease production through action upon soil bacteria. Such term shall not include—

(1) dicyandiamide;
(2) ammonium thiosulfate; or
(3) any substance or mixture of substances.—
    (A) that was not registered pursuant to section 3 prior to January 1, 1992; and
    (B) that was in commercial agronomic use prior to January 1, 1992, with respect to which after January 1, 1992, the distributor or seller of the substance or mixture has made no specific claim of prevention or hindering of the process of nitrification, denitrification, ammonia volatilization urease production regardless of the actual use or purpose for, or future use or purpose for, the substance or mixture.

Statements made in materials required to be submitted to any State legislative or regulatory authority, or required by such authority to be included in the labeling or other literature accompanying any such substance or mixture shall not be deemed a specific claim within the meaning of this subsection.

(jj) MAINTENANCE APPLICATOR.—The term “maintenance applicator” means any individual who, in the principal course of such individual’s employment, uses, or supervises the use of, a pesticide not classified for restricted use (other than a ready to use consumer products pesticide); for the purpose of providing structural pest control or lawn pest control including janitors, general maintenance personnel, sanitation personnel, and grounds maintenance personnel. The term “maintenance applicator” does not include private applicators as defined in section 2(e)(2); individuals who use antimicrobial pesticides, sanitizers or disinfectants; individuals em-

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4So in original (as added by sec. 120 of P.L. 104–170). This subsection should probably be “(ii)” and subsequent subsections should be redesignated accordingly.
ployed by Federal, State, and local governments or any political subdivisions thereof, or individuals who use pesticides not classified for restricted use in or around their homes, boats, sod farms, nurseries, greenhouses, or other noncommercial property.

(kk) **SERVICE TECHNICIAN.**—The term “service technician” means any individual who uses or supervises the use of pesticides (other than a ready to use consumer products pesticide) for the purpose of providing structural pest control or lawn pest control on the property of another for a fee. The term “service technician” does not include individuals who use antimicrobial pesticides, sanitizers or disinfectants; or who otherwise apply ready to use consumer products pesticides.

(ll) **MINOR USE.**—The term “minor use” means the use of a pesticide on an animal, on a commercial agricultural crop or site, or for the protection of public health where—

1. the total United States acreage for the crop is less than 300,000 acres, as determined by the Secretary of Agriculture; or
2. the Administrator, in consultation with the Secretary of Agriculture, determines that, based on information provided by an applicant for registration or a registrant, the use does not provide sufficient economic incentive to support the initial registration or continuing registration of a pesticide for such use and—
   (A) there are insufficient efficacious alternative registered pesticides available for the use;
   (B) the alternatives to the pesticide use pose greater risks to the environment or human health;
   (C) the minor use pesticide plays or will play a significant part in managing pest resistance; or
   (D) the minor use pesticide plays or will play a significant part in an integrated pest management program.

The status as a minor use under this subsection shall continue as long as the Administrator has not determined that, based on existing data, such use may cause an unreasonable adverse effect on the environment and the use otherwise qualifies for such status.

(mm) **ANTIMICROBIAL PESTICIDE.**—

1. **IN GENERAL.**—The term “antimicrobial pesticide” means a pesticide that—
   (A) is intended to—
      (i) disinfect, sanitize, reduce, or mitigate growth or development of microbiological organisms; or
      (ii) protect inanimate objects, industrial processes or systems, surfaces, water, or other chemical substances from contamination, fouling, or deterioration caused by bacteria, viruses, fungi, protozoa, algae, or slime; and
   (B) in the intended use is exempt from, or otherwise not subject to, a tolerance under section 408 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 346a and 348) or a food additive regulation under section 409 of such Act.

2. **EXCLUDED PRODUCTS.**—The term “antimicrobial pesticide” does not include —

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September 3, 2019

As Amended Through P.L. 116-8, Enacted March 08, 2019
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(A) a wood preservative or antifouling paint product for which a claim of pesticidal activity other than or in addition to an activity described in paragraph (1) is made;  
(B) an agricultural fungicide product; or  
(C) an aquatic herbicide product.

(3) INCLUDED PRODUCTS.—The term “antimicrobial pesticide” does include any other chemical sterilant product (other than liquid chemical sterilant products exempt under sub-section (u)), any other disinfectant product, any other industrial microbiocide product, and any other preservative product that is not excluded by paragraph (2).

(nn) PUBLIC HEALTH PESTICIDE.—The term “public health pesticide” means any minor use pesticide product registered for use and used predominantly in public health programs for vector control or for other recognized health protection uses, including the prevention or mitigation of viruses, bacteria, or other microorganisms (other than viruses, bacteria, or other microorganisms on or in living man or other living animal) that pose a threat to public health.

(o) VECTOR.—The term “vector” means any organism capable of transmitting the causative agent of human disease or capable of producing human discomfort or injury, including mosquitoes, flies, fleas, cockroaches, or other insects and ticks, mites, or rats.


(a) REQUIREMENT OF REGISTRATION.—Except as provided by this Act, no person in any State may distribute or sell to any person any pesticide that is not registered under this Act. To the extent necessary to prevent unreasonable adverse effects on the environment, the Administrator may by regulation limit the distribution, sale, or use in any State of any pesticide that is not registered under this Act and that is not the subject of an experimental use permit under section 5 or an emergency exemption under section 18.

(b) EXEMPTIONS.—A pesticide which is not registered with the Administrator may be transferred if—

(1) the transfer is from one registered establishment to another registered establishment operated by the same producer solely for packaging at the second establishment or for use as a constituent part of another pesticide produced at the second establishment; or

(2) the transfer is pursuant to and in accordance with the requirements of an experimental use permit.

(c) PROCEDURE FOR REGISTRATION.—

(1) STATEMENT REQUIRED.—Each applicant for registration of a pesticide shall file with the Administrator a statement which includes—

(A) the name and address of the applicant and of any other person whose name will appear on the labeling;  
(B) the name of the pesticide;  
(C) a complete copy of the labeling of the pesticide, a statement of all claims to be made for it, and any directions for its use;  
(D) the complete formula of the pesticide;
(E) a request that the pesticide be classified for general use or for restricted use, or for both; and

(F) except as otherwise provided in paragraph (2)(D), if requested by the Administrator, a full description of the tests made and the results thereof upon which the claims are based, or alternatively a citation to data that appear in the public literature or that previously had been submitted to the Administrator and that the Administrator may consider in accordance with the following provisions:

(i) With respect to pesticides containing active ingredients that are initially registered under this Act after the date of enactment of the Federal Pesticide Act of 1978 [September 30, 1978], data submitted to support the application for the original registration of the pesticide, or an application for an amendment adding any new use to the registration and that pertains solely to such new use, shall not, without the written permission of the original data submitter, be considered by the Administrator to support an application by another person during a period of ten years following the date the Administrator first registers the pesticide, except that such permission shall not be required in the case of defensive data.

(ii) The period of exclusive data use provided under clause (i) shall be extended 1 additional year for each 3 minor uses registered after the date of enactment of this clause [Aug. 3, 1996] and within 7 years of the commencement of the exclusive use period, up to a total of 3 additional years for all minor uses registered by the Administrator if the Administrator, in consultation with the Secretary of Agriculture, determines that, based on information provided by an applicant for registration or a registrant, that—

(I) there are insufficient efficacious alternative registered pesticides available for the use;

(II) the alternatives to the minor use pesticide pose greater risks to the environment or human health;

(III) the minor use pesticide plays or will play a significant part in managing pest resistance; or

(IV) the minor use pesticide plays or will play a significant part in an integrated pest management program.

The registration of a pesticide for a minor use on a crop grouping established by the Administrator shall be considered for purposes of this clause 1 minor use for each representative crop for which data are provided in the crop grouping. Any additional exclusive use period under this clause shall be modified as appropriate or terminated if the registrant voluntarily cancels the product or deletes from the registration the minor uses which formed the basis for the extension of the additional exclusive use period or if the Ad
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ministrator determines that the registrant is not actually marketing the product for such minor uses.

(iii) Except as otherwise provided in clause (i), with respect to data submitted after December 31, 1969, by an applicant or registrant to support an application for registration, experimental use permit, or amendment adding a new use to an existing registration, to support or maintain in effect an existing registration, or for reregistration, the Administrator may, without the permission of the original data submitter, consider any such item of data in support of an application by any other person (hereinafter in this subparagraph referred to as the “applicant”) within the fifteen-year period following the date the data were originally submitted only if the applicant has made an offer to compensate the original data submitter and submitted such offer to the Administrator accompanied by evidence of delivery to the original data submitter of the offer. The terms and amount of compensation may be fixed by agreement between the original data submitter and the applicant, or, failing such agreement, binding arbitration under this subparagraph. If, at the end of ninety days after the date of delivery to the original data submitter of the offer to compensate, the original data submitter and the applicant have neither agreed on the amount and terms of compensation nor on a procedure for reaching an agreement on the amount and terms of compensation, either person may initiate binding arbitration proceedings by requesting the Federal Mediation and Conciliation Service to appoint an arbitrator from the roster of arbitrators maintained by such Service. The procedure and rules of the Service shall be applicable to the selection of such arbitrator and to such arbitration proceedings, and the findings and determination of the arbitrator shall be final and conclusive, and no official or court of the United States shall have power or jurisdiction to review any such findings and determination, except for fraud, misrepresentation, or other misconduct by one of the parties to the arbitration or the arbitrator where there is a verified complaint with supporting affidavits attesting to specific instances of such fraud, misrepresentation, or other misconduct. The parties to the arbitration shall share equally in the payment of the fee and expenses of the arbitrator. If the Administrator determines that an original data submitter has failed to participate in a procedure for reaching an agreement or in an arbitration proceeding as required by this subparagraph, or failed to comply with the terms of an agreement or arbitration decision concerning compensation under this subparagraph, the original data submitter shall forfeit the right to compensation for the use of the data in support of the application. Notwithstanding any other provision of this
Act, if the Administrator determines that an applicant has failed to participate in a procedure for reaching an agreement or in an arbitration proceeding as required by this subparagraph, or failed to comply with the terms of an agreement or arbitration decision concerning compensation under this subparagraph, the Administrator shall deny the application or cancel the registration of the pesticide in support of which the data were used without further hearing. Before the Administrator takes action under either of the preceding two sentences, the Administrator shall furnish to the affected person, by certified mail, notice of intent to take action and allow fifteen days from the date of delivery of the notice for the affected person to respond. If a registration is denied or canceled under this subparagraph, the Administrator may make such order as the Administrator deems appropriate concerning the continued sale and use of existing stocks of such pesticide. Registration action by the Administrator shall not be delayed pending the fixing of compensation.

(iv) After expiration of any period of exclusive use and any period for which compensation is required for the use of an item of data under clauses (i), (ii), and (iii), the Administrator may consider such item of data in support of an application by any other applicant without the permission of the original data submitter and without an offer having been received to compensate the original data submitter for the use of such item of data.

(v) The period of exclusive use provided under clause (ii) shall not take effect until 1 year after enactment of this clause [Aug. 3, 1996], except where an applicant or registrant is applying for the registration of a pesticide containing an active ingredient not previously registered.

(vi) With respect to data submitted after the date of enactment of this clause [Aug. 3, 1996] by an applicant or registrant to support an amendment adding a new use to an existing registration that does not retain any period of exclusive use, if such data relates solely to a minor use of a pesticide, such data shall not, without the written permission of the original data submitter, be considered by the Administrator to support an application for a minor use by another person during the period of 10 years following the date of submission of such data. The applicant or registrant at the time the new minor use is requested shall notify the Administrator that to the best of their knowledge the exclusive use period for the pesticide has expired and that the data pertaining solely to the minor use of a pesticide is eligible for the provisions of this paragraph. If the minor use registration which is supported by data submitted pursuant to this subsection...
is voluntarily canceled or if such data are subsequently used to support a nonminor use, the data shall no longer be subject to the exclusive use provisions of this clause but shall instead be considered by the Administrator in accordance with the provisions of clause (i), as appropriate.

(G) If the applicant is requesting that the registration or amendment to the registration of a pesticide be expedited, an explanation of the basis for the request must be submitted, in accordance with paragraph (10) of this subsection.

(2) DATA IN SUPPORT OF REGISTRATION.—

(A) IN GENERAL.—The Administrator shall publish guidelines specifying the kinds of information which will be required to support the registration of a pesticide and shall revise such guidelines from time to time. If thereafter the Administrator requires any additional kind of information under subparagraph (B) of this paragraph, the Administrator shall permit sufficient time for applicants to obtain such additional information. The Administrator, in establishing standards for data requirements for the registration of pesticides with respect to minor uses, shall make such standards commensurate with the anticipated extent of use, pattern of use, the public health and agricultural need for such minor use, and the level and degree of potential beneficial or adverse effects on man and the environment. The Administrator shall not require a person to submit, in relation to a registration or reregistration of a pesticide for minor agricultural use under this Act, any field residue data from a geographic area where the pesticide will not be registered for such use. In the development of these standards, the Administrator shall consider the economic factors of potential national volume of use, extent of distribution, and the impact of the cost of meeting the requirements on the incentives for any potential registrant to undertake the development of the required data. Except as provided by section 10, within 30 days after the Administrator registers a pesticide under this Act, the Administrator shall make available to the public the data called for in the registration statement together with such other scientific information as the Administrator deems relevant to the Administrator’s decision.

(B) ADDITIONAL DATA.—(i) If the Administrator determines that additional data are required to maintain in effect an existing registration of a pesticide, the Administrator shall notify all existing registrants of the pesticide to which the determination relates and provide a list of such registrants to any interested person.

(ii) Each registrant of such pesticide shall provide evidence within ninety days after receipt of notification that it is taking appropriate steps to secure the additional data that are required. Two or more registrants may agree to develop jointly, or to share in the cost of developing, such data if they agree and advise the Administrator of their in-
tent within ninety days after notification. Any registrant who agrees to share in the cost of producing the data shall be entitled to examine and rely upon such data in support of maintenance of such registration. The Administrator shall issue a notice of intent to suspend the registration of a pesticide in accordance with the procedures prescribed by clause (iv) if a registrant fails to comply with this clause.

(iii) If, at the end of sixty days after advising the Administrator of their agreement to develop jointly, or share in the cost of developing data, the registrants have not further agreed on the terms of the data development arrangement or on a procedure for reaching such agreement, any of such registrants may initiate binding arbitration proceedings by requesting the Federal Mediation and Conciliation Service to appoint an arbitrator from the roster of arbitrators maintained by such Service. The procedure and rules of the Service shall be applicable to the selection of such arbitrator and to such arbitration proceedings, and the findings and determination of the arbitrator shall be final and conclusive, and no official or court of the United States shall have power or jurisdiction to review any such findings and determination, except for fraud, misrepresentation, or other misconduct by one of the parties to the arbitration or the arbitrator where there is a verified complaint with supporting affidavits attesting to specific instances of such fraud, misrepresentation, or other misconduct. All parties to the arbitration shall share equally in the payment of the fee and expenses of the arbitrator. The Administrator shall issue a notice of intent to suspend the registration of a pesticide in accordance with the procedures prescribed by clause (iv) if a registrant fails to comply with this clause.

(iv) Notwithstanding any other provision of this Act, if the Administrator determines that a registrant, within the time required by the Administrator, has failed to take appropriate steps to secure the data required under this subparagraph, to participate in a procedure for reaching agreement concerning a joint data development arrangement under this subparagraph or in an arbitration proceeding as required by this subparagraph, or to comply with the terms of an agreement or arbitration decision concerning a joint data development arrangement under this subparagraph, the Administrator may issue a notice of intent to suspend such registrant’s registration of the pesticide for which additional data is required. The Administrator may include in the notice of intent to suspend such provisions as the Administrator deems appropriate concerning the continued sale and use of existing stocks of such pesticide. Any suspension proposed under this subparagraph shall become final and effective at the end of thirty days from receipt by the registrant of the notice of intent to suspend, unless during that time a request for hearing is made by a person adversely affected by the no-
tice or the registrant has satisfied the Administrator that the registrant has complied fully with the requirements that served as a basis for the notice of intent to suspend. If a hearing is requested, a hearing shall be conducted under section 6(d) of this Act. The only matters for resolution at that hearing shall be whether the registrant has failed to take the action that served as the basis for the notice of intent to suspend the registration of the pesticide for which additional data is required, and whether the Administrator's determination with respect to the disposition of existing stocks is consistent with this Act. If a hearing is held, a decision after completion of such hearing shall be final. Notwithstanding any other provision of this Act, a hearing shall be held and a determination made within seventy-five days after receipt of a request for such hearing. Any registration suspended under this subparagraph shall be reinstated by the Administrator if the Administrator determines that the registrant has complied fully with the requirements that served as a basis for the suspension of the registration.

(v) Any data submitted under this subparagraph shall be subject to the provisions of paragraph (1)(D). Whenever such data are submitted jointly by two or more registrants, an agent shall be agreed on at the time of the joint submission to handle any subsequent data compensation matters for the joint submitters of such data.

(vi) Upon the request of a registrant the Administrator shall, in the case of a minor use, extend the deadline for the production of residue chemistry data under this subparagraph for data required solely to support that minor use until the final deadline for submission of data under section 4 for the other uses of the pesticide established as of the date of enactment of the Food Quality Protection Act of 1996 [Aug. 3, 1996], if—

(I) the data to support other uses of the pesticide on a food are being provided;

(II) the registrant, in submitting a request for such an extension, provides a schedule, including interim dates to measure progress, to assure that the data production will be completed before the expiration of the extension period;

(III) the Administrator has determined that such extension will not significantly delay the Administrator's schedule for issuing a reregistration eligibility determination required under section 4; and

(IV) the Administrator has determined that based on existing data, such extension would not significantly increase the risk of any unreasonable adverse effect on the environment. If the Administrator grants an extension under this clause, the Administrator shall monitor the development of the data and
shall ensure that the registrant is meeting the schedule for the production of the data. If the Administrator determines that the registrant is not meeting or has not met the schedule for the production of such data, the Administrator may proceed in accordance with clause (iv) regarding the continued registration of the affected products with the minor use and shall inform the public of such action. Notwithstanding the provisions of this clause, the Administrator may take action to modify or revoke the extension under this clause if the Administrator determines that the extension for the minor use may cause an unreasonable adverse effect on the environment. In such circumstance, the Administrator shall provide, in writing to the registrant, a notice revoking the extension of time for submission of data. Such data shall instead be due in accordance with the date established by the Administrator for the submission of the data.

(vii) If the registrant does not commit to support a specific minor use of the pesticide, but is supporting and providing data in a timely and adequate fashion to support uses of the pesticide on a food, or if all uses of the pesticide are nonfood uses and the registrant does not commit to support a specific minor use of the pesticide but is supporting and providing data in a timely and adequate fashion to support other nonfood uses of the pesticide, the Administrator, at the written request of the registrant, shall not take any action pursuant to this clause in regard to such unsupported minor use until the final deadline established as of the date of enactment of the Food Quality Protection Act of 1996 [Aug. 3, 1996], for the submission of data under section 4 for the supported uses identified pursuant to this clause unless the Administrator determines that the absence of the data is significant enough to cause human health or environmental concerns. On the basis of such determination, the Administrator may refuse the request for extension by the registrant. Upon receipt of the request from the registrant, the Administrator shall publish in the Federal Register a notice of the receipt of the request and the effective date upon which the uses not being supported will be voluntarily deleted from the registration pursuant to section 6(f)(1). If the Administrator grants an extension pursuant to section 6(f)(1). If the Administrator grants an extension under this clause, the Administrator shall monitor the development of the data for the uses being supported and shall ensure that the registrant is meeting the schedule for the production of such data. If the Administrator determines that the registrant is not meeting or has not met the schedule for the production of such data, the Administrator may proceed in accordance with clause (iv) of this subparagraph regarding the continued registration of the affected products with the minor and other uses and shall inform the public of such action in accordance with section 6(f)(2). Notwithstanding the provisions of this clause, the Administrator may deny,
modify, or revoke the temporary extension under this sub-
paragraph if the Administrator determines that the con-
tinuation of the minor use may cause an unreasonable ad-
verse effect on the environment. In the event of modifica-
tion or revocation, the Administrator shall provide, in writ-
ing, to the registrant a notice revoking the temporary ex-
tension and establish a new effective date by which the
minor use shall be deleted from the registration.

(viii)(I) If data required to support registration of a
pesticide under subparagraph (A) is requested by a Fed-
eral or State regulatory authority, the Administrator shall,
to the extent practicable, coordinate data requirements,
test protocols, timetables, and standards of review and re-
duce burdens and redundancy caused to the registrant by
multiple requirements on the registrant.

(II) The Administrator may enter into a cooperative
agreement with a State to carry out subclause (I).

(III) Not later than 1 year after the date of enactment
of this clause [Aug. 3, 1996], the Administrator shall de-
velop a process to identify and assist in alleviating future
disparities between Federal and State data requirements.

(C) SIMPLIFIED PROCEDURES.—Within nine months
after the date of enactment of this subparagraph [Sep-
tember 30, 1978], the Administrator shall, by regulation,
prescribe simplified procedures for the registration of pes-
ticides, which shall include the provisions of subparagraph
(D) of this paragraph.

(D) EXEMPTION.—No applicant for registration of a
pesticide who proposes to purchase a registered pesticide
from another producer in order to formulate such pur-
chased pesticide into the pesticide that is the subject of the
application shall be required to—

(i) submit or cite data pertaining to such pur-
chased product; or

(ii) offer to pay reasonable compensation otherwise
required by paragraph (1)(D) of this subsection for the
use of any such data.

(E) MINOR USE WAIVER.—In handling the registration
of a pesticide for a minor use, the Administrator may
waive otherwise applicable data requirements if the Ad-
ministrator determines that the absence of such data will
not prevent the Administrator from determining—

(i) the incremental risk presented by the minor
use of the pesticide; and

(ii) that such risk, if any, would not be an unre-
asonable adverse effect on the environment.

(3) TIME FOR ACTING WITH RESPECT TO APPLICATION.—

(A) IN GENERAL.—The Administrator shall review the
data after receipt of the application and shall, as expedi-
tiously as possible, either register the pesticide in accord-
ance with paragraph (5), or notify the applicant of the Ad-
ministrator's determination that it does not comply with
the provisions of the Act in accordance with paragraph (6).
(B) IDENTICAL OR SUBSTANTIALLY SIMILAR.—(i) The Administrator shall, as expeditiously as possible, review and act on any application received by the Administrator that—

(I) proposes the initial or amended registration of an end-use pesticide that, if registered as proposed, would be identical or substantially similar in composition and labeling to a currently-registered pesticide identified in the application, or that would differ in composition and labeling from such currently-registered pesticide only in ways that would not significantly increase the risk of unreasonable adverse effects on the environment; or

(II) proposes an amendment to the registration of a registered pesticide that does not require scientific review of data.

(ii) In expediting the review of an application for an action described in clause (i), the Administrator shall—

(I) review the application in accordance with section 33(f)(4)(B) and, if the application is found to be incomplete, reject the application;

(II) not later than the applicable decision review time established pursuant to section 33(f)(4)(B), or, if no review time is established, not later than 90 days after receiving a complete application, notify the registrant if the application has been granted or denied; and

(III) if the application is denied, notify the registrant in writing of the specific reasons for the denial of the application.

(C) MINOR USE REGISTRATION.—

(i) The Administrator shall, as expeditiously as possible, review and act on any complete application—

(I) that proposes the initial registration of a new pesticide active ingredient if the active ingredient is proposed to be registered solely for minor uses, or proposes a registration amendment solely for minor uses to an existing registration; or

(II) for a registration or a registration amendment that proposes significant minor uses.

(ii) For the purposes of clause (i)—

(I) the term “as expeditiously as possible” means that the Administrator shall, to the greatest extent practicable, complete a review and evaluation of all data, submitted with a complete application, within 12 months after the submission of the complete application, and the failure of the Administrator to complete such a review and evaluation under clause (i) shall not be subject to judicial review; and

(II) the term “significant minor uses” means 3 or more minor uses proposed for every nonminor use, a minor use that would, in the judgment of the Administrator, serve as a replacement for any
use which has been canceled in the 5 years pre-
ceeding the receipt of the application, or a minor
use that in the opinion of the Administrator would
avoid the reissuance of an emergency exemption
under section 18 for that minor use.

(D) ADEQUATE TIME FOR SUBMISSION OF MINOR USE
DATA.—If a registrant makes a request for a minor use
waiver, regarding data required by the Administrator, pur-
suant to paragraph (2)(E), and if the Administrator denies
in whole or in part such data waiver request, the reg-
istrant shall have a full-time period for providing such
data. For purposes of this subparagraph, the term “full-
time period” means the time period originally established
by the Administrator for submission of such data, begin-
ing with the date of receipt by the registrant of the Ad-
ministrator’s notice of denial.

(4) NOTICE OF APPLICATION.—The Administrator shall pub-
lish in the Federal Register, promptly after receipt of the state-
ment and other data required pursuant to paragraphs (1) and
(2), a notice of each application for registration of any pesticide
if it contains any new active ingredient or if it would entail a
changed use pattern. The notice shall provide for a period of
30 days in which any Federal agency or any other interested
person may comment.

(5) APPROVAL OF REGISTRATION.—The Administrator shall
register a pesticide if the Administrator determines that, when
considered with any restrictions imposed under subsection
(d)—

(A) its composition is such as to warrant the proposed
claims for it;
(B) its labeling and other material required to be sub-
mited comply with the requirements of this Act;
(C) it will perform its intended function without un-
reasonable adverse effects on the environment; and
(D) when used in accordance with widespread and
commonly recognized practice it will not generally cause
unreasonable adverse effects on the environment.

The Administrator shall not make any lack of essentiality a
criterion for denying registration of any pesticide. Where two
pesticides meet the requirements of this paragraph, one should
not be registered in preference to the other. In considering an
application for the registration of a pesticide, the Adminis-
trator may waive data requirements pertaining to efficacy, in
which event the Administrator may register the pesticide with-
out determining that the pesticide’s composition is such as to
warrant proposed claims of efficacy. If a pesticide is found to
be efficacious by any State under section 24(c) of this Act, a
presumption is established that the Administrator shall waive
data requirements pertaining to efficacy for use of the pesticide
in such State.

(6) DENIAL OF REGISTRATION.—If the Administrator deter-
mines that the requirements of paragraph (5) for registration
are not satisfied, the Administrator shall notify the applicant
for registration of the Administrator’s determination and of the
Administrator’s reasons (including the factual basis) therefor, and that, unless the applicant corrects the conditions and notifies the Administrator thereof during the 30-day period beginning with the day after the date on which the applicant receives the notice, the Administrator may refuse to register the pesticide. Whenever the Administrator refuses to register a pesticide, the Administrator shall notify the applicant of the Administrator’s decision and of the Administrator’s reasons (including the factual basis) therefor. The Administrator shall promptly publish in the Federal Register notice of such denial of registration and the reasons therefor. Upon such notification, the applicant for registration or other interested person with the concurrence of the applicant shall have the same remedies as provided for in section 6.

(7) Registration under special circumstances.—Notwithstanding the provisions of paragraph (5)—

(A) The Administrator may conditionally register or amend the registration of a pesticide if the Administrator determines that (i) the pesticide and proposed use are identical or substantially similar to any currently registered pesticide and use thereof, or differ only in ways that would not significantly increase the risk of unreasonable adverse effects on the environment, and (ii) approving the registration or amendment in the manner proposed by the applicant would not significantly increase the risk of any unreasonable adverse effect on the environment. An applicant seeking conditional registration or amended registration under this subparagraph shall submit such data as would be required to obtain registration of a similar pesticide under paragraph (5). If the applicant is unable to submit an item of data because it has not yet been generated, the Administrator may register or amend the registration of the pesticide under such conditions as will require the submission of such data not later than the time such data are required to be submitted with respect to similar pesticides already registered under this Act.

(B) The Administrator may conditionally amend the registration of a pesticide to permit additional uses of such pesticide notwithstanding that data concerning the pesticide may be insufficient to support an unconditional amendment, if the Administrator determines that (i) the applicant has submitted satisfactory data pertaining to the proposed additional use, and (ii) amending the registration in the manner proposed by the applicant would not significantly increase the risk of any unreasonable adverse effect on the environment. Notwithstanding the foregoing provisions of this subparagraph, no registration of a pesticide may be amended to permit an additional use of such pesticide if the Administrator has issued a notice stating that such pesticide, or any ingredient thereof, meets or exceeds risk criteria associated in whole or in part with human dietary exposure enumerated in regulations issued under this Act, and during the pendency of any risk-benefit evaluation initiated by such notice, if (I) the additional use of
such pesticide involves a major food or feed crop, or (II) the
additional use of such pesticide involves a minor food or
feed crop and the Administrator determines, with the con-
currence of the Secretary of Agriculture, there is available
an effective alternative pesticide that does not meet or ex-
ceed such risk criteria. An applicant seeking amended reg-
istration under this subparagraph shall submit such data
as would be required to obtain registration of a similar
pesticide under paragraph (5). If the applicant is unable to
submit an item of data (other than data pertaining to the
proposed additional use) because it has not yet been gen-
erated, the Administrator may amend the registration
under such conditions as will require the submission of
such data not later than the time such data are required
to be submitted with respect to similar pesticides already
registered under this Act.

(C) The Administrator may conditionally register a
pesticide containing an active ingredient not contained in
any currently registered pesticide for a period reasonably
sufficient for the generation and submission of required
data (which are lacking because a period reasonably suffi-
cient for generation of the data has not elapsed since the
Administrator first imposed the data requirement) on the
condition that by the end of such period the Administrator
receives such data and the data do not meet or exceed risk
criteria enumerated in regulations issued under this Act,
and on such other conditions as the Administrator may
prescribe. A conditional registration under this subpara-
graph shall be granted only if the Administrator deter-
mines that use of the pesticide during such period will not
cause any unreasonable adverse effect on the environment,
and that use of the pesticide is in the public interest.

(8) INTERIM ADMINISTRATIVE REVIEW.—Notwithstanding
any other provision of this Act, the Administrator may not ini-
tiate a public interim administrative review process to develop
a risk-benefit evaluation of the ingredients of a pesticide or any
of its uses prior to initiating a formal action to cancel, suspend,

or deny registration of such pesticide, required under this Act,
unless such interim administrative process is based on a vali-
dated test or other significant evidence raising prudent con-
cerns of unreasonable adverse risk to man or to the environ-
ment. Notice of the definition of the terms “validated test” and
“other significant evidence” as used herein shall be published
by the Administrator in the Federal Register.

(9) LABELING.—

(A) ADDITIONAL STATEMENTS.—Subject to subpara-
graphs (B) and (C), it shall not be a violation of this Act
for a registrant to modify the labeling of an antimicrobial
pesticide product to include relevant information on prod-
uct efficacy, product composition, container composition or
design, or other characteristics that do not relate to any
pesticidal claim or pesticidal activity.

(B) REQUIREMENTS.—Proposed labeling information
under subparagraph (A) shall not be false or misleading.
shall not conflict with or detract from any statement required by law or the Administrator as a condition of registration, and shall be substantiated on the request of the Administrator.

(C) Notification and Disapproval.—

(i) Notification.—A registration may be modified under subparagraph (A) if—

(I) the registrant notifies the Administrator in writing not later than 60 days prior to distribution or sale of a product bearing the modified labeling; and

(II) the Administrator does not disapprove of the modification under clause (ii).

(ii) Disapproval.—Not later than 30 days after receipt of a notification under clause (i), the Administrator may disapprove the modification by sending the registrant notification in writing stating that the proposed language is not acceptable and stating the reasons why the Administrator finds the proposed modification unacceptable.

(iii) Restriction on Sale.—A registrant may not sell or distribute a product bearing a disapproved modification.

(iv) Objection.—A registrant may file an objection in writing to a disapproval under clause (ii) not later than 30 days after receipt of notification of the disapproval.

(v) Final Action.—A decision by the Administrator following receipt and consideration of an objection filed under clause (iv) shall be considered a final agency action.

(D) Use Dilution.—The label or labeling required under this Act for an antimicrobial pesticide that is or may be diluted for use may have a different statement of caution or protective measures for use of the recommended diluted solution of the pesticide than for use of a concentrate of the pesticide if the Administrator determines that—

(i) adequate data have been submitted to support the statement proposed for the diluted solution uses; and

(ii) the label or labeling provides adequate protection for exposure to the diluted solution of the pesticide.

(10) Expedited Registration of Pesticides.—

(A) Not later than 1 year after the date of enactment of this paragraph [Aug. 3, 1996], the Administrator shall, utilizing public comment, develop procedures and guidelines, and expedite the review of an application for registration of a pesticide or an amendment to a registration that satisfies such guidelines.

(B) Any application for registration or an amendment, including biological and conventional pesticides, will be considered for expedited review under this paragraph. An application for registration or an amendment shall qualify
for expedited review if use of the pesticide proposed by the
application may reasonably be expected to accomplish 1 or
more of the following:

(i) Reduce the risks of pesticides to human health.
(ii) Reduce the risks of pesticides to nontarget or-
ganisms.
(iii) Reduce the potential for contamination of
groundwater, surface water, or other valued environ-
mental resources.
(iv) Broaden the adoption of integrated pest man-
agement strategies, or make such strategies more
available or more effective.

(C) The Administrator, not later than 30 days after re-
ceipt of an application for expedited review, shall notify
the applicant whether the application is complete. If it is
found to be incomplete, the Administrator may either re-
ject the request for expedited review or ask the applicant
for additional information to satisfy the guidelines devel-
oped under subparagraph (A).

(11) INTERAGENCY WORKING GROUP.—

(A) DEFINITION OF COVERED AGENCY.—In this para-
graph, the term “covered agency” means any of the fol-
lowing:

(i) The Department of Agriculture.
(ii) The Department of Commerce.
(iii) The Department of the Interior.
(v) The Environmental Protection Agency.

(B) ESTABLISHMENT.—The Administrator shall estab-
lish an interagency working group, to be comprised of rep-
resentatives from each covered agency, to provide rec-
ommendations regarding, and to implement a strategy for
improving, the consultation process required under section
for pesticide registration and registration review.

(C) DUTIES.—The interagency working group estab-
lished under subparagraph (B) shall—

(i) analyze relevant Federal law (including regula-
tions) and case law for purposes of providing an out-
line of the legal and regulatory framework for the con-
sultation process referred to in that subparagraph, in-
cluding—

(I) requirements under this Act and the En-
dangered Species Act of 1973 (16 U.S.C. 1531 et
seq.);
(II) Federal case law regarding the intersec-
tion of this Act and the Endangered Species Act of
1973 (16 U.S.C. 1531 et seq.); and
(III) Federal regulations relating to the pes-
ticide consultation process;
(ii) provide advice regarding methods of—

(I) defining the scope of actions of the covered
agencies that are subject to the consultation re-
quirement referred to in subparagraph (B); and
(II) properly identifying and classifying effects of actions of the covered agencies with respect to that consultation requirement;

(iii) identify the obligations and limitations under Federal law of each covered agency for purposes of providing a legal and regulatory framework for developing the recommendations referred to in subparagraph (B);

(iv) review practices for the consultation referred to in subparagraph (B) to identify problem areas, areas for improvement, and best practices for conducting that consultation among the covered agencies;

(v) develop scientific and policy approaches to increase the accuracy and timeliness of the process for that consultation, in accordance with requirements of this Act and the Endangered Species Act of 1973 (16 U.S.C. 1531 et seq.), including—

(I) processes to efficiently share data and coordinate analyses among the Department of Agriculture, the Department of Commerce, the Department of the Interior, and the Environmental Protection Agency;

(II) a streamlined process for identifying which actions require no consultation, informal consultation, or formal consultation;

(III) an approach that will provide clarity with respect to what constitutes the best scientific and commercial data available in the fields of pesticide use and ecological risk assessment, pursuant to section 7(a)(2) of the Endangered Species Act of 1973 (16 U.S.C. 1536(a)(2)); and

(IV) approaches that enable the Environmental Protection Agency to better assist the Department of the Interior and the Department of Commerce in carrying out obligations under that section in a timely and efficient manner; and

(vi) propose and implement a strategy to implement approaches to consultations under the Endangered Species Act of 1973 (16 U.S.C. 1531 et seq.) and document that strategy in a memorandum of understanding, revised regulations, or another appropriate format to promote durable cooperation among the covered agencies.

(D) REPORTS.—

(i) PROGRESS REPORTS.—

(I) IN GENERAL.—Not later than 18 months after the date of enactment of this paragraph, the Administrator, in coordination with the head of each other covered agency, shall submit to the Committee on Agriculture of the House of Representatives and the Committee on Agriculture, Nutrition, and Forestry of the Senate a report describing the progress of the working group in de-
(I) Requirements.—The report under this clause shall—
   (aa) reflect the perspectives of each covered agency; and
   (bb) identify areas of new consensus and continuing topics of disagreement and debate.

(ii) Results.—
   (I) In General.—Not later than 1 year after the date of enactment of this paragraph, the Administrator, in coordination with the head of each other covered agency, shall submit to the Committee on Agriculture of the House of Representatives and the Committee on Agriculture, Nutrition, and Forestry of the Senate a report describing—
      (aa) the recommendations developed under subparagraph (B); and
      (bb) plans for implementation of those recommendations.
   (II) Requirements.—The report under this clause shall—
      (aa) reflect the perspectives of each covered agency; and
      (bb) identify areas of consensus and continuing topics of disagreement and debate, if any.

(iii) Implementation.—Not later than 1 year after the date of submission of the report under clause (i), the Administrator, in coordination with the head of each other covered agency, shall submit to the Committee on Agriculture of the House of Representatives and the Committee on Agriculture, Nutrition, and Forestry of the Senate a report describing—
   (I) the implementation of the recommendations referred to in that clause;
   (II) the extent to which that implementation improved the consultation process referred to in subparagraph (B); and
   (III) any additional recommendations for improvements to the process described in subparagraph (B).

(iv) Other Reports.—Not later than the date that is 180 days after the date of submission of the report under clause (iii), and not less frequently than once every 180 days thereafter during the 5-year period beginning on that date, the Administrator, in coordination with the head of each other covered agency, shall submit to the Committee on Agriculture of the House of Representatives and the Committee on Agriculture, Nutrition, and Forestry of the Senate a report describing—
(I) the implementation of the recommendations referred to in that clause;
(II) the extent to which that implementation improved the consultation process referred to in subparagraph (B); and
(III) any additional recommendations for improvements to the process described in subparagraph (B).

(E) CONSULTATION WITH PRIVATE SECTOR.—In carrying out the duties under this paragraph, the working group shall, as appropriate—
   (i) consult with, representatives of interested industry stakeholders and nongovernmental organizations; and
   (ii) take into consideration factors, such as actual and potential differences in interest between, and the views of, those stakeholders and organizations.

(F) FEDERAL ADVISORY COMMITTEE ACT.—The Federal Advisory Committee Act (5 U.S.C. App.) shall not apply to the working group established under this paragraph.

(G) SAVINGS CLAUSE.—Nothing in this paragraph supersedes any provision of—
   (i) this Act; or

(d) CLASSIFICATION OF PESTICIDES.—

   (1) CLASSIFICATION FOR GENERAL USE, RESTRICTED USE, OR BOTH.—

   (A) As a part of the registration of a pesticide the Administrator shall classify it as being for general use or for restricted use. If the Administrator determines that some of the uses for which the pesticide is registered should be for general use and that other uses for which it is registered should be for restricted use, the Administrator shall classify it for both general use and restricted use. Pesticide uses may be classified by regulation on the initial classification and registered pesticides may be classified prior to reregistration. If some of the uses of the pesticide are classified for general use and other uses are classified for restricted use, the directions relating to its general uses shall be clearly separated and distinguished from those directions relating to its restricted uses. The Administrator may require that its packaging and labeling for restricted uses shall be clearly distinguishable from its packaging and labeling for general uses.

   (B) If the Administrator determines that the pesticide, when applied in accordance with its directions for use, warnings and cautions and for the uses for which it is registered, or for one or more of such uses, or in accordance with a widespread and commonly recognized practice, will not generally cause unreasonable adverse effects on the environment, the Administrator will classify the pesticide,
or the particular use or uses of the pesticide to which the determination applies, for general use.

(C) If the Administrator determines that the pesticide, when applied in accordance with its directions for use, warnings and cautions and for the uses for which it is registered, or for one or more of such uses, or in accordance with a widespread and commonly recognized practice, may generally cause, without additional regulatory restrictions, unreasonable adverse effects on the environment, including injury to the applicator, the Administrator shall classify the pesticide, or the particular use or uses to which the determination applies, for restricted use:

(i) If the Administrator classifies a pesticide, or one or more uses of such pesticide, for restricted use because of a determination that the acute dermal or inhalation toxicity of the pesticide presents a hazard to the applicator or other persons, the pesticide shall be applied for any use to which the restricted classification applies only by or under the direct supervision of a certified applicator.

(ii) If the Administrator classifies a pesticide, or one or more uses of such pesticide, for restricted use because of a determination that its use without additional regulatory restriction may cause unreasonable adverse effects on the environment, the pesticide shall be applied for any use to which the determination applies only by or under the direct supervision of a certified applicator, or subject to such other restrictions as the Administrator may provide by regulation. Any such regulation shall be reviewable in the appropriate court of appeals upon petition of a person adversely affected filed within 60 days of the publication of the regulation in final form.

(2) Change in Classification.—If the Administrator determines that a change in the classification of any use of a pesticide from general use to restricted use is necessary to prevent unreasonable adverse effects on the environment, the Administrator shall notify the registrant of such pesticide of such determination at least forty-five days before making the change and shall publish the proposed change in the Federal Register. The registrant, or other interested person with the concurrence of the registrant, may seek relief from such determination under section 6(b).

(3) Change in Classification from Restricted Use to General Use.—The registrant of any pesticide with one or more uses classified for restricted use may petition the Administrator to change any such classification from restricted to general use. Such petition shall set out the basis for the registrant’s position that restricted use classification is unnecessary because classification of the pesticide for general use would not cause unreasonable adverse effects on the environment. The Administrator, within sixty days after receiving such petition, shall notify the registrant whether the petition has been granted or denied. Any denial shall contain an expla-
nation therefor and any such denial shall be subject to judicial review under section 16 of this Act.

(e) Products with Same Formulation and Claims.—Products which have the same formulation, are manufactured by the same person, the labeling of which contains the same claims, and the labels of which bear a designation identifying the product as the same pesticide may be registered as a single pesticide; and additional names and labels shall be added to the registration by supplemental statements.

(f) Miscellaneous.—

(1) Effect of Change of Labeling or Formulation.—If the labeling or formulation for a pesticide is changed, the registration shall be amended to reflect such change if the Administrator determines that the change will not violate any provision of this Act.

(2) Registration Not a Defense.—In no event shall registration of an article be construed as a defense for the commission of any offense under this Act. As long as no cancellation proceedings are in effect registration of a pesticide shall be prima facie evidence that the pesticide, its labeling and packaging comply with the registration provisions of the Act.

(3) Authority to Consult Other Federal Agencies.—In connection with consideration of any registration or application for registration under this section, the Administrator may consult with any other Federal agency.

(4) Mixtures of Nitrogen Stabilizers and Fertilizer Products.—Any mixture or other combination of—

(A) 1 or more nitrogen stabilizers registered under this Act; and

(B) 1 or more fertilizer products,

shall not be subject to the provisions of this section or sections 4, 5, 7, 15, and 17(a)(2) if the mixture or other combination is accompanied by the labeling required under this Act for the nitrogen stabilizer contained in the mixture or other combination, the mixture or combination is mixed or combined in accordance with such labeling, and the mixture or combination does not contain any active ingredient other than the nitrogen stabilizer.

(g) Registration Review.—

(1)(A) General rule.—

(i) In General.—The registrations of pesticides are to be periodically reviewed.

(ii) Regulations.—In accordance with this subparagraph, the Administrator shall by regulation establish a procedure for accomplishing the periodic review of registrations.

(iii) Initial Registration Review.—The Administrator shall complete the registration review of each pesticide or pesticide case, which may be composed of 1 or more active ingredients and the products associated with the active ingredients, not later than the later of—

(I) October 1, 2022; or
(II) the date that is 15 years after the date on which the first pesticide containing a new active ingredient is registered.

(iv) Subsequent registration review.—Not later than 15 years after the date on which the initial registration review is completed under clause (iii) and each 15 years thereafter, the Administrator shall complete a subsequent registration review for each pesticide or pesticide case.

(v) Cancellation.—No registration shall be canceled as a result of the registration review process unless the Administrator follows the procedures and substantive requirements of section 6.

(B) Docketing.—

(i) In general.—Subject to clause (ii), after meeting with 1 or more individuals that are not government employees to discuss matters relating to a registration review, the Administrator shall place in the docket minutes of the meeting, a list of attendees, and any documents exchanged at the meeting, not later than the earlier of—

(I) the date that is 45 days after the meeting; or

(II) the date of issuance of the registration review decision.

(ii) Protected information.—The Administrator shall identify, but not include in the docket, any confidential business information the disclosure of which is prohibited by section 10.

(C) Limitation.—Nothing in this subsection shall prohibit the Administrator from undertaking any other review of a pesticide pursuant to this Act.

(2)(A) Data.—The Administrator shall use the authority in subsection (c)(2)(B) to require the submission of data when such data are necessary for a registration review.

(B) Data submission, compensation, and exemption.—For purposes of this subsection, the provisions of subsections (c)(1), (c)(2)(B), and (c)(2)(D) shall be utilized for and be applicable to any data required for registration review.

(h) Registration Requirements for Antimicrobial Pesticides.—

(1) Evaluation of process.—To the maximum extent practicable consistent with the degrees of risk presented by an antimicrobial pesticide and the type of review appropriate to evaluate the risks, the Administrator shall identify and evaluate reforms to the antimicrobial registration process that would reduce review periods existing as of the date of enactment of this subsection [Aug. 3, 1996] for antimicrobial pesticide product registration applications and applications for amended registration of antimicrobial pesticide products, including—

(A) new antimicrobial active ingredients;

(B) new antimicrobial end-use products;

(C) substantially similar or identical antimicrobial pesticides; and
(D) amendments to antimicrobial pesticide registrations.

(2) REVIEW TIME PERIOD REDUCTION GOAL.—Each reform identified under paragraph (1) shall be designed to achieve the goal of reducing the review period following submission of a complete application, consistent with the degree of risk, to a period of not more than—

(A) 540 days for a new antimicrobial active ingredient pesticide registration;
(B) 270 days for a new antimicrobial use of a registered active ingredient;
(C) 120 days for any other new antimicrobial product;
(D) 90 days for a substantially similar or identical antimicrobial product;
(E) 90 days for an amendment to an antimicrobial registration that does not require scientific review of data; and
(F) 120 days for an amendment to an antimicrobial registration that requires scientific review of data and that is not otherwise described in this paragraph.

(3) IMPLEMENTATION.—

(A) PROPOSED RULEMAKING.—

(i) ISSUANCE.—Not later than 270 days after the date of enactment of this subsection [Aug. 3, 1996], the Administrator shall publish in the Federal Register proposed regulations to accelerate and improve the review of antimicrobial pesticide products designed to implement, to the extent practicable, the goals set forth in paragraph (2).

(ii) REQUIREMENTS.—Proposed regulations issued under clause (i) shall—

(I) define the various classes of antimicrobial use patterns, including household, industrial, and institutional disinfectants and sanitizing pesticides, preservatives, water treatment, and pulp and paper mill additives, and other such products intended to disinfect, sanitize, reduce, or mitigate growth or development of microbiological organisms, or protect inanimate objects, industrial processes or systems, surfaces, water, or other chemical substances from contamination, fouling, or deterioration caused by bacteria, viruses, fungi, protozoa, algae, or slime;

(II) differentiate the types of review undertaken for antimicrobial pesticides;

(III) conform the degree and type of review to the risks and benefits presented by antimicrobial pesticides and the function of review under this Act, considering the use patterns of the product, toxicity, expected exposure, and product type;

(IV) ensure that the registration process is sufficient to maintain antimicrobial pesticide efficacy and that antimicrobial pesticide products continue to meet product performance standards
and effectiveness levels for each type of label claim made; and
(V) implement effective and reliable deadlines for process management.

(iii) COMMENTS.—In developing the proposed regulations, the Administrator shall solicit the views from registrants and other affected parties to maximize the effectiveness of the rule development process.

(B) FINAL REGULATIONS.—
(i) ISSUANCE.—The Administrator shall issue final regulations not later than 240 days after the close of the comment period for the proposed regulations.
(ii) FAILURE TO MEET GOAL.—If a goal described in paragraph (2) is not met by the final regulations, the Administrator shall identify the goal, explain why the goal was not attained, describe the element of the regulations included instead, and identify future steps to attain the goal.
(iii) REQUIREMENTS.—In issuing final regulations, the Administrator shall—
(I) consider the establishment of a certification process for regulatory actions involving risks that can be responsibly managed, consistent with the degree of risk, in the most cost-efficient manner;
(II) consider the establishment of a certification process by approved laboratories as an adjunct to the review process;
(III) use all appropriate and cost-effective review mechanisms, including—
(aa) expanded use of notification and non-notification procedures;
(bb) revised procedures for application review; and
(cc) allocation of appropriate resources to ensure streamlined management of antimicrobial pesticide registrations; and
(IV) clarify criteria for determination of the completeness of an application.

(C) EXPEDITED REVIEW.—This subsection does not affect the requirements or extend the deadlines or review periods contained in subsection (c)(3).

(D) ALTERNATIVE REVIEW PERIODS.—If the final regulations to carry out this paragraph are not effective 630 days after the date of enactment of this subsection [Aug. 3, 1996], until the final regulations become effective, the review period, beginning on the date of receipt by the Agency of a complete application, shall be—
(i) 2 years for a new antimicrobial active ingredient pesticide registration;
(ii) 1 year for a new antimicrobial use of a registered active ingredient;
(iii) 180 days for any other new antimicrobial product;
(iv) 90 days for a substantially similar or identical antimicrobial product;
(v) 90 days for an amendment to an antimicrobial registration that does not require scientific review of data; and
(vi) 120 days for an amendment to an antimicrobial registration that requires scientific review of data and that is not otherwise described in this subparagraph.

(E) Wood preservatives.—An application for the registration, or for an amendment to the registration, of a wood preservative product for which a claim of pesticidal activity listed in section 2(mm) is made (regardless of any other pesticidal claim that is made with respect to the product) shall be reviewed by the Administrator within the same period as that established under this paragraph for an antimicrobial pesticide product application, consistent with the degree of risk posed by the use of the wood preservative product, if the application requires the applicant to satisfy the same data requirements as are required to support an application for a wood preservative product that is an antimicrobial pesticide.

(F) Notification.—
(i) In general.—Subject to clause (iii), the Administrator shall notify an applicant whether an application has been granted or denied not later than the final day of the appropriate review period under this paragraph, unless the applicant and the Administrator agree to a later date.

(ii) Final decision.—If the Administrator fails to notify an applicant within the period of time required under clause (i), the failure shall be considered an agency action unlawfully withheld or unreasonably delayed for purposes of judicial review under chapter 7 of title 5, United States Code.

(iii) Exemption.—This subparagraph does not apply to an application for an antimicrobial pesticide that is filed under subsection (c)(3)(B) prior to 90 days after the date of enactment of this subsection.

[Aug. 3, 1996.]

(iv) Limitation.—Notwithstanding clause (ii), the failure of the Administrator to notify an applicant for an amendment to a registration for an antimicrobial pesticide shall not be judicially reviewable in a Federal or State court if the amendment requires scientific review of data within—

(I) the time period specified in subparagraph (D)(vi), in the absence of a final regulation under subparagraph (B); or

(II) the time period specified in paragraph (2)(F), if adopted in a final regulation under subparagraph (B).

(4) Annual report.—As Amended Through P.L. 116-8, Enacted March 08, 2019.
Sec. 4. | 7 U.S.C. 136a–11 | Reregistration of Registered Pesticides.

(a) General Rule.—The Administrator shall reregister, in accordance with this section, each registered pesticide containing any active ingredient contained in any pesticide first registered before November 1, 1984, except for any pesticide as to which the Administrator has determined, after November 1, 1984, and before the effective date of this section [December 24, 1988], that—

(1) there are no outstanding data requirements; and

(2) the requirements of section 3(c)(5) have been satisfied.

(b) Reregistration Phases.—Reregistrations of pesticides under this section shall be carried out in the following phases:

(1) The first phase shall include the listing under subsection (c) of the active ingredients of the pesticides that will be reregistered.

(2) The second phase shall include the submission to the Administrator under subsection (d) of notices by registrants respecting their intention to seek reregistration, identification by registrants of missing and inadequate data for such pesticides, and commitments by registrants to replace such missing or inadequate data within the applicable time period.

(3) The third phase shall include submission to the Administrator by registrants of the information required under subsection (e).

(4) The fourth phase shall include an independent, initial review by the Administrator under subsection (f) of submissions under phases two and three, identification of outstanding data requirements, and the issuance, as necessary, of requests for additional data.

(5) The fifth phase shall include the review by the Administrator under subsection (g) of data submitted for reregistration and appropriate regulatory action by the Administrator.

(c) Phase One.—

(1) Priority for Reregistration.—For purposes of the reregistration of the pesticides described in subsection (a), the Administrator shall list the active ingredients of pesticides and shall give priority to, among others, active ingredients (other than active ingredients for which registration standards have
been issued before the effective date of this section [December 24, 1988] that—
(A) are in use on or in food or feed and may result in postharvest residues;
(B) may result in residues of potential toxicological concern in potable ground water, edible fish, or shellfish;
(C) have been determined by the Administrator before the effective date of this section [December 24, 1988] to have significant outstanding data requirements; or
(D) are used on crops, including in greenhouses and nurseries, where worker exposure is most likely to occur.

(2) Reregistration Lists.—For purposes of reregistration under this section, the Administrator shall by order—
(A) not later than 70 days after the effective date of this section [December 24, 1988], list pesticide active ingredients for which registration standards have been issued before such effective date;
(B) not later than 4 months after such effective date, list the first 150 pesticide active ingredients, as determined under paragraph (1);
(C) not later than 7 months after such effective date, list the second 150 pesticide active ingredients, as determined under paragraph (1); and
(D) not later than 10 months after such effective date, list the remainder of the pesticide active ingredients, as determined under paragraph (1).

Each list shall be published in the Federal Register.

(3) Judicial Review.—The content of a list issued by the Administrator under paragraph (2) shall not be subject to judicial review.

(4) Notice to Registrants.—On the publication of a list of pesticide active ingredients under paragraph (2), the Administrator shall send by certified mail to the registrants of the pesticides containing such active ingredients a notice of the time by which the registrants are to notify the Administrator under subsection (d) whether the registrants intend to seek or not to seek reregistration of such pesticides.

(d) Phase Two.—
(1) In General.—The registrant of a pesticide that contains an active ingredient listed under subparagraph (B), (C), or (D) of subsection (c)(2) shall submit to the Administrator, within the time period prescribed by paragraph (4), the notice described in paragraph (2) and any information, commitment, or offer described in paragraph (3).

(2) Notice of Intent to Seek or Not to Seek Reregistration.—
(A) The registrant of a pesticide containing an active ingredient listed under subparagraph (B), (C), or (D) of subsection (c)(2) shall notify the Administrator by certified mail whether the registrant intends to seek or does not intend to seek reregistration of the pesticide.

(B) If a registrant submits a notice under subparagraph (A) of an intention not to seek reregistration of a pesticide, the Administrator shall publish a notice in the...
Federal Register stating that such a notice has been submitted.

(3) Missing or Inadequate Data.—Each registrant of a pesticide that contains an active ingredient listed under subparagraph (B), (C), or (D) of subsection (c)(2) and for which the registrant submitted a notice under paragraph (2) of an intention to seek reregistration of such pesticide shall submit to the Administrator—

(A) in accordance with regulations issued by the Administrator under section 3, an identification of—

(i) all data that are required by regulation to support the registration of the pesticide with respect to such active ingredient;

(ii) data that were submitted by the registrant previously in support of the registration of the pesticide that are inadequate to meet such regulations; and

(iii) data identified under clause (i) that have not been submitted to the Administrator; and

(B) either—

(i) a commitment to replace the data identified under subparagraph (A)(ii) and submit the data identified under subparagraph (A)(iii) within the applicable time period prescribed by paragraph (4)(B); or

(ii) an offer to share in the cost to be incurred by a person who has made a commitment under clause (i) to replace or submit the data and an offer to submit to arbitration as described by section 3(c)(2)(B) with regard to such cost sharing.

For purposes of a submission by a registrant under subparagraph (A)(ii), data are inadequate if the data are derived from a study with respect to which the registrant is unable to make the certification prescribed by subsection (e)(1)(G) that the registrant possesses or has access to the raw data used in or generated by such study. For purposes of a submission by a registrant under such subparagraph, data shall be considered to be inadequate if the data are derived from a study submitted before January 1, 1970, unless it is demonstrated to the satisfaction of the Administrator that such data should be considered to support the registration of the pesticide that is to be reregistered.

(4) Time Periods.—

(A) A submission under paragraph (2) or (3) shall be made—

(i) in the case of a pesticide containing an active ingredient listed under subsection (c)(2)(B), not later than 3 months after the date of publication of the listing of such active ingredient;

(ii) in the case of a pesticide containing an active ingredient listed under subsection (c)(2)(C), not later than 3 months after the date of publication of the listing of such active ingredient; and

(iii) in the case of a pesticide containing an active ingredient listed under subsection (c)(2)(D), not later
than 3 months after the date of publication of the listing of such active ingredient.

On application, the Administrator may extend a time period prescribed by this subparagraph if the Administrator determines that factors beyond the control of the registrant prevent the registrant from complying with such period.

(B) A registrant shall submit data in accordance with a commitment entered into under paragraph (3)(B) within a reasonable period of time, as determined by the Administrator, but not more than 48 months after the date the registrant submitted the commitment. The Administrator, on application of a registrant, may extend the period prescribed by the preceding sentence by no more than 2 years if extraordinary circumstances beyond the control of the registrant prevent the registrant from submitting data within such prescribed period. Upon application of a registrant, the Administrator shall, in the case of a minor use, extend the deadline for the production of residue chemistry data under this subparagraph for data required solely to support that minor use until the final deadline for submission of data under this section for the other uses of the pesticide established as of the date of enactment of the Food Quality Protection Act of 1996 [Aug. 3, 1996] if—

(i) the data to support other uses of the pesticide on a food are being provided;

(ii) the registrant, in submitting a request for such an extension provides a schedule, including interim dates to measure progress, to assure that the data production will be completed before the expiration of the extension period;

(iii) the Administrator has determined that such extension will not significantly delay the Administrator’s schedule for issuing a reregistration eligibility determination required under this section; and

(iv) the Administrator has determined that based on existing data, such extension would not significantly increase the risk of any unreasonable adverse effect on the environment.\(^6\) If the Administrator grants an extension under this subparagraph, the Administrator shall monitor the development of the data and shall ensure that the registrant is meeting the schedule for the production of the data. If the Administrator determines that the registrant is not meeting or has not met the schedule for the production of such data, the Administrator may proceed in accordance with clause (iv) of section 3(c)(2)(B) or other provisions of this section, as appropriate, regarding the continued registration of the affected products with the minor use and shall inform the public of such action. Notwithstanding the provisions of this subparagraph, the

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\(^6\) Indentation of the following sentences of this subparagraph is so in original (as added by sec. 201(c)(2) of P.L. 104–170). Probably should be indented the same as flush matter of this subparagraph.
Administrator may take action to modify or revoke the extension under this subparagraph if the Administrator determines that the extension for the minor use may cause an unreasonable adverse effect on the environment. In such circumstance, the Administrator shall provide written notice to the registrant revoking the extension of time for submission of data. Such data shall instead be due in accordance with the date then established by the Administrator for submission of the data.

(5) CANCELLATION AND REMOVAL.—
(A) If the registrant of a pesticide does not submit a notice under paragraph (2) or (3) within the time prescribed by paragraph (4)(A), the Administrator shall issue a notice of intent to cancel the registration of such registrant for such pesticide and shall publish the notice in the Federal Register and allow 60 days for the submission of comments on the notice. On expiration of such 60 days, the Administrator, by order and without a hearing, may cancel the registration or take such other action, including extension of applicable time periods, as may be necessary to enable reregistration of such pesticide by another person.

(B)(i) If—
(I) no registrant of a pesticide containing an active ingredient listed under subsection (c)(2) notifies the Administrator under paragraph (2) that the registrant intends to seek reregistration of any pesticide containing that active ingredient;
(II) no such registrant complies with paragraph (3)(A); or
(III) no such registrant makes a commitment under paragraph (3)(B) to replace or submit all data described in clauses (ii) and (iii) of paragraph (3)(A);
the Administrator shall publish in the Federal Register a notice of intent to remove the active ingredient from the list established under subsection (c)(2) and a notice of intent to cancel the registrations of all pesticides containing such active ingredient and shall provide 60 days for comment on such notice.

(ii) After the 60-day period has expired, the Administrator, by order, may cancel any such registration without hearing, except that the Administrator shall not cancel a registration under this subparagraph if—
(I) during the comment period a person acquires the rights of the registrant in that registration;
(II) during the comment period that person furnishes a notice of intent to reregister the pesticide in accordance with paragraph (2); and
(III) not later than 120 days after the publication of the notice under this subparagraph, that person has complied with paragraph (3) and the fee prescribed by this section has been paid.
(6) SUSPENSIONS AND PENALTIES.—The Administrator shall issue a notice of intent to suspend the registration of a pesticide in accordance with the procedures prescribed by section 3(c)(2)(B)(iv) if the Administrator determines that (A) progress is insufficient to ensure the submission of the data required for such pesticide under a commitment made under paragraph (3)(B) within the time period prescribed by paragraph (4)(B) or (B) the registrant has not submitted such data to the Administrator within such time period. If the registrant does not commit to support a specific minor use of the pesticide, but is supporting and providing data in a timely and adequate fashion to support uses of the pesticide on a food, or if all uses of the pesticide are nonfood uses and the registrant does not commit to support a specific minor use of the pesticide but is supporting and providing data in a timely and adequate fashion to support other nonfood uses of the pesticide, the Administrator, at the written request of the registrant, shall not take any action pursuant to this paragraph in regard to such unsupported minor use until the final deadline established as of the date of enactment of the Food Quality Protection Act of 1996 [Aug. 3, 1996], for the submission of data under this section for the supported uses identified pursuant to this paragraph unless the Administrator determines that the absence of the data is significant enough to cause human health or environmental concerns. On such a determination the Administrator may refuse the request for extension by the registrant. Upon receipt of the request from the registrant, the Administrator shall publish in the Federal Register a notice of the receipt of the request and the effective date upon which the uses not being supported will be voluntarily deleted from the registration pursuant to section 6(f)(1). If the Administrator grants an extension under this paragraph, the Administrator shall monitor the development of the data for the uses being supported and shall ensure that the registrant is meeting the schedule for the production of such data. If the Administrator determines that the registrant is not meeting or has not met the schedule for the production of such data, the Administrator may proceed in accordance with section 3(c)(2)(B)(iv) regarding the continued registration of the affected products with the minor and other uses and shall inform the public of such action in accordance with section 6(f)(2). Notwithstanding this subparagraph, the Administrator may deny, modify, or revoke the temporary extension under this paragraph if the Administrator determines that the continuation of the minor use may cause an unreasonable adverse effect on the environment. In the event of modification or revocation, the Administrator shall provide, in writing, to the registrant a notice revoking the temporary extension and establish a new effective date by which the minor use shall be deleted from the registration.

(e) PHASE THREE.—

(1) INFORMATION ABOUT STUDIES.—Each registrant of a pesticide that contains an active ingredient listed under subparagraph (B), (C), or (D) of subsection (c)(2) who has submitted a notice under subsection (d)(2) of an intent to seek the
reregistration of such pesticide shall submit, in accordance
with the guidelines issued under paragraph (4), to the Admin-
istrator—

(A) a summary of each study concerning the active in-
gredient previously submitted by the registrant in support
of the registration of a pesticide containing such active in-
gredient and considered by the registrant to be adequate
to meet the requirements of section 3 and the regulations
issued under such section;

(B) a summary of each study concerning the active in-
gredient previously submitted by the registrant in support
of the registration of a pesticide containing such active in-
gredient that may not comply with the requirements of
section 3 and the regulations issued under such section but
which the registrant asserts should be deemed to comply
with such requirements and regulations;

(C) a reformat of the data from each study summa-
rized under subparagraph (A) or (B) by the registrant con-
cerning chronic dosing, oncogenicity, reproductive effects,
mutagenicity, neurotoxicity, teratogenicity, or residue
chemistry of the active ingredient that were submitted to
the Administrator before January 1, 1982;

(D) where data described in subparagraph (C) are not
required for the active ingredient by regulations issued
under section 3, a reformat of acute and subchronic dosing
data submitted by the registrant to the Administrator be-
fore January 1, 1982, that the registrant considers to be
adequate to meet the requirements of section 3 and the
regulations issued under such section;

(E) an identification of data that are required to be
submitted to the Administrator under section 6(a)(2) indi-
cating an adverse effect of the pesticide;

(F) an identification of any other information available
that in the view of the registrant supports the registration;

(G) a certification that the registrant or the Adminis-
trator possesses or has access to the raw data used in or
generated by the studies that the registrant summarized
under subparagraph (A) or (B);

(H) either—

(i) a commitment to submit data to fill each out-
standing data requirement identified by the reg-
istrant; or

(ii) an offer to share in the cost of developing such
data to be incurred by a person who has made a com-
mitment under clause (i) to submit such data, and an
offer to submit to arbitration as described by section
3(c)(2)(B) with regard to such cost sharing; and

(I) evidence of compliance with section 3(c)(1)(D)(ii)
and regulations issued thereunder with regard to pre-
viously submitted data as if the registrant were now seek-
ing the original registration of the pesticide.

A registrant who submits a certification under subparagraph
(G) that is false shall be considered to have violated this Act
and shall be subject to the penalties prescribed by section 14.
(2) Time periods.—

(A) The information required by paragraph (1) shall be submitted to the Administrator—

(i) in the case of a pesticide containing an active ingredient listed under subsection (c)(2)(B), not later than 12 months after the date of publication of the listing of such active ingredient;

(ii) in the case of a pesticide containing an active ingredient listed under subsection (c)(2)(C), not later than 12 months after the date of publication of the listing of such active ingredient; and

(iii) in the case of a pesticide containing an active ingredient listed under subsection (c)(2)(D), not later than 12 months after the date of publication of the listing of such active ingredient.

(B) A registrant shall submit data in accordance with a commitment entered into under paragraph (1)(H) within a reasonable period of time, as determined by the Administrator, but not more than 48 months after the date the registrant submitted the commitment under such paragraph. The Administrator, on application of a registrant, may extend the period prescribed by the preceding sentence by no more than 2 years if extraordinary circumstances beyond the control of the registrant prevent the registrant from submitting data within such prescribed period. Upon application of a registrant, the Administrator shall, in the case of a minor use, extend the deadline for the production of residue chemistry data under this subparagraph for data required solely to support that minor use until the final deadline for submission of data under this section for the other uses of the pesticide as of the date of enactment of the Food Quality Protection Act of 1996 [Aug. 3, 1996] if—

(i) the data to support other uses of the pesticide on a food are being provided;

(ii) the registrant, in submitting a request for such an extension provides a schedule, including interim dates to measure progress, to assure that the data production will be completed before the expiration of the extension period;

(iii) the Administrator has determined that such extension will not significantly delay the Administrator’s schedule for issuing a reregistration eligibility determination required under this section; and

(iv) the Administrator has determined that based on existing data, such extension would not significantly increase the risk of any unreasonable adverse effect on the environment.7 If the Administrator grants an extension under this subparagraph, the Administrator shall monitor the development of the data and shall ensure that the registrant is meeting the...
schedule for the production of the data. If the Administrator determines that the registrant is not meeting or has not met the schedule for the production of such data, the Administrator may proceed in accordance with clause (iv) of section 3(c)(2)(B) or other provisions of this section, as appropriate, regarding the continued registration of the affected products with the minor use and shall inform the public of such action. Notwithstanding the provisions of this subparagraph, the Administrator may take action to modify or revoke the extension under this subparagraph if the Administrator determines that the extension for the minor use may cause an unreasonable adverse effect on the environment. In such circumstance, the Administrator shall provide written notice to the registrant revoking the extension of time for submission of data. Such data shall instead be due in accordance with the date then established by the Administrator for submission of the data.

(3) CANCELLATION.—
(A) If the registrant of a pesticide fails to submit the information required by paragraph (1) within the time prescribed by paragraph (2), the Administrator, by order and without hearing, shall cancel the registration of such pesticide. If the registrant does not commit to support a specific minor use of the pesticide, but is supporting and providing data in a timely and adequate fashion to support uses of the pesticide on a food, or if all uses of the pesticide are nonfood uses and the registrant does not commit to support a specific minor use of the pesticide but is supporting and providing data in a timely and adequate fashion to support other nonfood uses of the pesticide, the Administrator, at the written request of the registrant, shall not take any action pursuant to this subparagraph in regard to such unsupported minor use until the final deadline established as of the date of enactment of the Food Quality Protection Act of 1996 [Aug. 3, 1996] for the submission of data under this section for the supported uses identified pursuant to this subparagraph unless the Administrator determines that the absence of the data is significant enough to cause human health or environmental concerns. On the basis of such determination, the Administrator may refuse the request for extension by the registrant. Upon receipt of the request from the registrant, the Administrator shall publish in the Federal Register a notice of the receipt of the request and the effective date upon which the uses not being supported will be voluntarily deleted from the registration pursuant to section 6(f)(1). If the Administrator grants an extension under this subparagraph, the Administrator shall monitor the development of the data for the uses being supported and shall ensure that the registrant is meeting the schedule for the production of such data. If the Administrator determines that the registrant is not meeting or has not met the
schedule for the production of such data, the Administrator may proceed in accordance with section 3(c)(2)(B)(iv) regarding the continued registration of the affected products with the minor and other uses and shall inform the public of such action in accordance with section 6(f)(2). Notwithstanding this subparagraph, the Administrator may deny, modify, or revoke the temporary extension under this subparagraph if the Administrator determines that the continuation of the minor use may cause an unreasonable adverse effect on the environment. In the event of modification or revocation, the Administrator shall provide, in writing, to the registrant a notice revoking the temporary extension and establish a new effective date by which the minor use shall be deleted from the registration.

(B)(i) If the registrant of a pesticide submits the information required by paragraph (1) within the time prescribed by paragraph (2) and such information does not conform to the guidelines for submissions established by the Administrator, the Administrator shall determine whether the registrant made a good faith attempt to conform its submission to such guidelines.

(ii) If the Administrator determines that the registrant made a good faith attempt to conform its submission to such guidelines, the Administrator shall provide the registrant a reasonable period of time to make any necessary changes or corrections.

(iii)(I) If the Administrator determines that the registrant did not make a good faith attempt to conform its submission to such guidelines, the Administrator may issue a notice of intent to cancel the registration. Such a notice shall be sent to the registrant by certified mail.

(II) The registration shall be canceled without a hearing or further notice at the end of 30 days after receipt by the registrant of the notice unless during that time a request for a hearing is made by the registrant.

(III) If a hearing is requested, a hearing shall be conducted under section 6(d), except that the only matter for resolution at the hearing shall be whether the registrant made a good faith attempt to conform its submission to such guidelines. The hearing shall be held and a determination made within 75 days after receipt of a request for hearing.

(4) GUIDELINES.—

(A) Not later than 1 year after the effective date of this section [December 24, 1988], the Administrator, by order, shall issue guidelines to be followed by registrants in—

(i) summarizing studies;
(ii) reformatting studies;
(iii) identifying adverse information; and
(iv) identifying studies that have been submitted previously that may not meet the requirements of section 3 or regulations issued under such section, under paragraph (1).
Guidelines issued under subparagraph (A) shall not be subject to judicial review.

(5) MONITORING.—The Administrator shall monitor the progress of registrants in acquiring and submitting the data required under paragraph (1).

(f) PHASE FOUR.—

(1) INDEPENDENT REVIEW AND IDENTIFICATION OF OUTSTANDING DATA REQUIREMENTS.—

(A) The Administrator shall review the submissions of all registrants of pesticides containing a particular active ingredient under subsections (d)(3) and (e)(1) to determine if such submissions identified all the data that are missing or inadequate for such active ingredient. To assist the review of the Administrator under this subparagraph, the Administrator may require a registrant seeking reregistration to submit complete copies of studies summarized under subsection (e)(1).

(B) The Administrator shall independently identify and publish in the Federal Register the outstanding data requirements for each active ingredient that is listed under subparagraph (B), (C), or (D) of subsection (c)(2) and that is contained in a pesticide to be reregistered under this section. The Administrator, at the same time, shall issue a notice under section 3(c)(2)(B) for the submission of the additional data that are required to meet such requirements.

(2) TIME PERIODS.—

(A) The Administrator shall take the action required by paragraph (1)—

(i) in the case of a pesticide containing an active ingredient listed under subsection (c)(2)(B), not later than 18 months after the date of the listing of such active ingredient;

(ii) in the case of a pesticide containing an active ingredient listed under subsection (c)(2)(C), not later than 24 months after the date of the listing of such active ingredient; and

(iii) in the case of a pesticide containing an active ingredient listed under subsection (c)(2)(D), not later than 33 months after the date of the listing of such active ingredient.

(B) If the Administrator issues a notice to a registrant under paragraph (1)(B) for the submission of additional data, the registrant shall submit such data within a reasonable period of time, as determined by the Administrator, but not to exceed 48 months after the issuance of such notice. The Administrator, on application of a registrant, may extend the period prescribed by the preceding sentence by no more than 2 years if extraordinary circumstances beyond the control of the registrant prevent the registrant from submitting data within such prescribed period. Upon application of a registrant, the Administrator shall, in the case of a minor use, extend the deadline for the production of residue chemistry data under this sub-
paragraph for data required solely to support that minor use until the final deadline for submission of data under this section for the other uses of the pesticide established as of the date of enactment of the Food Quality Protection Act of 1996 [Aug. 3, 1996] if—

(i) the data to support other uses of the pesticide on a food are being provided;
(ii) the registrant, in submitting a request for such an extension provides a schedule, including interim dates to measure progress, to assure that the data production will be completed before the expiration of the extension period;
(iii) the Administrator has determined that such extension will not significantly delay the Administrator’s schedule for issuing a reregistration eligibility determination required under this section; and
(iv) the Administrator has determined that based on existing data, such extension would not significantly increase the risk of any unreasonable adverse effect on the environment.\(^8\) If the Administrator grants an extension under this subparagraph, the Administrator shall monitor the development of the data and shall ensure that the registrant is meeting the schedule for the production of the data. If the Administrator determines that the registrant is not meeting or has not met the schedule for the production of such data, the Administrator may proceed in accordance with clause (iv) of section 3(c)(2)(B) or other provisions of this section, as appropriate, regarding the continued registration of the affected products with the minor use and shall inform the public of such action. Notwithstanding the provisions of this subparagraph, the Administrator may take action to modify or revoke the extension under this subparagraph if the Administrator determines that the extension for the minor use may cause an unreasonable adverse effect on the environment. In such circumstance, the Administrator shall provide written notice to the registrant revoking the extension of time for submission of data. Such data shall instead be due in accordance with the date then established by the Administrator for submission of the data.

(3) SUSPENSIONS AND PENALTIES.—The Administrator shall issue a notice of intent to suspend the registration of a pesticide in accordance with the procedures prescribed by section 3(c)(2)(B)(iv) if the Administrator determines that (A) tests necessary to fill an outstanding data requirement for such pesticide have not been initiated within 1 year after the issuance of a notice under paragraph (1)(B), or (B) progress is insufficient to ensure submission of the data referred to in clause (A) within the time period prescribed by paragraph (2)(B) or the

\(^8\) Indentation of the following sentences of this subparagraph is so in original (as added by sec. 201(c)(2) of P.L. 104–170). Probably should be indented the same as flush matter of this subparagraph.
required data have not been submitted to the Administrator within such time period. If the registrant does not commit to support a specific minor use of the pesticide, but is supporting and providing data in a timely and adequate fashion to support uses of the pesticide on a food, or if all uses of the pesticide are nonfood uses and the registrant does not commit to support a specific minor use of the pesticide but is supporting and providing data in a timely and adequate fashion to support other nonfood uses of the pesticide, the Administrator, at the written request of the registrant, shall not take any action pursuant to this paragraph in regard to such unsupported minor use until the final deadline established as of the date of enactment of the Food Quality Protection Act of 1996 [Aug. 3, 1996] for the submission of data under this section for the supported uses identified pursuant to this paragraph unless the Administrator determines that the absence of the data is significant enough to cause human health or environmental concerns. On such a determination the Administrator may refuse the request for extension by the registrant. Upon receipt of the request from the registrant, the Administrator shall publish in the Federal Register a notice of the receipt of the request and the effective date upon which the uses not being supported will be voluntarily deleted from the registration pursuant to section 6(f)(1). If the Administrator grants an extension under this paragraph, the Administrator shall monitor the development of the data for the uses being supported and shall ensure that the registrant is meeting the schedule for the production of such data. If the Administrator determines that the registrant is not meeting or has not met the schedule for the production of such data, the Administrator may proceed in accordance with section 3(c)(2)(B)(iv) regarding the continued registration of the affected products with the minor and other uses and shall inform the public of such action in accordance with section 6(f)(2). Notwithstanding this subparagraph, the Administrator may deny, modify, or revoke the temporary extension under this paragraph if the Administrator determines that the continuation of the minor use may cause an unreasonable adverse effect on the environment. In the event of modification or revocation, the Administrator shall provide, in writing, to the registrant a notice revoking the temporary extension and establish a new effective date by which the minor use shall be deleted from the registration.

(g) PHASE FIVE.—

(1) DATA REVIEW.—The Administrator shall conduct a thorough examination of all data submitted under this section concerning an active ingredient listed under subsection (c)(2) and of all other available data found by the Administrator to be relevant.

(2) Reregistration and other actions.—

(A) IN GENERAL.—The Administrator shall make a determination as to eligibility for reregistration—

(i) for all active ingredients subject to reregistration under this section for which tolerances or exemptions from tolerances are required under the Federal
Food, Drug, and Cosmetic Act (21 U.S.C. 301 et seq.), not later than the last date for tolerance reassessment established under section 408(q)(1)(C) of that Act (21 U.S.C. 346a(q)(1)(C)); and

(ii) for all other active ingredients subject to reregistration under this section, not later than October 3, 2008.

(B) PRODUCT-SPECIFIC DATA.—

(i) IN GENERAL.—Before reregistering a pesticide, the Administrator shall obtain any needed product-specific data regarding the pesticide by use of section 3(c)(2)(B) and shall review such data within 90 days after its submission.

(ii) TIMING.—

(II) EXTRAORDINARY CIRCUMSTANCES.—In the case of extraordinary circumstances, the Administrator may provide such a longer period, of not more than 2 additional years, for submission of data to the Administrator under this subparagraph.

(C) After conducting the review required by paragraph (1) for each active ingredient of a pesticide and the review required by subparagraph (B) of this paragraph, the Administrator shall determine whether to reregister a pesticide by determining whether such pesticide meets the requirements of section 3(c)(5). If the Administrator determines that a pesticide is eligible to be reregistered, the Administrator shall reregister such pesticide within 6 months after the submission of the data concerning such pesticide under subparagraph (B).

(D) DETERMINATION TO NOT REREGISTER.—

(i) IN GENERAL.—If after conducting a review under paragraph (1) or subparagraph (B) of this paragraph the Administrator determines that a pesticide should not be reregistered, the Administrator shall take appropriate regulatory action.

(ii) TIMING FOR REGULATORY ACTION.—Regulatory action under clause (i) shall be completed as expeditiously as possible.

(E) As soon as the Administrator has sufficient information with respect to the dietary risk of a particular active ingredient, but in any event no later than the time the Administrator makes a determination under subparagraph (C) or (D) with respect to pesticides containing a particular active ingredient, the Administrator shall—
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(i) reassess each associated tolerance and exemption from the requirement for a tolerance issued under section 408 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 346a);

(ii) determine whether such tolerance or exemption meets the requirements of that Act;

(iii) determine whether additional tolerances or exemptions should be issued;

(iv) publish in the Federal Register a notice setting forth the determinations made under this subparagraph; and

(v) commence promptly such proceedings under this Act and section 408 of the Federal Food, Drug, and Cosmetic Act as are warranted by such determinations.

(h) COMPENSATION OF DATA SUBMITTER.—If data that are submitted by a registrant under subsection (d), (e), (f), or (g) are used to support the application of another person under section 3, the registrant who submitted such data shall be entitled to compensation for the use of such data as prescribed by section 3(c)(1)(D). In determining the amount of such compensation, the fees paid by the registrant under this section shall be taken into account.

(i) FEES.—

(1) MAINTENANCE FEE.—

(A) IN GENERAL.—Subject to other provisions of this paragraph, each registrant of a pesticide shall pay an annual fee by January 15 of each year for each registration, except that no fee shall be charged for more than 200 registrations held by any registrant.

(B) In the case of a pesticide that is registered for a minor agricultural use, the Administrator may reduce or waive the payment of the fee imposed under this paragraph if the Administrator determines that the fee would significantly reduce the availability of the pesticide for the use.

(C) TOTAL AMOUNT OF FEES.—The amount of each fee prescribed under subparagraph (A) shall be adjusted by the Administrator to a level that will result in the collection under this paragraph of, to the extent practicable, an average amount of $31,000,000 for each of fiscal years 2019 through 2023.

(D) Maximum amount of fees for registrants.—The maximum annual fee payable under this paragraph by—

(i) a registrant holding not more than 50 pesticide registrations shall be $129,400 for each of fiscal years 2019 through 2023; and

Title II of the Emergency Wartime Supplemental Appropriations Act, 2003, P.L. 108–11, 117 Stat. 603, provided that “Within 30 days of enactment of this Act, the Administrator of the Environmental Protection Agency shall adjust each maximum annual fee payable pursuant to section 136a–1(i)(5)(D) and (E) in a manner such that maintenance fee collections made to reach the level authorized in division K of Public Law 108–7 shall be established in the same proportion as those maintenance fee collections authorized in Public Law 107–73.”
(ii) a registrant holding over 50 registrations shall be $207,000 for each of fiscal years 2019 through 2023.

(E) Maximum amount of fees for small businesses.—

(i) In general.—For a small business, the maximum annual fee payable under this paragraph by—

(I) a registrant holding not more than 50 pesticide registrations shall be $79,100 for each of fiscal years 2019 through 2023; and

(II) a registrant holding over 50 pesticide registrations shall be $136,800 for each of fiscal years 2019 through 2023.

(ii) Definition of small business.—

(I) In general.—In clause (i), the term “small business” means a corporation, partnership, or unincorporated business that—

(aa) has 500 or fewer employees; and

(bb) during the 3-year period prior to the most recent maintenance fee billing cycle, had an average annual global gross revenue from pesticides that did not exceed $60,000,000.

(II) Affiliates.—

(aa) In general.—In the case of a business entity with 1 or more affiliates, the gross revenue limit under subclause (I)(bb) shall apply to the gross revenue for the entity and all of the affiliates of the entity, including parents and subsidiaries, if applicable.

(bb) Affiliated persons.—For the purpose of item (aa), persons are affiliates of each other if, directly or indirectly, either person controls or has the power to control the other person, or a third person controls or has the power to control both persons.

(cc) Indicia of control.—For the purpose of item (aa), indicia of control include interlocking management or ownership, identity of interests among family members, shared facilities and equipment, and common use of employees.

(F) Fee reduction for certain small businesses.—

(i) Definition.—In this subparagraph, the term “qualified small business entity” means a corporation, partnership, or unincorporated business that—

(I) has 500 or fewer employees;

(II) during the 3-year period prior to the most recent maintenance fee billing cycle, had an average annual global gross revenue from all sources that did not exceed $10,000,000; and

(III) holds not more than 5 pesticide registrations under this paragraph.

(ii) Waiver.—Except as provided in clause (iii), the Administrator shall waive 25 percent of the fee under this paragraph applicable to the first registra-
tion of any qualified small business entity under this paragraph.

(iii) LIMITATION.—The Administrator shall not grant a waiver under clause (ii) to a qualified small business entity if the Administrator determines that the entity has been formed or manipulated primarily for the purpose of qualifying for the waiver.

(G) The Administrator shall exempt any public health pesticide from the payment of the fee prescribed under this paragraph if, in consultation with the Secretary of Health and Human Services, the Administrator determines, based on information supplied by the registrant, that the economic return to the registrant from sales of the pesticide does not support the registration or reregistration of the pesticide.

(H) If any fee prescribed by this paragraph with respect to the registration of a pesticide is not paid by a registrant by the time prescribed, the Administrator, by order and without hearing, may cancel the registration.

(I) The authority provided under this paragraph shall terminate on September 30, 2023.

(2) OTHER FEES.—Except as provided in section 33, during the period beginning on the effective date of the Pesticide Registration Improvement Extension Act of 2018 and ending on September 30, 2025, the Administrator may not levy any other fees for the registration of a pesticide under this Act or any other action covered under a table specified in section 33(b)(3), except as provided in paragraph (1).

(j) EXEMPTION OF CERTAIN REGISTRANTS.—The requirements of subsections (d), (e), (f), and (i) (other than subsection (i)(1)) regarding data concerning an active ingredient and fees for review of such data shall not apply to any person who is the registrant of a pesticide to the extent that, under section 3(c)(2)(D), the person would not be required to submit or cite such data to obtain an initial registration of such pesticide.

(k) REREGISTRATION AND EXPEDITED PROCESSING FUND.—

(1) ESTABLISHMENT.—There shall be established in the Treasury of the United States a reregistration and expedited processing fund which shall be known as the Reregistration and Expedited Processing Fund.10

(2) SOURCE AND USE.—

(A) All moneys derived from fees collected by the Administrator under subsection (i) shall be deposited in the Reregistration and Expedited Processing Fund and shall be available to the Administrator, without fiscal year limitation, specifically to offset the costs of reregistration and expedited processing of the applications specified in paragraph (3), to offset the costs of registration review under...
section 3(g), including the costs associated with any review under the Endangered Species Act of 1973 (16 U.S.C. 1531 et seq.) required as part of the registration review, to offset the costs associated with tracking and implementing registration review decisions, including registration review decisions designed to reduce risk, for the purposes specified in paragraphs (4) and (5), and to enhance the information systems capabilities to improve the tracking of pesticide registration decisions. The Administrator shall, prior to expending any such moneys derived from fees—

(i) effective October 1, 1997, adopt specific and cost accounting rules and procedures as approved by the General Accounting Office and the Inspector General of the Environmental Protection Agency to ensure that moneys derived from fees are allocated solely for the purposes specified in the first sentence of this subparagraph;

(ii) prohibit the use of such moneys derived from fees to pay for any costs other than those necessary to achieve the purposes specified in the first sentence of this subparagraph; and

(iii) ensure that personnel and facility costs associated with the functions to be carried out under this paragraph do not exceed agency averages for comparable personnel and facility costs.

(B) The Administrator shall also—

(i) complete the review of unreviewed reregistration studies required to support the reregistration eligibility decisions scheduled for completion in accordance with subsection (l)(2); and

(ii) contract for such outside assistance as may be necessary for review of required studies, using a generally accepted competitive process for the selection of vendors of such assistance.

(3) REVIEW OF INERT INGREDIENTS; EXPEDITED PROCESSING OF SIMILAR APPLICATIONS.—

(A) For each of fiscal years 2018 through 2023, the Administrator shall use between 1/9 and 1/8 of the maintenance fees collected in such fiscal year to obtain sufficient personnel and resources—

(i) to review and evaluate inert ingredients; and

(ii) to ensure the expedited processing and review of any application that—

(I) proposes the initial or amended registration of an end-use pesticide that, if registered as proposed, would be identical or substantially similar in composition and labeling to a currently-registered pesticide identified in the application, or that would differ in composition and labeling from any such currently-registered pesticide only in ways that would not significantly increase the risk of unreasonable adverse effects on the environment;
(II) proposes an amendment to the registration of a registered pesticide that does not require scientific review of data; or

(III) proposes the initial or amended registration of an end use pesticide that, if registered as proposed, would be used for a public health pesticide.

(B) Any amounts made available under subparagraph (A) shall be used to obtain sufficient personnel and resources to carry out the activities described in such subparagraph that are in addition to the personnel and resources available to carry out such activities on the date of enactment of this section.

[October 25, 1988.]

(C) So long as the Administrator has not met the time frames specified in clause (ii) of section 3(c)(3)(B) with respect to any application subject to section 3(c)(3)(B) that was received prior to the date of enactment of the Food Quality Protection Act of 1996 [Aug. 3, 1996], the Administrator shall use the full amount of the fees specified in subparagraph (A) for the purposes specified therein. Once all applications subject to section 3(c)(3)(B) that were received prior to such date of enactment have been acted upon, no limitation shall be imposed by the preceding sentence of this subparagraph so long as the Administrator meets the time frames specified in clause (ii) of section 3(c)(3)(B) on 90 percent of affected applications in a fiscal year. Should the Administrator not meet such time frames in a fiscal year, the limitations imposed by the first sentence of this subparagraph shall apply until all overdue applications subject to section 3(c)(3)(B) have been acted upon.

(4) EXPEDITED RULEMAKING AND GUIDANCE DEVELOPMENT FOR CERTAIN PRODUCT PERFORMANCE DATA REQUIREMENTS.—

(A) SET-ASIDE.—For each of fiscal years 2018 through 2023, the Administrator shall use not more than $500,000 of the amounts made available to the Administrator in the Reregistration and Expedited Processing Fund for the activities described in subparagraph (B).

(B) PRODUCTS CLAIMING EFFICACY AGAINST INVERTEBRATE PESTS OF SIGNIFICANT PUBLIC HEALTH OR ECONOMIC IMPORTANCE.—The Administrator shall use amounts made available under subparagraph (A) to develop, receive comments with respect to, finalize, and implement the necessary rulemaking and guidance for product performance data requirements to evaluate products claiming efficacy against the following invertebrate pests of significant public health or economic importance (in order of importance):

(i) Bed bugs.

Footnote:
12 Sec. 501(d)(2) of P.L. 104–170 added subpara. (C) to sec. 4(k)(3) (7 U.S.C. 136a–1(k)(3)), without specifying the Act that was being amended. The amendment was executed to this Act to effectuate the probable intent of Congress.
(ii) Premise (including crawling insects, flying insects, and baits).

(iii) Pests of pets (including pet pests controlled by spot-ons, collars, shampoos, powders, or dips).

(iv) Fire ants.

(C) DEADLINES FOR GUIDANCE.—The Administrator shall develop, and publish guidance required by subparagraph (B), with respect to claims of efficacy against pests described in such subparagraph as follows:

(i) With respect to bed bugs, issue final guidance not later than 30 days after the effective date of the Pesticide Registration Improvement Extension Act of 2018.

(ii) With respect to pests specified in clause (ii) of such subparagraph—

(I) submit draft guidance to the Scientific Advisory Panel and for public comment not later than June 30, 2018; and

(II) complete any response to comments received with respect to such draft guidance and finalize the guidance not later than September 30, 2019.

(iii) With respect to pests specified in clauses (iii) and (iv) of such subparagraph—

(I) submit draft guidance to the Scientific Advisory Panel and for public comment not later than June 30, 2019; and

(II) complete any response to comments received with respect to such draft guidance and finalize the guidance not later than March 31, 2021.

(D) REVISION.—The Administrator shall revise the guidance required by subparagraph (B) from time to time, but shall permit applicants and registrants sufficient time to obtain data that meet the requirements specified in such revised guidance.

(E) DEADLINE FOR PRODUCT PERFORMANCE DATA REQUIREMENTS.—The Administrator shall, not later than September 30, 2021, issue regulations prescribing product performance data requirements for any pesticide intended for preventing, destroying, repelling, or mitigating any invertebrate pest of significant public health or economic importance specified in clauses (i) through (iv) of subparagraph (B).

(5) GOOD LABORATORY PRACTICES INSPECTIONS.—

(A) SET-ASIDE.—For each of fiscal years 2018 through 2023, the Administrator shall use not more than $500,000 of the amounts made available to the Administrator in the Reregistration and Expedited Processing Fund for the activities described in subparagraph (B).

(B) ACTIVITIES.—The Administrator shall use amounts made available under subparagraph (A) for enhancements to the good laboratory practices standards compliance monitoring program established under part 160 of title 40.
of the Code of Federal Regulations (or successor regulations), with respect to laboratory inspections and data audits conducted in support of pesticide product registrations under this Act. As part of such monitoring program, the Administrator shall make available to each laboratory inspected under such program in support of such registrations a preliminary summary of inspection observations not later than 60 days after the date on which such an inspection is completed.

(6) UNUSED FUNDS.—Money in the fund not currently needed to carry out this section shall be—

(A) maintained on hand or on deposit;

(B) invested in obligations of the United States or guaranteed thereby; or

(C) invested in obligations, participations, or other instruments that are lawful investments for fiduciary, trust, or public funds.

(7) ACCOUNTING AND PERFORMANCE.—The Administrator shall take all steps necessary to ensure that expenditures from fees authorized by subsection (i)(1)(C)(ii) are used only for the purposes described in paragraphs (2), (3), (4), and (5) and to carry out the goals established under subsection (l). The Reregistration and Expedited Processing Fund shall be designated as an Environmental Protection Agency component for purposes of section 3515(c) of title 31, United States Code. The annual audit required under section 3521 of such title of the financial statements of activities under this Act under section 3515(b) of such title shall include an audit of the fees collected under subsection (i)(1)(C) and disbursed, of the amount appropriated to match such fees, and of the Administrator’s attainment of performance measures and goals established under subsection (l). Such an audit shall also include a review of the reasonableness of the overhead allocation and adequacy of disclosures of direct and indirect costs associated with carrying out the reregistration and expedited processing of the applications specified in paragraph (3), and the basis for and accuracy of all costs paid with moneys derived from such fees. The Inspector General shall conduct the annual audit and report the findings and recommendations of such audit to the Administrator and to the Committees on Agriculture of the House of Representatives and the Senate. The cost of such audit shall be paid for out of the fees collected under subsection (i)(1)(C).

(l) PERFORMANCE MEASURES AND GOAL.—The Administrator shall establish and publish annually in the Federal Register performance measures and goals. Such measures and goals shall include—

(1) the number of products reregistered, canceled, or amended, the status of reregistration, the number and type of
data requests under section 3(c)(2)(B) issued to support product reregistration by active ingredient, the progress in reducing the number of unreviewed, required reregistration studies, the aggregate status of tolerances reassessed, and the number of applications for registration submitted under subsection (k)(3) that were approved or disapproved;

(2) the future schedule for reregistrations, including the projection for such schedules that will be issued under subsection (g)(2)(A) and (B) in the current fiscal year and the succeeding fiscal year; and

(3) the projected year of completion of the reregistrations under this section.

(m) JUDICIAL REVIEW.—Any failure of the Administrator to take any action required by this section shall be subject to judicial review under the procedures prescribed by section 16(b).

(n) AUTHORIZATION OF FUNDS TO DEVELOP PUBLIC HEALTH DATA.—

(1) DEFINITION.—For the purposes of this section, “Secretary” means the Secretary of Health and Human Services, acting through the Public Health Service.

(2) CONSULTATION.—In the case of a pesticide registered for use in public health programs for vector control or for other uses the Administrator determines to be human health protection uses, the Administrator shall, upon timely request by the registrant or any other interested person, or on the Administrator's own initiative may, consult with the Secretary prior to taking final action to suspend registration under section 3(c)(2)(B)(iv), or cancel a registration under section 4, 6(e), or 6(f). In consultation with the Secretary, the Administrator shall prescribe the form and content of requests under this section.

(3) BENEFITS TO SUPPORT FAMILY.—The Administrator, after consulting with the Secretary, shall make a determination whether the potential benefits of continued use of the pesticide for public health or health protection purposes are of such significance as to warrant a commitment by the Secretary to conduct or to arrange for the conduct of the studies required by the Administrator to support continued registration under section 3 or reregistration under section 4.

(4) ADDITIONAL TIME.—If the Administrator determines that such a commitment is warranted and in the public interest, the Administrator shall notify the Secretary and shall, to the extent necessary, amend a notice issued under section 3(c)(2)(B) to specify additional reasonable time periods for submission of the data.

(5) ARRANGEMENTS.—The Secretary shall make such arrangements for the conduct of required studies as the Secretary finds necessary and appropriate to permit submission of data in accordance with the time periods prescribed by the Administrator. Such arrangements may include Public Health Service intramural research activities, grants, contracts, or cooperative agreements with academic, public health, or other organizations qualified by experience and training to conduct such studies.
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(6) SUPPORT.—The Secretary may provide for support of the required studies using funds authorized to be appropriated under this section, the Public Health Service Act, or other appropriate authorities. After a determination is made under subsection (d), the Secretary shall notify the Committees on Appropriations of the House of Representatives and the Senate of the sums required to conduct the necessary studies.

(7) AUTHORIZATION OF APPROPRIATIONS.—There is authorized to be appropriated to carry out the purposes of this section $12,000,000 for fiscal year 1997, and such sums as may be necessary for succeeding fiscal years.

SEC. 5. [7 U.S.C. 136c] EXPERIMENTAL USE PERMITS.

(a) ISSUANCE.—Any person may apply to the Administrator for an experimental use permit for a pesticide. An application for an experimental use permit for a covered application under section 33(b) shall conform with the requirements of that section. The Administrator shall review the application. After completion of the review, but not later than one hundred and twenty days after receipt of the application and all required supporting data (or in the case of an application for an experimental use permit for a covered application under section 33(b), not later than the last day of the applicable timeframe for such application specified in such section), the Administrator shall either issue the permit or notify the applicant of the Administrator's determination not to issue the permit and the reasons therefor. The applicant may correct the application or request a waiver of the conditions for such permit within thirty days of receipt by the applicant of such notification. The Administrator may issue an experimental use permit only if the Administrator determines that the applicant needs such permit in order to accumulate information necessary to register a pesticide under section 3 of this Act. An application for an experimental use permit may be filed at any time.

(b) TEMPORARY TOLERANCE LEVEL.—If the Administrator determines that the use of a pesticide may reasonably be expected to result in any residue on or in food or feed, the Administrator may establish a temporary tolerance level for the residue of the pesticide before issuing the experimental use permit.

(c) USE UNDER PERMIT.—Use of a pesticide under an experimental use permit shall be under the supervision of the Administrator, and shall be subject to such terms and conditions and be for such period of time as the Administrator may prescribe in the permit.

(d) STUDIES.—When any experimental use permit is issued for a pesticide containing any chemical or combination of chemicals which has not been included in any previously registered pesticide, the Administrator may specify that studies be conducted to detect whether the use of the pesticide under the permit may cause unreasonable adverse effects on the environment. All results of such studies shall be reported to the Administrator before such pesticide may be registered under section 3.

(e) REVOCATION.—The Administrator may revoke any experimental use permit, at any time, if the Administrator finds that its terms or conditions are being violated, or that its terms and condi-
Sec. 6. [7 U.S.C. 136d] ADMINISTRATIVE REVIEW; SUSPENSION.

(a) EXISTING STOCKS AND INFORMATION.—15

(1) EXISTING STOCKS.—The Administrator may permit the continued sale and use of existing stocks of a pesticide whose registration is suspended or canceled under this section, or section 3 or 4, to such extent, under such conditions, and for such uses as the Administrator determines that such sale or use is not inconsistent with the purposes of this Act.

(2) INFORMATION.—If at any time after the registration of a pesticide the registrant has additional factual information regarding unreasonable adverse effects on the environment of the pesticide, the registrant shall submit such information to the Administrator.

(b) CANCELLATION AND CHANGE IN CLASSIFICATION.—If it appears to the Administrator that a pesticide or its labeling or other material required to be submitted does not comply with the provisions of this Act or, when used in accordance with widespread and commonly recognized practice, generally causes unreasonable adverse effects on the environment, the Administrator may issue a notice of the Administrator’s intent either—

(1) to cancel its registration or to change its classification together with the reasons (including the factual basis) for the Administrator’s action, or

(2) to hold a hearing to determine whether or not its registration should be canceled or its classification changed.

Such notice shall be sent to the registrant and made public. In determining whether to issue any such notice, the Administrator shall include among those factors to be taken into account the impact of the action proposed in such notice on production and prices of agricultural commodities, retail food prices, and otherwise on the

15 Sec. 106(a)(1) of P.L. 104–170 amended subsec. (a) by striking the heading and inserting “(a) EXISTING STOCKS AND INFORMATION.—”. The second subsec. designation for “(a)” was omitted to effectuate the probable intent of Congress.
agricultural economy. At least 60 days prior to sending such notice to the registrant or making public such notice, whichever occurs first, the Administrator shall provide the Secretary of Agriculture with a copy of such notice and an analysis of such impact on the agricultural economy. If the Secretary comments in writing to the Administrator regarding the notice and analysis within 30 days after receiving them, the Administrator shall publish in the Federal Register (with the notice) the comments of the Secretary and the response of the Administrator with regard to the Secretary’s comments. If the Secretary does not comment in writing to the Administrator regarding the notice and analysis within 30 days after receiving them, the Administrator may notify the registrant and make public the notice at any time after such 30-day period notwithstanding the foregoing 60-day time requirement. The time requirements imposed by the preceding 3 sentences may be waived or modified to the extent agreed upon by the Administrator and the Secretary. Notwithstanding any other provision of this subsection (b) and section 25(d), in the event that the Administrator determines that suspension of a pesticide registration is necessary to prevent an imminent hazard to human health, then upon such a finding the Administrator may waive the requirement of notice to and consultation with the Secretary of Agriculture pursuant to subsection (b) and of submission to the Scientific Advisory Panel pursuant to section 25(d) and proceed in accordance with subsection (c). When a public health use is affected, the Secretary of Health and Human Services should provide available benefits and use information, or an analysis thereof, in accordance with the procedures followed and subject to the same conditions as the Secretary of Agriculture in the case of agricultural pesticides. The proposed action shall become final and effective at the end of 30 days from receipt by the registrant, or publication, of a notice issued under paragraph (1), whichever occurs later, unless within that time either (i) the registrant makes the necessary corrections, if possible, or (ii) a request for a hearing is made by a person adversely affected by the notice. In the event a hearing is held pursuant to such a request or to the Administrator’s determination under paragraph (2), a decision pertaining to registration or classification issued after completion of such hearing shall be final. In taking any final action under this subsection, the Administrator shall consider restricting a pesticide’s use or uses as an alternative to cancellation and shall fully explain the reasons for these restrictions, and shall include among those factors to be taken into account the impact of such final action on production and prices of agricultural commodities, retail food prices, and otherwise on the agricultural economy, and the Administrator shall publish in the Federal Register an analysis of such impact.

(c) Suspension.—

(1) Order.—If the Administrator determines that action is necessary to prevent an imminent hazard during the time required for cancellation or change in classification proceedings, the Administrator may, by order, suspend the registration of the pesticide immediately. Except as provided in paragraph (3), no order of suspension may be issued under this subsection unless the Administrator has issued, or at the same time issues,
a notice of intention to cancel the registration or change the classification of the pesticide under subsection (b). Except as provided in paragraph (3), the Administrator shall notify the registrant prior to issuing any suspension order. Such notice shall include findings pertaining to the question of “imminent hazard”. The registrant shall then have an opportunity, in accordance with the provisions of paragraph (2), for an expedited hearing before the Administrator on the question of whether an imminent hazard exists.

(2) EXPEDITED HEARING.—If no request for a hearing is submitted to the Administrator within five days of the registrant's receipt of the notification provided for by paragraph (1), the suspension order may be issued and shall take effect and shall not be reviewable by a court. If a hearing is requested, it shall commence within five days of the receipt of the request for such hearing unless the registrant and the Administrator agree that it shall commence at a later time. The hearing shall be held in accordance with the provisions of subchapter II of title 5 of the United States Code, except that the presiding officer need not be a certified hearing examiner. The presiding officer shall have ten days from the conclusion of the presentation of evidence to submit recommended findings and conclusions to the Administrator, who shall then have seven days to render a final order on the issue of suspension.

(3) EMERGENCY ORDER.—Whenever the Administrator determines that an emergency exists that does not permit the Administrator to hold a hearing before suspending, the Administrator may issue a suspension order in advance of notification to the registrant. The Administrator may issue an emergency order under this paragraph before issuing a notice of intention to cancel the registration or change the classification of the pesticide under subsection (b) and the Administrator shall proceed to issue the notice under subsection (b) within 90 days of issuing an emergency order. If the Administrator does not issue a notice under subsection (b) within 90 days of issuing an emergency order, the emergency order shall expire. In the case of an emergency order, paragraph (2) shall apply except that (A) the order of suspension shall be in effect pending the expeditious completion of the remedies provided by that paragraph and the issuance of a final order on suspension, and (B) no party other than the registrant and the Administrator shall participate except that any person adversely affected may file briefs within the time allotted by the Administrator’s rules. Any person so filing briefs shall be considered a party to such proceeding for the purposes of section 16(b).

(4) JUDICIAL REVIEW.—A final order on the question of suspension following a hearing shall be reviewable in accordance with Section 16 of this Act, notwithstanding the fact that any related cancellation proceedings have not been completed. Any order of suspension entered prior to a hearing before the Administrator shall be subject to immediate review in an action by the registrant or other interested person with the concurrence of the registrant in an appropriate district court, solely to determine whether the order of suspension was arbitrary.
capricious or an abuse of discretion, or whether the order was issued in accordance with the procedures established by law. The effect of any order of the court will be only to stay the effectiveness of the suspension order, pending the Administrator's final decision with respect to cancellation or change in classification. This action may be maintained simultaneously with any administrative review proceeding under this section. The commencement of proceedings under this paragraph shall not operate as a stay of order, unless ordered by the court.

(d) Public Hearings and Scientific Review.—In the event a hearing is requested pursuant to subsection (b) or determined upon by the Administrator pursuant to subsection (b), such hearing shall be held after due notice for the purpose of receiving evidence relevant and material to the issues raised by the objections filed by the applicant or other interested parties, or to the issues stated by the Administrator, if the hearing is called by the Administrator rather than by the filing of objections. Upon a showing of relevance and reasonable scope of evidence sought by any party to a public hearing, the Hearing Examiner shall issue a subpoena to compel testimony or production of documents from any person. The Hearing Examiner shall be guided by the principles of the Federal Rules of Civil Procedure in making any order for the protection of the witness or the content of documents produced and shall order the payment of reasonable fees and expenses as a condition to requiring testimony of the witness. On contest, the subpoena may be enforced by an appropriate United States district court in accordance with the principles stated herein. Upon the request of any party to a public hearing and when in the Hearing Examiner's judgment it is necessary or desirable, the Hearing Examiner shall at any time before the hearing record is closed refer to a Committee of the National Academy of Sciences the relevant questions of scientific fact involved in the public hearing. No member of any committee of the National Academy of Sciences established to carry out the functions of this section shall have a financial or other conflict of interest with respect to any matter considered by such committee. The Committee of the National Academy of Sciences shall report in writing to the Hearing Examiner within 60 days after such referral on these questions of scientific fact. The report shall be made public and shall be considered as part of the hearing record. The Administrator shall enter into appropriate arrangements with the National Academy of Sciences to assure an objective and competent scientific review of the questions presented to Committees of the Academy and to provide such other scientific advisory services as may be required by the Administrator for carrying out the purposes of this Act. As soon as practicable after completion of the hearing (including the report of the Academy) but not later than 90 days thereafter, the Administrator shall evaluate the data and reports before the Administrator and issue an order either revoking the Administrator's notice of intention issued pursuant to this section, or shall issue an order either canceling the registration, changing the classification, denying the registration, or requiring modification of the labeling or packaging of the article. Such order shall be based only on substantial evidence of record of such hearing and
shall set forth detailed findings of fact upon which the order is based.

(e) Conditional Registration.—

(1) The Administrator shall issue a notice of intent to cancel a registration issued under section 3(c)(7) of this Act if (A) the Administrator, at any time during the period provided for satisfaction of any condition imposed, determines that the registrant has failed to initiate and pursue appropriate action toward fulfilling any condition imposed, or (B) at the end of the period provided for satisfaction of any condition imposed, that condition has not been met. The Administrator may permit the continued sale and use of existing stocks of a pesticide whose conditional registration has been canceled under this subsection to such extent, under such conditions, and for such uses as the Administrator may specify if the Administrator determines that such sale or use is not inconsistent with the purposes of this Act and will not have unreasonable adverse effects on the environment.  

(2) A cancellation proposed under this subsection shall become final and effective at the end of thirty days from receipt by the registrant of the notice of intent to cancel unless during that time a request for hearing is made by a person adversely affected by the notice. If a hearing is requested, a hearing shall be conducted under subsection (d) of this section. The only matters for resolution at that hearing shall be whether the registrant has initiated and pursued appropriate action to comply with the condition or conditions within the time provided or whether the condition or conditions have been satisfied within the time provided, and whether the Administrator's determination with respect to the disposition of existing stocks is consistent with this Act. A decision after completion of such hearing shall be final. Notwithstanding any other provision of this section, a hearing shall be held and a determination made within seventy-five days after receipt of a request for such hearing.

(f) General Provisions.—

(1) Voluntary Cancellation.—

(A) A registrant may, at any time, request that a pesticide registration of the registrant be canceled or amended to terminate one or more pesticide uses.

(B) Before acting on a request under subparagraph (A), the Administrator shall publish in the Federal Register a notice of the receipt of the request and provide for a 30-day period in which the public may comment.

(C) In the case of a pesticide that is registered for a minor agricultural use, if the Administrator determines that the cancellation or termination of uses would adversely affect the availability of the pesticide for use, the Administrator—

(i) shall publish in the Federal Register a notice of the receipt of the request and make reasonable efforts to inform persons who so use the pesticide of the request; and
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(ii) may not approve or reject the request until the termination of the 180-day period beginning on the date of publication of the notice in the Federal Register, except that the Administrator may waive the 180-day period upon the request of the registrant or if the Administrator determines that the continued use of the pesticide would pose an unreasonable adverse effect on the environment.

(D) Subject to paragraph (3)(B), after complying with this paragraph, the Administrator may approve or deny the request.

(2) Publication of Notice.—A notice of denial of registration, intent to cancel, suspension, or intent to suspend issued under this Act or a notice issued under subsection (c)(4) or (d)(5)(A) of section 4 shall be published in the Federal Register and shall be sent by certified mail, return receipt requested, to the registrant’s or applicant’s address of record on file with the Administrator. If the mailed notice is returned to the Administrator as undeliverable at that address, if delivery is refused, or if the Administrator otherwise is unable to accomplish delivery of the notice to the registrant or applicant after making reasonable efforts to do so, the notice shall be deemed to have been received by the registrant or applicant on the date the notice was published in the Federal Register.

(3) Transfer of Registration of Pesticides Registered for Minor Agricultural Uses.—In the case of a pesticide that is registered for a minor agricultural use:

(A) During the 180-day period referred to in paragraph (1)(C)(ii), the registrant of the pesticide may notify the Administrator of an agreement between the registrant and a person or persons (including persons who so use the pesticide) to transfer the registration of the pesticide, in lieu of canceling or amending the registration to terminate the use.

(B) An application for transfer of registration, in conformance with any regulations the Administrator may adopt with respect to the transfer of the pesticide registrations, must be submitted to the Administrator within 30 days of the date of notification provided pursuant to subparagraph (A). If such an application is submitted, the Administrator shall approve the transfer and shall not approve the request for voluntary cancellation or amendment to terminate use unless the Administrator determines that the continued use of the pesticide would cause an unreasonable adverse effect on the environment.

(C) If the Administrator approves the transfer and the registrant transfers the registration of the pesticide, the Administrator shall not cancel or amend the registration to delete the use or rescind the transfer of the registration, during the 180-day period beginning on the date of the approval of the transfer unless the Administrator determines that the continued use of the pesticide would cause an unreasonable adverse effect on the environment.
(D) The new registrant of the pesticide shall assume the outstanding data and other requirements for the pesticide that are pending at the time of the transfer.

(4) Utilization of data for voluntarily canceled pesticide.—When an application is filed with the Administrator for the registration of a pesticide for a minor use and another registrant subsequently voluntarily cancels its registration for an identical or substantially similar pesticide for an identical or substantially similar use, the Administrator shall process, review, and evaluate the pending application as if the voluntary cancellation had not yet taken place except that the Administrator shall not take such action if the Administrator determines that such minor use may cause an unreasonable adverse effect on the environment. In order to rely on this subsection, the applicant must certify that it agrees to satisfy any outstanding data requirements necessary to support the reregistration of the pesticide in accordance with the data submission schedule established by the Administrator.

(g) Notice for stored pesticides with canceled or suspended registrations.—

(1) In general.—Any producer or exporter of pesticides, registrant of a pesticide, applicant for registration of a pesticide, applicant for or holder of an experimental use permit, commercial applicator, or any person who distributes or sells any pesticide, who possesses any pesticide which has had its registration canceled or suspended under this section shall notify the Administrator and appropriate State and local officials of—

(A) such possession,
(B) the quantity of such pesticide such person possesses, and
(C) the place at which such pesticide is stored.

(2) Copies.—The Administrator shall transmit a copy of each notice submitted under this subsection to the regional office of the Environmental Protection Agency which has jurisdiction over the place of pesticide storage identified in the notice.

(h) Judicial review.—Final orders of the Administrator under this section shall be subject to judicial review pursuant to section 16.


(a) Requirement.—No person shall produce any pesticide subject to this Act or active ingredient used in producing a pesticide subject to this Act in any State unless the establishment in which it is produced is registered with the Administrator. The application for registration of any establishment shall include the name and address of the establishment and of the producer who operates such establishment.

(b) Registration.—Whenever the Administrator receives an application under subsection (a), the Administrator shall register the establishment and assign it an establishment number.

(c) Information required.—
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(1) Any producer operating an establishment registered under this section shall inform the Administrator within 30 days after it is registered of the types and amounts of pesticides and, if applicable, active ingredients used in producing pesticides—

(A) which the producer is currently producing;
(B) which the producer has produced during the past year; and
(C) which the producer has sold or distributed during the past year.

The information required by this paragraph shall be kept current and submitted to the Administrator annually as required under such regulations as the Administrator may prescribe.

(2) Any such producer shall, upon the request of the Administrator for the purpose of issuing a stop sale order pursuant to section 13, inform the Administrator of the name and address of any recipient of any pesticide produced in any registered establishment which the producer operates.

(d) CONFIDENTIAL RECORDS AND INFORMATION.—Any information submitted to the Administrator pursuant to subsection (c) other than the names of the pesticides or active ingredients used in producing pesticides produced, sold, or distributed at an establishment shall be considered confidential and shall be subject to the provisions of section 10.


(a) REQUIREMENTS.—The Administrator may prescribe regulations requiring producers, registrants, and applicants for registration to maintain such records with respect to their operations and the pesticides and devices produced as the Administrator determines are necessary for the effective enforcement of this Act and to make the records available for inspection and copying in the same manner as provided in subsection (b). No records required under this subsection shall extend to financial data, sales data other than shipment data, pricing data, personnel data, and research data (other than data relating to registered pesticides or to a pesticide for which an application for registration has been filed).

(b) INSPECTION.—For the purposes of enforcing the provisions of this Act, any producer, distributor, carrier, dealer, or any other person who sells or offers for sale, delivers or offers for delivery any pesticide or device subject to this Act, shall, upon request of any officer or employee of the Environmental Protection Agency or of any State or political subdivision, duly designated by the Administrator, furnish or permit such person at all reasonable times to have access to, and to copy: (1) all records showing the delivery, movement, or holding of such pesticide or device, including the quantity, the date of shipment and receipt, and the name of the consignor and consignee; or (2) in the event of the inability of any person to produce records containing such information, all other records and information relating to such delivery, movement, or holding of the pesticide or device. Any inspection with respect to any records and information referred to in this subsection shall not extend to financial data, sales data other than shipment data, pricing data, personnel data, and research data (other than data relat-
ing to registered pesticides or to a pesticide for which an application for registration has been filed). Before undertaking an inspection under this subsection, the officer or employee must present to the owner, operator, or agent in charge of the establishment or other place where pesticides or devices are held for distribution or sale, appropriate credentials and a written statement as to the reason for the inspection, including a statement as to whether a violation of the law is suspected. If no violation is suspected, an alternate and sufficient reason shall be given in writing. Each such inspection shall be commenced and completed with reasonable promptness.

SEC. 9. [7 U.S.C. 136g] INSPECTION OF ESTABLISHMENTS, ETC.

(a) IN GENERAL.—(1) For purposes of enforcing the provisions of this Act, officers or employees of the Environmental Protection Agency or of any State duly designated by the Administrator are authorized to enter at reasonable times (A) any establishment or other place where pesticides or devices are held for distribution or sale for the purpose of inspecting and obtaining samples of any pesticides or devices, packaged, labeled, and released for shipment, and samples of any containers or labeling for such pesticides or devices, or (B) any place where there is being held any pesticide the registration of which has been suspended or canceled for the purpose of determining compliance with section 19.

(2) Before undertaking such inspection, the officers or employees must present to the owner, operator, or agent in charge of the establishment or other place where pesticides or devices are held for distribution or sale, appropriate credentials and a written statement as to the reason for the inspection, including a statement as to whether a violation of the law is suspected. If no violation is suspected, an alternate and sufficient reason shall be given in writing. Each such inspection shall be commenced and completed with reasonable promptness. If the officer or employee obtains any samples, prior to leaving the premises, the officer or employee shall give to the owner, operator, or agent in charge a receipt describing the samples obtained and, if requested, a portion of each such sample equal in volume or weight to the portion retained. If an analysis is made of such samples, a copy of the results of such analysis shall be furnished promptly to the owner, operator, or agent in charge.

(b) WARRANTS.—For purposes of enforcing the provisions of this Act and upon a showing to an officer or court of competent jurisdiction that there is reason to believe that the provisions of this Act have been violated, officers or employees duly designated by the Administrator are empowered to obtain and to execute warrants authorizing—

(1) entry, inspection, and copying of records for purposes of this section or section 8;

(2) inspection and reproduction of all records showing the quantity, date of shipment, and the name of consignor and consignee of any pesticide or device found in the establishment which is adulterated, misbranded, not registered (in the case of a pesticide) or otherwise in violation of this Act and in the event of the inability of any person to produce records containing such information, all other records and information re-
lating to such delivery, movement, or holding of the pesticide or device; and
(3) the seizure of any pesticide or device which is in violation of this Act.
(c) Enforcement.—
(1) Certification of Facts to Attorney General.—The examination of pesticides or devices shall be made in the Environmental Protection Agency or elsewhere as the Administrator may designate for the purpose of determining from such examinations whether they comply with the requirements of this Act. If it shall appear from any such examination that they fail to comply with the requirements of this Act, the Administrator shall cause notice to be given to the person against whom criminal or civil proceedings are contemplated. Any person so notified shall be given an opportunity to present the person’s views, either orally or in writing, with regard to such contemplated proceedings, and if in the opinion of the Administrator it appears that the provisions of this Act have been violated by such person, then the Administrator shall certify the facts to the Attorney General, with a copy of the results of the analysis or the examination of such pesticide for the institution of a criminal proceeding pursuant to section 14(b) or a civil proceeding under section 14(a), when the Administrator determines that such action will be sufficient to effectuate the purposes of this Act.
(2) Notice Not Required.—The notice of contemplated proceedings and opportunity to present views set forth in this subsection are not prerequisites to the institution of any proceeding by the Attorney General.
(3) Warning Notices.—Nothing in this Act shall be construed as requiring the Administrator to institute proceedings for prosecution of minor violations of this Act whenever the Administrator believes that the public interest will be adequately served by a suitable written notice of warning.

(a) In General.—In submitting data required by this Act, the applicant may (1) clearly mark any portions thereof which in the applicant’s opinion are trade secrets or commercial or financial information and (2) submit such marked material separately from other material required to be submitted under this Act.
(b) Disclosure.—Notwithstanding any other provision of this Act and subject to the limitations in subsections (d) and (e) of this section, the Administrator shall not make public information which in the Administrator’s judgment contains or relates to trade secrets or commercial or financial information obtained from a person and privileged or confidential, except that, when necessary to carry out the provisions of this Act, information relating to formulas of products acquired by authorization of this Act may be revealed to any Federal agency consulted and may be revealed at a public hearing or in findings of fact issued by the Administrator.
(c) Disputes.—If the Administrator proposes to release for inspection information which the applicant or registrant believes to be protected from disclosure under subsection (b), the Adminis-
trator shall notify the applicant or registrant, in writing, by certified mail. The Administrator shall not thereafter make available for inspection such data until thirty days after receipt of the notice by the applicant or registrant. During this period, the applicant or registrant may institute an action in an appropriate district court for a declaratory judgment as to whether such information is subject to protection under subsection (b).

(d) LIMITATIONS.—

(1) All information concerning the objectives, methodology, results, or significance of any test or experiment performed on or with a registered or previously registered pesticide or its separate ingredients, impurities, or degradation products, and any information concerning the effects of such pesticide on any organism or the behavior of such pesticide in the environment, including, but not limited to, data on safety to fish and wildlife, humans and other mammals, plants, animals, and soil, and studies on persistence, translocation and fate in the environment, and metabolism, shall be available for disclosure to the public. The use of such data for any registration purpose shall be governed by section 3 of this Act. This paragraph does not authorize the disclosure of any information that—

(A) discloses manufacturing or quality control processes,

(B) discloses the details of any methods for testing, detecting, or measuring the quantity of any deliberately added inert ingredient of a pesticide, or

(C) discloses the identity or percentage quantity of any deliberately added inert ingredient of a pesticide, unless the Administrator has first determined that disclosure is necessary to protect against an unreasonable risk of injury to health or the environment.

(2) Information concerning production, distribution, sale, or inventories of a pesticide that is otherwise entitled to confidential treatment under subsection (b) of this section may be publicly disclosed in connection with a public proceeding to determine whether a pesticide, or any ingredient of a pesticide, causes unreasonable adverse effects on health or the environment, if the Administrator determines that such disclosure is necessary in the public interest.

(3) If the Administrator proposes to disclose information described in clause (A), (B), or (C) of paragraph (1) or in paragraph (2) of this subsection, the Administrator shall notify by certified mail the submitter of such information of the intent to release such information. The Administrator may not release such information, without the submitter's consent, until thirty days after the submitter has been furnished such notice. Where the Administrator finds that disclosure of information described in clause (A), (B), or (C) of paragraph (1) of this subsection is necessary to avoid or lessen an imminent and substantial risk of injury to the public health, the Administrator may set such shorter period of notice (but not less than ten days) and such method of notice as the Administrator finds appropriate. During such period the data submitter may institute an action in an appropriate district court to enjoin or limit the
proposed disclosure. The court may enjoin disclosure, or limit the disclosure or the parties to whom disclosure shall be made, to the extent that—

(A) in the case of information described in clause (A), (B), or (C) of paragraph (1) of this subsection, the proposed disclosure is not required to protect against an unreasonable risk of injury to health or the environment; or

(B) in the case of information described in paragraph (2) of this subsection, the public interest in availability of the information in the public proceeding does not outweigh the interests in preserving the confidentiality of the information.

(e) Disclosure to contractors.—Information otherwise protected from disclosure to the public under subsection (b) of this section may be disclosed to contractors with the United States and employees of such contractors if, in the opinion of the Administrator, such disclosure is necessary for the satisfactory performance by the contractor of a contract with the United States for the performance of work in connection with this Act and under such conditions as the Administrator may specify. The Administrator shall require as a condition to the disclosure of information under this subsection that the person receiving it take such security precautions respecting the information as the Administrator shall by regulation prescribe.

(f) Penalty for disclosure by federal employees.—(1) Any office or employee of the United States or former officer or employee of the United States who, by virtue of such employment or official position, has obtained possession of, or has access to, material the disclosure of which is prohibited by subsection (b) of this section, and who, knowing that disclosure of such material is prohibited by such subsection, willfully discloses the material in any manner to any person not entitled to receive it, shall be fined not more than $10,000 or imprisoned for not more than one year, or both. Section 1905 of title 18 of the United States Code shall not apply with respect to the publishing, divulging, disclosure, or making known of, or making available, information reported or otherwise obtained under this Act. Nothing in this Act shall preempt any civil remedy under State or Federal law for wrongful disclosure of trade secrets.

(2) For the purposes of this section, any contractor with the United States who is furnished information as authorized by subsection (e) of this section, or any employee of any such contractor, shall be considered to be an employee of the United States.

(g) Disclosure to foreign and multinational pesticide producers.—(1) The Administrator shall not knowingly disclose information submitted by an applicant or registrant under this Act to any employee or agent of any business or other entity engaged in the production, sale, or distribution of pesticides in countries other than the United States or in addition to the United States or to any other person who intends to deliver such data to such foreign or multinational business or entity unless the applicant or registrant has consented to such disclosure. The Administrator shall require an affirmation from any person who intends to inspect data that such person does not seek access to the data for purposes of
delivering it or offering it for sale to any such business or entity or its agents or employees and will not purposefully deliver or negligently cause the data to be delivered to such business or entity or its agents or employees. Notwithstanding any other provision of this subsection, the Administrator may disclose information to any person in connection with a public proceeding under law or regulation, subject to restrictions on the availability of information contained elsewhere in this Act, which information is relevant to a determination by the Administrator with respect to whether a pesticide, or any ingredient of a pesticide, causes unreasonable adverse effects on health or the environment.

(2) The Administrator shall maintain records of the names of persons to whom data are disclosed under this subsection and the persons or organizations they represent and shall inform the applicant or registrant of the names and affiliations of such persons.

(3) Section 1001 of title 18 of the United States Code shall apply to any affirmation made under paragraph (1) of this subsection.

SEC. 11. [7 U.S.C. 136i] USE OF RESTRICTED USE PESTICIDES; APPLICATORS.

(a) Certification Procedure.—

(1) Federal certification.—In any State for which a State plan for applicator certification has not been approved by the Administrator, the Administrator, in consultation with the Governor of such State, shall conduct a program for the certification of applicators of pesticides. Such program shall conform to the requirements imposed upon the States under the provisions of subsection (a)(2) of this section and shall not require private applicators to take any examination to establish competency in the use of pesticides. Prior to the implementation of the program, the Administrator shall publish in the Federal Register for review and comment a summary of the Federal plan for applicator certification and shall make generally available within the State copies of the plan. The Administrator shall hold public hearings at one or more locations within the State if so requested by the Governor of such State during the thirty days following publication of the Federal Register notice inviting comment on the Federal plan. The hearings shall be held within thirty days following receipt of the request from the Governor. In any State in which the Administrator conducts a certification program, the Administrator may require any person engaging in the commercial application, sale, offering for sale, holding for sale, or distribution of any pesticide one or more uses of which have been classified for restricted use to maintain such records and submit such reports concerning the commercial application, sale, or distribution of such pesticide as the Administrator may be regulation prescribe. Subject to paragraph (2), the Administrator shall prescribe standards for the certification of applicators of pesticides. Such standards shall provide that to be certified, an individual must be determined to be competent with respect to the use and handling of pesticides, or to the use and handling of the pesticide or class of pesticides covered by such individual’s certification. The certification standard for a private ap-
Applicator shall, under a State plan submitted for approval, be deemed fulfilled by the applicator completing a certification form. The Administrator shall further assure that such form contains adequate information and affirmations to carry out the intent of this Act, and may include in the form an affirmation that the private applicator has completed a training program approved by the Administrator so long as the program does not require the private applicator to take, pursuant to a requirement prescribed by the Administrator, any examination to establish competency in the use of the pesticide. The Administrator may require any pesticide dealer participating in a certification program to be licensed under a State licensing program approved by the Administrator.

(2) State Certification.—If any State, at any time, desires to certify applicators of pesticides, the Governor of such State shall submit a State plan for such purpose. The Administrator shall approve the plan submitted by any State, or any modification thereof, if such plan in the Administrator’s judgment—

(A) designates a State agency as the agency responsible for administering the plan throughout the State;

(B) contains satisfactory assurances that such agency has or will have the legal authority and qualified personnel necessary to carry out the plan;

(C) gives satisfactory assurances that the State will devote adequate funds to the administration of the plan;

(D) provides that the State agency will make such reports to the Administrator in such form and containing such information as the Administrator may from time to time require; and

(E) contains satisfactory assurances that State standards for the certification of applicators of pesticides conform with those standards prescribed by the Administrator under paragraph (1).

Any State certification program under this section shall be maintained in accordance with the State plan approved under this section.

(b) State Plans.—If the Administrator rejects a plan submitted under subsection (a)(2), the Administrator shall afford the State submitting the plan due notice and opportunity for hearing before so doing. If the Administrator approves a plan submitted under subsection (a)(2), then such State shall certify applicators of pesticides with respect to such State. Whenever the Administrator determines that a State is not administering the certification program in accordance with the plan approved under this section, the Administrator shall so notify the State and provide for a hearing at the request of the State, and, if appropriate corrective action is not taken within a reasonable time, not to exceed ninety days, the Administrator shall withdraw approval of such plan.

(c) Instruction in Integrated Pest Management Techniques.—Standards prescribed by the Administrator for the certification of applicators of pesticides under subsection (a), and the State plans submitted to the Administrator under subsection (a), shall include provisions for making instructional materials con-
cerning integrated pest management techniques available to individuals at their request in accordance with the provisions of section 23(c) of this Act, but such plans may not require that any individual receive instruction concerning such techniques or be shown to be competent with respect to the use of such techniques. The Administrator and States implementing such plans shall provide that all interested individuals are notified of the availability of such instructional materials.

(d) IN GENERAL.—No regulations prescribed by the Administrator for carrying out the provisions of this Act shall require any private applicator to maintain any records or file any reports or other documents.

(e) SEPARATE STANDARDS.—When establishing or approving standards for licensing or certification, the Administrator shall establish separate standards for commercial and private applicators.

SEC. 12. 17 U.S.C. 136j UNLAWFUL ACTS.

(a) IN GENERAL.—

(1) Except as provided by subsection (b), it shall be unlawful for any person in any State to distribute or sell to any person—

(A) any pesticide that is not registered under section 3 or whose registration has been canceled or suspended, except to the extent that distribution or sale otherwise has been authorized by the Administrator under this Act;

(B) any registered pesticide if any claims made for it as a part of its distribution or sale substantially differ from any claims made for it as a part of the statement required in connection with its registration under section 3;

(C) any registered pesticide the composition of which differs at the time of its distribution or sale from its composition as described in the statement required in connection with its registration under section 3;

(D) any pesticide which has not been colored or discolored pursuant to the provisions of section 25(c)(5);

(E) any pesticide which is adulterated or misbranded;

or

(F) any device which is misbranded.

(2) It shall be unlawful for any person—

(A) to detach, alter, deface, or destroy, in whole or in part, any labeling required under this Act;

(B) to refuse to—

(i) prepare, maintain, or submit any records required by or under section 5, 7, 8, 11, or 19;

(ii) submit any reports required by or under section 5, 6, 7, 8, 11, or 19; or

(iii) allow any entry, inspection, copying of records, or sampling authorized by this Act;

(C) to give a guaranty or undertaking provided for in subsection (b) which is false in any particular, except that a person who receives and relies upon a guaranty authorized under subsection (b) may give a guaranty to the same effect, which guaranty shall contain, in addition to the person’s own name and address, the name and address of the
person residing in the United States from whom the person received the guaranty or undertaking;

(D) to use for the person's own advantage or to reveal, other than to the Administrator, or officials or employees of the Environmental Protection Agency or other Federal executive agencies, or to the courts, or to physicians, pharmacists, and other qualified persons, needing such information for the performance of their duties, in accordance with such directions as the Administrator may prescribe, any information acquired by authority of this Act which is confidential under this Act;

(E) who is a registrant, wholesaler, dealer, retailer, or other distributor to advertise a product registered under this Act for restricted use without giving the classification of the product assigned to it under section 3;

(F) to distribute or sell, or to make available for use, or to use, any registered pesticide classified for restricted use for some or all purposes other than in accordance with section 3(d) and any regulations thereunder, except that it shall not be unlawful to sell, under regulations issued by the Administrator, a restricted use pesticide to a person who is not a certified applicator for application by a certified applicator;

(G) to use any registered pesticide in a manner inconsistent with its labeling;

(H) to use any pesticide which is under an experimental use permit contrary to the provisions of such permit;

(I) to violate any order issued under section 13;

(J) to violate any suspension order issued under section 3(c)(2)(B), 4, or 6;

(K) to violate any cancellation order issued under this Act or to fail to submit a notice in accordance with section 6(g);

(L) who is a producer to violate any of the provisions of section 7;

(M) to knowingly falsify all or part of any application for registration, application for experimental use permit, any information submitted to the Administrator pursuant to section 7, any records required to be maintained pursuant to this Act, any report filed under this Act, or any information marked as confidential and submitted to the Administrator under any provision of this Act;

(N) who is a registrant, wholesaler, dealer, retailer, or other distributor to fail to file reports required by this Act;

(O) to add any substance to, or take any substance from, any pesticide in a manner that may defeat the purpose of this Act;

(P) to use any pesticide in tests on human beings unless such human beings (i) are fully informed of the nature and purposes of the test and of any physical and mental health consequences which are reasonably foreseeable therefrom, and (ii) freely volunteer to participate in the test;
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| Q | to falsify all or part of any information relating to the testing of any pesticide (or any ingredient, metabolite, or degradation product thereof), including the nature of any protocol, procedure, substance, organism, or equipment used, observation made, or conclusion or opinion formed, submitted to the Administrator, or that the person knows will be furnished to the Administrator or will become a part of any records required to be maintained by this Act;

| R | to submit to the Administrator data known to be false in support of a registration; or

| S | to violate any regulation issued under section 3(a) or 19.

(b) Exemptions.—The penalties provided for a violation of paragraph (1) of subsection (a) shall not apply to—

1. any person who establishes a guaranty signed by, and containing the name and address of, the registrant or person residing in the United States from whom the person purchased or received in good faith the pesticide in the same unbroken package, to the effect that the pesticide was lawfully registered at the time of sale and delivery to the person, and that it complies with the other requirements of this Act, and in such case the guarantor shall be subject to the penalties which would otherwise attach to the person holding the guaranty under the provisions of this Act;

2. any carrier while lawfully shipping, transporting, or delivering for shipment any pesticide or device, if such carrier upon request of any officer or employee duly designated by the Administrator shall permit such officer or employee to copy all of its records concerning such pesticide or device;

3. any public official while engaged in the performance of the official duties of the public official;

4. any person using or possessing any pesticide as provided by an experimental use permit in effect with respect to such pesticide and such use or possession; or

5. any person who ships a substance or mixture of substances being put through tests in which the purpose is only to determine its value for pesticide purposes or to determine its toxicity or other properties and from which the user does not expect to receive any benefit in pest control from its use.


(a) Stop Sale, Etc., Orders.—Whenever any pesticide or device is found by the Administrator in any State and there is reason to believe on the basis of inspection or tests that such pesticide or device is in violation of any of the provisions of this Act, or that such pesticide or device has been or is intended to be distributed or sold in violation of any such provisions, or when the registration of the pesticide has been canceled by a final order or has been suspended, the Administrator may issue a written or printed “stop sale, use, or removal” order to any person who owns, controls, or has custody of such pesticide or device, and after receipt of such order no person shall sell, use, or remove the pesticide or device de-
scribed in the order except in accordance with the provisions of the order.

(b) Seizure.—Any pesticide or device that is being transported or, having been transported, remains unsold or in original unbroken packages, or that is sold or offered for sale in any State, or that is imported from a foreign country, shall be liable to be proceeding against in any district court in the district where it is found and seized for confiscation by a process in rem for condemnation if—

(1) in the case of a pesticide—
   (A) it is adulterated or misbranded;
   (B) it is not registered pursuant to the provisions of section 3;
   (C) its labeling fails to bear the information required by this Act;
   (D) it is not colored or discolored and such coloring or discoloring is required under this Act; or
   (E) any of the claims made for it or any of the directions for its use differ in substance from the representations made in connection with its registration;
(2) in the case of a device, it is misbranded; or
(3) in the case of a pesticide or device, when used in accordance with the requirements imposed under this Act and as directed by the labeling, it nevertheless causes unreasonable adverse effects on the environment.

In the case of a plant regulator, defoliant, or desiccant, used in accordance with the label claims and recommendations, physical or physiological effects on plants or parts thereof shall not be deemed to be injury, when such effects are the purpose for which the plant regulator, defoliant, or desiccant was applied.

(c) Disposition After Condemnation.—If the pesticide or device is condemned it shall, after entry of the decree, be disposed of by destruction or sale as the court may direct and the proceeds, if sold, less the court costs, shall be paid into the Treasury of the United States, but the pesticide or device shall not be sold contrary to the provisions of this Act or the laws of the jurisdiction in which it is sold. On payment of the costs of the condemnation proceedings and the execution and delivery of a good and sufficient bond conditioned that the pesticide or device shall not be sold or otherwise disposed of contrary to the provisions of the Act or the laws of any jurisdiction in which sold, the court may direct that such pesticide or device be delivered to the owner thereof. The proceedings of such condemnation cases shall conform, as near as may be to the proceedings in admiralty, except that either party may demand trial by jury of any issue of fact joined in any case, and all such proceedings shall be at the suit of and in the name of the United States.

(d) Court Costs, Etc.—When a decree of condemnation is entered against the pesticide or device, court costs and fees, storage, and other proper expenses shall be awarded against the person, if any, intervening as claimant of the pesticide or device.


(a) Civil Penalties.—
(1) **In General.**—Any registrant, commercial applicator, wholesaler, dealer, retailer, or other distributor who violates any provision of this Act may be assessed a civil penalty by the Administrator of not more than $5,000 for each offense.

(2) **Private Applicator.**—Any private applicator or other person not included in paragraph (1) who violates any provision of this Act subsequent to receiving a written warning from the Administrator or following a citation for a prior violation, may be assessed a civil penalty by the Administrator of not more than $1,000 for each offense, except that any applicator not included under paragraph (1) of this subsection who holds or applies registered pesticides, or uses dilutions of registered pesticides, only to provide a service of controlling pests without delivering any unapplied pesticide to any person so served, and who violates any provision of this Act may be assessed a civil penalty by the Administrator of not more than $500 for the first offense nor more than $1,000 for each subsequent offense.

(3) **Hearing.**—No civil penalty shall be assessed unless the person charged shall have been given notice and opportunity for a hearing on such charge in the county, parish, or incorporated city of the residence of the person charged.

(4) **Determination of Penalty.**—In determining the amount of the penalty, the Administrator shall consider the appropriateness of such penalty to the size of the business of the person charged, the effect on the person’s ability to continue in business, and the gravity of the violation. Whenever the Administrator finds that the violation occurred despite the exercise of due care or did not cause significant harm to health or the environment, the Administrator may issue a warning in lieu of assessing a penalty.

(5) **References to Attorney General.**—In case of inability to collect such civil penalty or failure of any person to pay all, or such portion of such civil penalty as the Administrator may determine, the Administrator shall refer the matter to the Attorney General, who shall recover such amount by action in the appropriate United States district court.

(b) **Criminal Penalties.**—

(1) **In General.**—

   (A) Any registrant, applicant for a registration, or producer who knowingly violates any provision of this Act shall be fined not more than $50,000 or imprisoned for not more than 1 year, or both.

   (B) Any commercial applicator of a restricted use pesticide, or any other person not described in subparagraph (A) who distributes or sells pesticides or devices, who knowingly violates any provision of this Act shall be fined not more than $25,000 or imprisoned for not more than 1 year, or both.

(2) **Private Applicator.**—Any private applicator or other person not included in paragraph (1) who knowingly violates any provision of this Act shall be guilty of a misdemeanor and shall on conviction be fined not more than $1,000, or imprisoned for not more than 30 days, or both.
(3) DISCLOSURE OF INFORMATION.—Any person, who, with intent to defraud, uses or reveals information relative to formulas of products acquired under the authority of section 3, shall be fined not more than $10,000, or imprisoned for not more than three years, or both.

(4) ACTS OF OFFICERS, AGENTS, ETC.—When construing and enforcing the provisions of this Act, the act, omission, or failure of any officer, agent, or other person acting for or employed by any person shall in every case be also deemed to be the act, omission, or failure of such person as well as that of the person employed.

SEC. 15. [7 U.S.C. 136m] INDEMNITIES.

(a) GENERAL INDEMNIFICATION.—

(1) IN GENERAL.—Except as otherwise provided in this section, if—

(A) the Administrator notifies a registrant under section 6(c)(1) that the Administrator intends to suspend a registration or that an emergency order of suspension of a registration under section 6(c)(3) has been issued;

(B) the registration in question is suspended under section 6(c), and thereafter is canceled under section 6(b), 6(d), or 6(f); and

(C) any person who owned any quantity of the pesticide immediately before the notice to the registrant under subparagraph (A) suffered losses by reason of suspension or cancellation of the registration;

the Administrator shall make an indemnity payment to the person.

(2) EXCEPTION.—Paragraph (1) shall not apply if the Administrator finds that the person—

(A) had knowledge of facts that, in themselves, would have shown that the pesticide did not meet the requirements of section 3(c)(5) for registration; and

(B) continued thereafter to produce the pesticide without giving timely notice of such facts to the Administrator.

(3) REPORT.—If the Administrator takes an action under paragraph (1) that requires the payment of indemnification, the Administrator shall report to the Committee on Agriculture of the House of Representatives, the Committee on Agriculture, Nutrition, and Forestry of the Senate, and the Committees on Appropriations of the House of Representatives and the Senate on—

(A) the action taken that requires the payment of indemnification;

(B) the reasons for taking the action;

(C) the estimated cost of the payment; and

(D) a request for the appropriation of funds for the payment.

(4) APPROPRIATION.—The Administrator may not make a payment of indemnification under paragraph (1) unless a specific line item appropriation of funds has been made in advance for the payment.
(b) Indemnification of End Users, Dealers, and Distributors.—

(1) END USERS.—If—

(A) the Administrator notifies a registrant under section 6(c)(1) that the Administrator intends to suspend a registration or that an emergency order of suspension of a registration under section 6(c)(3) has been issued;

(B) the registration in question is suspended under section 6(c), and thereafter is canceled under section 6(b), 6(d), or 6(f); and

(C) any person who, immediately before the notice to the registrant under subparagraph (A), owned any quantity of the pesticide for purposes of applying or using the pesticide as an end user, rather than for purposes of distributing or selling it or further processing it for distribution or sale, suffered a loss by reason of the suspension or cancellation of the pesticide;

the person shall be entitled to an indemnity payment under this subsection for such quantity of the pesticide.

(2) DEALERS AND DISTRIBUTORS.—

(A) Any registrant, wholesaler, dealer, or other distributor (hereinafter in this paragraph referred to as a “seller”) of a registered pesticide who distributes or sells the pesticide directly to any person not described as an end user in paragraph (1)(C) shall, with respect to any quantity of the pesticide that such person cannot use or resell as a result of the suspension or cancellation of the pesticide, reimburse such person for the cost of first acquiring the pesticide from the seller (other than the cost of transportation, if any), unless the seller provided to the person at the time of distribution or sale a notice, in writing, that the pesticide is not subject to reimbursement by the seller.

(B) If—

(i) the Administrator notifies a registrant under section 6(c)(1) that the Administrator intends to suspend a registration or that an emergency order of suspension of a registration under section 6(c)(3) has been issued;

(ii) the registration in question is suspended under section 6(c), and thereafter is canceled under section 6(b), 6(d), or 6(f);

(iii) any person who, immediately before the notice to the registrant under clause (i)—

(I) had not been notified in writing by the seller, as provided under subparagraph (A), that any quantity of the pesticide owned by such person is not subject to reimbursement by the seller in the event of suspension or cancellation of the pesticide; and

(II) owned any quantity of the pesticide for purposes of—

(aa) distributing or selling it; or

(bb) further processing it for distribution or sale directly to an end user;
suffered a loss by reason of the suspension or cancellation of the pesticide; and

(iv) the Administrator determines on the basis of a claim of loss submitted to the Administrator by the person, that the seller—

(I) did not provide the notice specified in subparagraph (A) to such person; and

(II) is and will continue to be unable to provide reimbursement to such person, as provided under subparagraph (A), for the loss referred to in clause (iii), as a result of the insolvency or bankruptcy of the seller and the seller's resulting inability to provide such reimbursement;

the person shall be entitled to an indemnity payment under this subsection for such quantity of the pesticide.

(C) If an indemnity payment is made by the United States under this paragraph, the United States shall be subrogated to any right that would otherwise be held under this paragraph by a seller who is unable to make a reimbursement in accordance with this paragraph with regard to reimbursements that otherwise would have been made by the seller.

(3) SOURCE.—Any payment required to be made under paragraph (1) or (2) shall be made from the appropriation provided under section 1304 of title 31, United States Code.

(4) ADMINISTRATIVE SETTLEMENT.—An administrative settlement of a claim for such indemnity may be made in accordance with the third paragraph of section 2414 of title 28, United States Code, and shall be regarded as if it were made under that section for purposes of section 1304 of title 31, United States Code.

(c) AMOUNT OF PAYMENT.—

(1) IN GENERAL.—The amount of an indemnity payment under subsection (a) or (b) to any person shall be determined on the basis of the cost of the pesticide owned by the person (other than the cost of transportation, if any) immediately before the issuance of the notice to the registrant referred to in subsection (a)(1)(A), (b)(1)(A), or (b)(2)(B)(i), except that in no event shall an indemnity payment to any person exceed the fair market value of the pesticide owned by the person immediately before the issuance of the notice.

(2) SPECIAL RULE.—Notwithstanding any other provision of this Act, the Administrator may provide a reasonable time for use or other disposal of the pesticide. In determining the quantity of any pesticide for which indemnity shall be paid under this section, proper adjustment shall be made for any pesticide used or otherwise disposed of by the owner.

SEC. 16. [7 U.S.C. 136n] ADMINISTRATIVE PROCEDURE; JUDICIAL REVIEW.

(a) DISTRICT COURT REVIEW.—Except as otherwise provided in this Act, the refusal of the Administrator to cancel or suspend a registration or to change a classification not following a hearing and other final actions of the Administrator not committed to the
discretion of the Administrator by law are judicially reviewable by the district courts of the United States.

(b) REVIEW BY COURT OF APPEALS.—In the case of actual controversy as to the validity of any order issued by the Administrator following a public hearing, any person who will be adversely affected by such order and who had been a party to the proceedings may obtain judicial review by filing in the United States court of appeals for the circuit wherein such person resides or has a place of business, within 60 days after the entry of such order, a petition praying that the order be set aside in whole or in part. A copy of the petition shall be forthwith transmitted by the clerk of the court to the Administrator or any officer designated by the Administrator for that purpose, and thereupon the Administrator shall file in the court the record of the proceedings on which the Administrator based the Administrator’s order, as provided in section 2112 of title 28, United States Code. Upon the filing of such petition the court shall have exclusive jurisdiction to affirm or set aside the order complained of in whole or in part. The court shall consider all evidence of record. The order of the Administrator shall be sustained if it is supported by substantial evidence when considered on the record as a whole. The judgment of the court affirming or setting aside, in whole or in part, any order under this section shall be final, subject to review by the Supreme Court of the United States upon certiorari or certification as provided in section 1254 of title 28 of the United States Code. The commencement of proceedings under this section shall not, unless specifically ordered by the court to the contrary, operate as a stay of an order.

(c) JURISDICTION OF DISTRICT COURTS.—The district courts of the United States are vested with jurisdiction specifically to enforce, and to prevent and restrain violations of, this Act.

(d) NOTICE OF JUDGMENTS.—The Administrator shall, by publication in such manner as the Administrator may prescribe, give notice of all judgments entered in actions instituted under the authority of this Act.

SEC. 17. [7 U.S.C. 136c] IMPORTS AND EXPORTS.

(a) PESTICIDES AND DEVICES INTENDED FOR EXPORT.—Notwithstanding any other provision this Act, no pesticide or device or active ingredient used in producing a pesticide intended solely for export to any foreign country shall be deemed in violation of this Act—

(1) when prepared or packed according to the specifications or directions of the foreign purchaser, except that producers of such pesticides and devices and active ingredients used in producing pesticides shall be subject to sections 2(p), 2(q) (1) (A), (C), (D), (E), (G), and (H), 2(q) (2) (A), (B), (C) (i) and (iii), and (D), 7, and 8 of this Act; and

(2) in the case of any pesticide other than a pesticide registered under section 3 or sold under section 6(a) (1) of this Act, if, prior to export, the foreign purchaser has signed a statement acknowledging that the purchaser understands that such pesticide is not registered for use in the United States and cannot be sold in the United States under this Act.
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A copy of that statement shall be transmitted to an appropriate official of the government of the importing country.

(b) CANCELLATION NOTICES FURNISHED TO FOREIGN GOVERNMENTS.—Whenever a registration, or a cancellation or suspension of the registration of a pesticide becomes effective, or ceases to be effective, the Administrator shall transmit through the State Department notification thereof to the governments of other countries and to appropriate international agencies. Such notification shall, upon request, include all information related to the cancellation or suspension of the registration of the pesticide and information concerning other pesticides that are registered under section 3 of this Act and that could be used in lieu of such pesticide.

(c) IMPORTATION OF PESTICIDES AND DEVICES.—

(1) IN GENERAL.—The Secretary of the Treasury shall notify the Administrator of the arrival of pesticides and devices and shall deliver to the Administrator, upon the Administrator’s request, samples of pesticides or devices which are being imported into the United States, giving notice to the owner or consignee, who may appear before the Administrator and have the right to introduce testimony. If it appears from the examination of a sample that it is adulterated, or misbranded or otherwise violates the provisions set forth in this Act, or is otherwise injurious to health or the environment, the pesticide or device may be refused admission, and the Secretary of the Treasury shall refuse delivery to the consignee and shall cause the destruction of any pesticide or device refused delivery which shall not be exported by the consignee within 90 days from the date of notice of such refusal under such regulations as the Secretary of the Treasury may prescribe. The Secretary of the Treasury may deliver to the consignee such pesticide or device pending examination and decision in the matter on execution of bond for the amount of the full invoice value of such pesticide or device, together with the duty thereon, and on refusal to return such pesticide or device for any cause to the custody of the Secretary of the Treasury, when demanded, for the purpose of excluding them from the country, or for any other purpose, said consignee shall forfeit the full amount of said bond. All charges for storage, cartage, and labor on pesticides or devices which are refused admission or delivery shall be paid by the owner or consignee, and in default of such payment shall constitute a lien against any future importation made by such owner or consignee.

(2) IMPORTATION OF SEED.—Notwithstanding any other provision of law, no person is required to notify the Administrator of the arrival of a plant-incorporated protectant (as defined in section 174.3 of title 40, Code of Federal Regulations (or any successor regulation)) that is contained in a seed, if—

(A) that plant-incorporated protectant is registered under section 3;

(B) the Administrator has issued an experimental use permit for that plant-incorporated protectant under section 5; or
(C) the seed is covered by a permit (as defined in part 340 of title 7, Code of Federal Regulations (or any successor regulation)) or a notification.

(3) COOPERATION.—

(A) IN GENERAL.—In response to a request from the Administrator, the Secretary of Agriculture shall provide to the Administrator a list of seed containing plant-incorporated protectants (as defined in section 174.3 of title 40, Code of Federal Regulations (or any successor regulation)) if the importation of that seed into the United States has been approved under a permit or notification referred to in paragraph (2).

(B) CONTENTS.—The list under subparagraph (A) shall be provided in a form and at such intervals as may be agreed to by the Secretary and the Administrator.

(4) APPLICABILITY.—Nothing in this subsection precludes or limits the authority of the Secretary of Agriculture with respect to the importation or movement of plants, plant products, or seeds under—

(A) the Plant Protection Act (7 U.S.C. 7701 et seq.); and

(B) the Federal Seed Act (7 U.S.C. 1551 et seq.).

(d) COOPERATION IN INTERNATIONAL EFFORTS.—

(1) IN GENERAL.—The Administrator shall, in cooperation with the Department of State and any other appropriate Federal agency, participate and cooperate in any international efforts to develop improved pesticide research and regulations.

(2) DEPARTMENT OF STATE EXPENSES.—Any expenses incurred by an employee of the Environmental Protection Agency who participates in any international technical, economic, or policy review board, committee, or other official body that is meeting in relation to an international treaty shall be paid by the Department of State.

(e) REGULATIONS.—The Secretary of the Treasury, in consultation with the Administrator, shall prescribe regulations for the enforcement of subsection (c) of this section.


The Administrator may, at the Administrator’s discretion, exempt any Federal or State agency from any provision of this Act if the Administrator determines that emergency conditions exist which require such exemption. The Administrator, in determining whether or not such emergency conditions exist, shall consult with the Secretary of Agriculture and the Governor of any State concerned if they request such determination.


(a) STORAGE, DISPOSAL, AND TRANSPORTATION.—

(1) DATA REQUIREMENTS AND REGISTRATION OF PESTICIDES.—The Administrator may require under section 3 or 6 that—

(A) the registrant or applicant for registration of a pesticide submit or cite data or information regarding meth-
ods for the safe storage and disposal of excess quantities of the pesticide to support the registration or continued registration of a pesticide;

(B) the labeling of a pesticide contain requirements and procedures for the transportation, storage, and disposal of the pesticide, any container of the pesticide, any rinsate containing the pesticide, or any other material used to contain or collect excess or spilled quantities of the pesticide; and

(C) the registrant of a pesticide provide evidence of sufficient financial and other resources to carry out a recall plan under subsection (b), and provide for the disposal of the pesticide, in the event of suspension and cancellation of the pesticide.

(2) PESTICIDES.—The Administrator may by regulation, or as part of an order issued under section 6 or an amendment to such an order—

(A) issue requirements and procedures to be followed by any person who stores or transports a pesticide the registration of which has been suspended or canceled;

(B) issue requirements and procedures to be followed by any person who disposes of stocks of a pesticide the registration of which has been suspended; and

(C) issue requirements and procedures for the disposal of any pesticide the registration of which has been canceled.

(3) CONTAINERS, RINSATES, AND OTHER MATERIALS.—The Administrator may by regulation, or as part of an order issued under section 6 or an amendment to such an order—

(A) issue requirements and procedures to be followed by any person who stores or transports any container of a pesticide the registration of which has been suspended or canceled, any rinsate containing the pesticide, or any other material used to contain or collect excess or spilled quantities of the pesticide;

(B) issue requirements and procedures to be followed by any person who disposes of stocks of any container of a pesticide the registration of which has been suspended, any rinsate containing the pesticide, or any other material used to contain or collect excess or spilled quantities of the pesticide; and

(C) issue requirements and procedures for the disposal of any container of a pesticide the registration of which has been canceled, any rinsate containing the pesticide, or any other material used to contain or collect excess or spilled quantities of the pesticide.

(4) CONTAINER RECYCLING.—The Secretary may promulgate a regulation for the return and recycling of disposable pesticide containers used for the distribution or sale of registered pesticide products in interstate commerce. Any such regulation requiring recycling of disposable pesticide containers shall not apply to antimicrobial pesticides (as defined in section 2) or other pesticide products intended for non-agricultural uses.

(b) RECALLS.—
(1) In general.—If the registration of a pesticide has been suspended and canceled under section 6, and if the Administrator finds that recall of the pesticide is necessary to protect health or the environment, the Administrator shall order a recall of the pesticide in accordance with this subsection.

(2) Voluntary recall.—If, after determining under paragraph (1) that a recall is necessary, the Administrator finds that voluntary recall by the registrant and others in the chain of distribution may be as safe and effective as a mandatory recall, the Administrator shall request the registrant of the pesticide to submit, within 60 days of the request, a plan for the voluntary recall of the pesticide. If such a plan is requested and submitted, the Administrator shall approve the plan and order the registrant to conduct the recall in accordance with the plan unless the Administrator determines, after an informal hearing, that the plan is inadequate to protect health or the environment.

(3) Mandatory recall.—If, after determining under paragraph (1) that a recall is necessary, the Administrator does not request the submission of a plan under paragraph (2) or finds such a plan to be inadequate, the Administrator shall issue a regulation that prescribes a plan for the recall of the pesticide. A regulation issued under this paragraph may apply to any person who is or was a registrant, distributor, or seller of the pesticide, or any successor in interest to such a person.

(4) Recall procedure.—A regulation issued under this subsection may require any person that is subject to the regulation to—

(A) arrange to make available one or more storage facilities to receive and store the pesticide to which the recall program applies, and inform the Administrator of the location of each such facility;

(B) accept and store at such a facility those existing stocks of such pesticide that are tendered by any other person who obtained the pesticide directly or indirectly from the person that is subject to such regulation;

(C) on the request of a person making such a tender, provide for proper transportation of the pesticide to a storage facility; and

(D) take such reasonable steps as the regulation may prescribe to inform persons who may be holders of the pesticide of the terms of the recall regulation and how those persons may tender the pesticide and arrange for transportation of the pesticide to a storage facility.

(5) Contents of recall plan.—A recall plan established under this subsection shall include—

(A) the level in the distribution chain to which the recall is to extend, and a schedule for recall; and

(B) the means to be used to verify the effectiveness of the recall.

(6) Requirements or procedures.—No requirement or procedure imposed in accordance with paragraph (2) of subsection (a) may require the recall of existing stocks of the pesticide except as provided by this subsection.
(c) Storage Costs.—

(1) Submission of Plan.—A registrant who wishes to become eligible for reimbursement of storage costs incurred as a result of a recall prescribed under subsection (b) for a pesticide whose registration has been suspended and canceled shall, as soon as practicable after the suspension of the registration of the pesticide, submit to the Administrator a plan for the storage and disposal of the pesticide that meets criteria established by the Administrator by regulation.

(2) Reimbursement.—Within a reasonable period of time after such storage costs are incurred and paid by the registrant, the Administrator shall reimburse the registrant, on request, for—

(A) none of the costs incurred by the registrant before the date of submission of the plan referred to in paragraph (1) to the Administrator;

(B) 100 percent of the costs incurred by the registrant after the date of submission of the plan to the Administrator or the date of cancellation of the registration of the pesticide, whichever is later, but before the approval of the plan by the Administrator;

(C) 50 percent of the costs incurred by the registrant during the 1-year period beginning on the date of the approval of the plan by the Administrator or the date of cancellation of the registration of the pesticide, whichever is later;

(D) none of the costs incurred by the registrant during the 3-year period beginning on the 366th day following approval of the plan by the Administrator or the date of cancellation of the registration of the pesticide, whichever is later; and

(E) 25 percent of the costs incurred by the registrant during the period beginning on the first day of the 5th year following the date of the approval of the plan by the Administrator or the date of cancellation of the registration of the pesticide, whichever is later, and ending on the date that a disposal permit for the pesticide is issued by a State or an alternative plan for disposal of the pesticide in accordance with applicable law has been developed.

(d) Administration of Storage, Disposal, Transportation, and Recall Programs.—

(1) Voluntary Agreements.—Nothing in this section shall be construed as preventing or making unlawful any agreement between a seller and a buyer of any pesticide or other substance regarding the ultimate allocation of the costs of storage, transportation, or disposal of a pesticide.

(2) Rule and Regulation Review.—Section 25(a)(4) shall not apply to any regulation issued under subsection (a)(2) or (b).

(3) Limitations.—No registrant shall be responsible under this section for a pesticide the registration of which is held by another person. No distributor or seller shall be responsible under this section for a pesticide that the distributor or seller did not hold or sell.
(4) **Seizure and Penalties.**—If the Administrator finds that a person who is subject to a regulation or order under subsection (a)(2) or (b) has failed substantially to comply with that regulation or order, the Administrator may take action under section 13 or 14 or obtain injunctive relief under section 16(c) against such person or any successor in interest of any such person.

(e) **Container Design.**—

(1) **Procedures.**—

(A) Not later than 3 years after the effective date of this subsection [December 24, 1988], the Administrator shall, in consultation with the heads of other interested Federal agencies, promulgate regulations for the design of pesticide containers that will promote the safe storage and disposal of pesticides.

(B) The regulations shall ensure, to the fullest extent practicable, that the containers—

(i) accommodate procedures used for the removal of pesticides from the containers and the rinsing of the containers;

(ii) facilitate the safe use of the containers, including elimination of splash and leakage of pesticides from the containers;

(iii) facilitate the safe disposal of the containers; and

(iv) facilitate the safe refill and reuse of the containers.

(2) **Compliance.**—The Administrator shall require compliance with the regulations referred to in paragraph (1) not later than 5 years after the effective date of this subsection [December 24, 1988].

(f) **Pesticide Residue Removal.**—

(1) **Procedures.**—

(A) Not later than 3 years after the effective date of this subsection [December 24, 1988], the Administrator shall, in consultation with the heads of other interested Federal agencies, promulgate regulations prescribing procedures and standards for the removal of pesticides from containers prior to disposal.

(B) The regulations may—

(i) specify, for each major type of pesticide container, procedures and standards providing for, at a minimum, triple rinsing or the equivalent degree of pesticide removal;

(ii) specify procedures that can be implemented promptly and easily in various circumstances and conditions;

(iii) provide for reuse, whenever practicable, or disposal of rinse water and residue; and

(iv) be coordinated with requirements for the rinsing of containers imposed under the Solid Waste Disposal Act (42 U.S.C. 6901 et seq.).
(C) The Administrator may, at the discretion of the Administrator, exempt products intended solely for household use from the requirements of this subsection.

(2) COMPLIANCE.—Effective beginning 5 years after the effective date of this subsection [December 24, 1988], a State may not exercise primary enforcement responsibility under section 26, or certify an applicator under section 11, unless the Administrator determines that the State is carrying out an adequate program to ensure compliance with this subsection.

(3) SOLID WASTE DISPOSAL ACT.—Nothing in this subsection shall affect the authorities or requirements concerning pesticide containers under the Solid Waste Disposal Act (42 U.S.C. 6901).

(g) PESTICIDE CONTAINER STUDY.—

(1) STUDY.—

(A) The Administrator shall conduct a study of options to encourage or require—

(i) the return, refill, and reuse of pesticide containers;

(ii) the development and use of pesticide formulations that facilitate the removal of pesticide residues from containers; and

(iii) the use of bulk storage facilities to reduce the number of pesticide containers requiring disposal.

(B) In conducting the study, the Administrator shall—

(i) consult with the heads of other interested Federal agencies, State agencies, industry groups, and environmental organizations; and

(ii) assess the feasibility, costs, and environmental benefits of encouraging or requiring various measures or actions.

(2) REPORT.—Not later than 2 years after the effective date of this subsection [December 24, 1988], the Administrator shall submit to Congress a report describing the results of the study required under paragraph (1).

(h) RELATIONSHIP TO SOLID WASTE DISPOSAL ACT.—

(1) IN GENERAL.—Nothing in this section shall diminish the authorities or requirements of the Solid Waste Disposal Act (42 U.S.C. 6901 et seq.).

(2) ANTIMICROBIAL PRODUCTS.—A household, industrial, or institutional antimicrobial product that is not subject to regulation under the Solid Waste Disposal Act (42 U.S.C. 6901 et seq.) shall not be subject to the provisions of subsections (a), (e), and (f), unless the Administrator determines that such product must be subject to such provisions to prevent an unreasonable adverse effect on the environment.


(a) RESEARCH.—The Administrator shall undertake research, including research by grant or contract with other Federal agencies, universities, or others as may be necessary to carry out the purposes of this Act, and the Administrator shall conduct research into integrated pest management in coordination with the Secretary of Agriculture. The Administrator shall also take care to en-
sure that such research does not duplicate research being undertaken by any other Federal agency.

(b) NATIONAL MONITORING PLAN.—The Administrator shall formulate and periodically revise, in cooperation with other Federal, State, or local agencies, a national plan for monitoring pesticides.

(c) MONITORING.—The Administrator shall undertake such monitoring activities, including, but not limited to monitoring in air, soil, water, man, plants, and animals, as may be necessary for the implementation of this Act and of the national pesticide monitoring plan. The Administrator shall establish procedures for the monitoring of man and animals and their environment for incidental pesticide exposure, including, but not limited to, the quantification of incidental human and environmental pesticide pollution and the secular trends thereof, and identification of the sources of contamination and their relationship to human and environmental effects. Such activities shall be carried out in cooperation with other Federal, State, and local agencies.

SEC. 21. [7 U.S.C. 136s] SOLICITATION OF COMMENTS; NOTICE OF PUBLIC HEARINGS.

(a) SECRETARY OF AGRICULTURE.—The Administrator, before publishing regulations under this Act, shall solicit the views of the Secretary of Agriculture in accordance with the procedure described in section 25(a).

(b) SECRETARY OF HEALTH AND HUMAN SERVICES.—The Administrator, before publishing regulations under this Act for any public health pesticide, shall solicit the views of the Secretary of Health and Human Services in the same manner as the views of the Secretary of Agriculture are solicited under section 25(a)(2).

(c) VIEWS.—In addition to any other authority relating to public hearings and solicitation of views, in connection with the suspension or cancellation of a pesticide registration or any other actions authorized under this Act, the Administrator may, at the Administrator's discretion, solicit the views of all interested persons, either orally or in writing, and seek such advice from scientists, farmers, farm organizations, and other qualified persons as the Administrator deems proper.

(d) NOTICE.—In connection with all public hearings under this Act the Administrator shall publish timely notice of such hearings in the Federal Register.

SEC. 22. [7 U.S.C. 136t] DELEGATION AND COOPERATION.

(a) DELEGATION.—All authority vested in the Administrator by virtue of the provisions of this Act may with like force and effect be executed by such employees of the Environmental Protection Agency as the Administrator may designate for the purpose.

(b) COOPERATION.—The Administrator shall cooperate with the Department of Agriculture, any other Federal agency, and any appropriate agency of any State or any political subdivision thereof, in carrying out the provisions of this Act, and in securing uniformity of regulations.

16So in original. Probably should be “incidental”.

September 3, 2019
As Amended Through P.L. 116-8, Enacted March 08, 2019

(a) COOPERATIVE AGREEMENTS.—The Administrator may enter into cooperative agreements with States and Indian tribes—

(1) to delegate to any State or Indian tribe the authority to cooperate in the enforcement of this Act through the use of its personnel or facilities, to train personnel of the State or Indian tribe to cooperate in the enforcement of this Act, and to assist States and Indian tribes in implementing cooperative enforcement programs through grants-in-aid; and

(2) to assist States in developing and administering State programs, and Indian tribes that enter into cooperative agreements, to train and certify applicators consistent with the standards the Administrator prescribes.

Effective with the fiscal year beginning October 1, 1978, there are authorized to be appropriated annually such funds as may be necessary for the Administrator to provide through cooperative agreements an amount equal to 50 percent of the anticipated cost to each State or Indian tribe, as agreed to under such cooperative agreements, of conducting training and certification programs during such fiscal year. If funds sufficient to pay 50 percent of the costs for any year are not appropriated, the share of each State and Indian tribe shall be reduced in a like proportion in allocating available funds.

(b) CONTRACTS FOR TRAINING.—In addition, the Administrator may enter into contracts with Federal, State, or Indian tribal agencies for the purpose of encouraging the training of certified applicators.

(c) INFORMATION AND EDUCATION.—The Administrator shall, in cooperation with the Secretary of Agriculture, use the services of the cooperative State extension services to inform and educate pesticide users about accepted uses and other regulations made under this Act.


(a) IN GENERAL.—A State may regulate the sale or use of any federally registered pesticide or device in the State, but only if and to the extent the regulation does not permit any sale or use prohibited by this Act.

(b) UNIFORMITY.—Such State shall not impose or continue in effect any requirements for labeling or packaging in addition to or different from those required under this Act.

(c) ADDITIONAL USES.—

(1) A State may provide registration for additional uses of federally registered pesticides formulated for distribution and use within that State to meet special local needs in accord with the purposes of this Act and if registration for such use has not previously been denied, disapproved, or canceled by the Administrator. Such registration shall be deemed registration under section 3 for all purposes of this Act, but shall authorize distribution and use only within such State.

(2) A registration issued by a State under this subsection shall not be effective for more than ninety days if disapproved by the Administrator within that period. Prior to disapproval, the Administrator shall, except as provided in paragraph (3) of this subsection, advise the State of the Administrator’s inten-
tion to disapprove and the reasons therefor, and provide the State time to respond. The Administrator shall not prohibit or disapprove a registration issued by a State under this subsection (A) on the basis of lack of essentiality of a pesticide or (B) except as provided in paragraph (3) of this subsection, if its composition and use patterns are similar to those of a federally registered pesticide.

(3) In no instance may a State issue a registration for a food or feed use unless there exists a tolerance or exemption under the Federal Food, Drug, and Cosmetic Act that permits the residues of the pesticide on the food or feed. If the Administrator determines that a registration issued by a State is inconsistent with the Federal Food, Drug, and Cosmetic Act, or the use of, a pesticide under a registration issued by a State constitutes an imminent hazard, the Administrator may immediately disapprove the registration.

(4) If the Administrator finds, in accordance with standards set forth in regulations issued under section 25 of this Act, that a State is not capable of exercising adequate controls to assure that State registration under this section will be in accord with the purposes of this Act or has failed to exercise adequate controls, the Administrator may suspend the authority of the State to register pesticides until such time as the Administrator is satisfied that the State can and will exercise adequate controls. Prior to any such suspension, the Administrator shall advise the State of the Administrator’s intention to suspend and the reasons therefor and provide the State time to respond.


(a) IN GENERAL.—

(1) REGULATIONS.—The Administrator is authorized in accordance with the procedure described in paragraph (2), to prescribe regulations to carry out the provisions of this Act. Such regulations shall take into account the difference in concept and usage between various classes of pesticides, including public health pesticides, and differences in environmental risk and the appropriate data for evaluating such risk between agricultural, nonagricultural, and public health pesticides.

(2) PROCEDURE.—

(A) PROPOSED REGULATIONS.—At least 60 days prior to signing any proposed regulation for publication in the Federal Register, the Administrator shall provide the Secretary of Agriculture with a copy of such regulation. If the Secretary comments in writing to the Administrator regarding any such regulation within 30 days after receiving it, the Administrator shall publish in the Federal Register (with the proposed regulation) the comments of the Secretary and the response of the Administrator with regard to the Secretary’s comments. If the Secretary does not comment in writing to the Administrator regarding the regulation within 30 days after receiving it, the Administrator may sign such regulation for publication in the Fed-
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(a) NOTIFICATION TO SECRETARY.—At any time after such 30-day period notwithstanding the foregoing 60-day time requirement.

(B) Final Regulations.—At least 30 days prior to signing any regulation in final form for publication in the Federal Register, the Administrator shall provide the Secretary of Agriculture with a copy of such regulation. If the Secretary comments in writing to the Administrator regarding any such final regulation within 15 days after receiving it, the Administrator shall publish in the Federal Register (with the final regulation) the comments of the Secretary, if requested by the Secretary, and the response of the Administrator concerning the Secretary’s comments. If the Secretary does not comment in writing to the Administrator regarding the regulation within 15 days after receiving it, the Administrator may sign such regulation for publication in the Federal Register at any time after such 15-day period notwithstanding the foregoing 30-day time requirement. In taking any final action under this subsection, the Administrator shall include among those factors to be taken into account the effect of the regulation on production and prices of agricultural commodities, retail food prices, and otherwise on the agricultural economy, and the Administrator shall publish in the Federal Register an analysis of such effect.

(C) Time Requirements.—The time requirements imposed by subparagraphs (A) and (B) may be waived or modified to the extent agreed upon by the Administrator and the Secretary.

(D) Publication in the Federal Register.—The Administrator shall, simultaneously with any notification to the Secretary of Agriculture under this paragraph prior to the issuance of any proposed or final regulation, publish such notification in the Federal Register.

(3) Congressional Committees.—At such time as the Administrator is required under paragraph (2) of this subsection to provide the Secretary of Agriculture with a copy of proposed regulations and a copy of the final form of regulations, the Administrator shall also furnish a copy of such regulations to the Committee on Agriculture of the House of Representatives and the Committee on Agriculture, Nutrition, and Forestry of the Senate.

(4) Congressional Review of Regulations.—Simultaneously with the promulgation of any rule or regulation under this Act, the Administrator shall transmit a copy thereof to the Secretary of the Senate and the Clerk of the House of Representatives. The rule or regulation shall not become effective until the passage of 60 calendar days after the rule or regulation is so transmitted.

(b) Exemption of Pesticides.—The Administrator may exempt from the requirements of this Act by regulation any pesticide which the Administrator determines either (1) to be adequately regulated by another Federal agency, or (2) to be of a character which is unnecessary to be subject to this Act in order to carry out the purposes of this Act.
(c) Other Authority.—The Administrator, after notice and opportunity for hearing, is authorized—

(1) to declare a pest any form of plant or animal life (other than man and other than bacteria, virus, and other micro-organisms on or in living man or other living animals) which is injurious to health or the environment;

(2) to determine any pesticide which contains any substance or substances in quantities highly toxic to man;

(3) to establish standards (which shall be consistent with those established under the authority of the Poison Prevention Packaging Act (Public Law 91–601)) with respect to the package, container, or wrapping in which a pesticide or device is enclosed for use or consumption, in order to protect children and adults from serious injury or illness resulting from accidental ingestion or contact with pesticides or devices regulated by this Act as well as to accomplish the other purposes of this Act;

(4) to specify those classes of devices which shall be subject to any provision of paragraph (1) or section 7 of this Act upon the Administrator's determination that application of such provision is necessary to effectuate the purposes of this Act;

(5) to prescribe regulations requiring any pesticide to be colored or discolored if the Administrator determines that such requirement is feasible and is necessary for the protection of health and the environment; and

(6) to determine and establish suitable names to be used in the ingredient statement.

(d) Scientific Advisory Panel.—

(1) In general.—The Administrator shall submit to an advisory panel for comment as to the impact on health and the environment of the action proposed in notices of intent issued under section 6(b) and of the proposed and final form of regulations issued under section 25(a) within the same time periods as provided for the comments of the Secretary of Agriculture under such sections. The time requirements for notices of intent and proposed and final forms of regulation may not be modified or waived unless in addition to meeting the requirements of section 6(b) or 25(a), as applicable, the advisory panel has failed to comment on the proposed action within the prescribed time period or has agreed to the modification or waiver. The Administrator shall also solicit from the advisory panel comments, evaluations, and recommendations for operating guidelines to improve the effectiveness and quality of scientific analyses made by personnel of the Environmental Protection Agency that lead to decisions by the Administrator in carrying out the provisions of this Act. The comments, evaluations, and recommendations of the advisory panel submitted under this subsection and the response of the Administrator shall be published in the Federal Register in the same manner as provided for publication of the comments of the Secretary of Agriculture under such sections. The chairman of the advisory panel, after consultation with the Administrator, may create temporary subpanels on specific projects to assist the full advisory panel
in expediting and preparing its evaluations, comments, and recommendations. The subpanels may be composed of scientists other than members of the advisory panel, as deemed necessary for the purpose of evaluating scientific studies relied upon by the Administrator with respect to proposed action. Such additional scientists shall be selected by the advisory panel. The panel referred to in this subsection shall consist of 7 members appointed by the Administrator from a list of 12 nominees, 6 nominated by the National Institutes of Health and 6 by the National Science Foundation, utilizing a system of staggered terms of appointment. Members of the panel shall be selected on the basis of their professional qualifications to assess the effects of the impact of pesticides on health and the environment. To the extent feasible to insure multidisciplinary representation, the panel membership shall include representation from the disciplines of toxicology, pathology, environmental biology, and related sciences. If a vacancy occurs on the panel due to expiration of a term, resignation, or any other reason, each replacement shall be selected by the Administrator from a group of 4 nominees, 2 submitted by each of the nominating entities named in this subsection. The Administrator may extend the term of a panel member until the new member is appointed to fill the vacancy. If a vacancy occurs due to resignation, or reason other than expiration of a term, the Administrator shall appoint a member to serve during the unexpired term utilizing the nomination process set forth in this subsection. Should the list of nominees provided under this subsection be unsatisfactory, the Administrator may request an additional set of nominees from the nominating entities. The Administrator may require such information from the nominees to the advisory panel as the Administrator deems necessary, and the Administrator shall publish in the Federal Register the name, address, and professional affiliations of each nominee. Each member of the panel shall receive per diem compensation at a rate not in excess of that fixed for GS–18 of the General Schedule as may be determined by the Administrator, except that any such member who holds another office or position under the Federal Government the compensation for which exceeds such rate may elect to receive compensation at the rate provided for such other office or position in lieu of the compensation provided by this subsection. In order to assure the objectivity of the advisory panel, the Administrator shall promulgate regulations regarding conflicts of interest with respect to the members of the panel. The advisory panel established under this section shall be permanent. In performing the functions assigned by this Act, the panel shall consult and coordinate its activities with the Science Advisory Board established under the Environmental Research, Development, and Demonstration Authorization Act of 1978. Whenever the Administrator exercises authority under section 6(c) of this Act to immediately suspend the registration of any pesticide to prevent an imminent hazard, the Administrator shall promptly submit to the advisory panel for comment, as
to the impact on health and the environment, the action taken to suspend the registration of such pesticide.

(2) **SCIENCE REVIEW BOARD.**—There is established a Science Review Board to consist of 60 scientists who shall be available to the Scientific Advisory Panel to assist in reviews conducted by the Panel. Members of the Board shall be selected in the same manner as members of temporary subpanels created under paragraph (1). Members of the Board shall be compensated in the same manner as members of the Panel.

(e) **PEER REVIEW.**—The Administrator shall, by written procedures, provide for peer review with respect to the design, protocols, and conduct of major scientific studies conducted under this Act by the Environmental Protection Agency or by any other Federal agency, any State or political subdivision thereof, or any institution or individual under grant, contract, or cooperative agreement from or with the Environmental Protection Agency. In such procedures, the Administrator shall also provide for peer review, using the advisory panel established under subsection (d) of this section or appropriate experts appointed by the Administrator from a current list of nominees maintained by such panel, with respect to the results of any such scientific studies relied upon by the Administrator with respect to actions the Administrator may take relating to the change in classification, suspension, or cancellation of a pesticide.

Whenever the Administrator determines that circumstances do not permit the peer review of the results of any such scientific study prior to the Administrator’s exercising authority under section 6(c) of this Act to immediately suspend the registration of any pesticide to prevent an imminent hazard, the Administrator shall promptly thereafter provide for the conduct of peer review as provided in this sentence. The evaluations and relevant documentation constituting the peer review that relate to the proposed scientific studies and the results of the completed scientific studies shall be included in the submission for comment forwarded by the Administrator to the advisory panel as provided in subsection (d). As used in this subsection, the term “peer review” shall mean an independent evaluation by scientific experts, either within or outside the Environmental Protection Agency, in the appropriate disciplines.


(a) **IN GENERAL.**—For the purposes of this Act, a State shall have primary enforcement responsibility for pesticide use violations during any period for which the Administrator determines that such State—

(1) has adopted adequate pesticide use laws and regulations, except that the Administrator may not require a State to have pesticide use laws that are more stringent than this Act;

(2) has adopted and is implementing adequate procedures for the enforcement of such State laws and regulations; and

(3) will keep such records and make such reports showing compliance with paragraphs (1) and (2) of this subsection as the Administrator may require by regulation.

(b) **SPECIAL RULES.**—Notwithstanding the provisions of subsection (a) of this section, any State that enters into a cooperative...
agreement with the Administrator under section 23 of this Act for the enforcement of pesticide use restrictions shall have the primary enforcement responsibility for pesticide use violations. Any State that has a plan approved by the Administrator in accordance with the requirements of section 11 of this Act that the Administrator determines meets the criteria set out in subsection (a) of this section shall have the primary enforcement responsibility for pesticide use violations. The Administrator shall make such determinations with respect to State plans under section 11 of this Act in effect on the date of enactment of the Federal Pesticide Act of 1978 [September 30, 1978] not later than six months after that date.

(c) ADMINISTRATOR.—The Administrator shall have primary enforcement responsibility for those States that do not have primary enforcement responsibility under this Act. Notwithstanding the provisions of section 2(e)(1) of this Act, during any period when the Administrator has such enforcement responsibility, section 8(b) of this Act shall apply to the books and records of commercial applicators and to any applicator who holds or applies pesticides, or uses dilutions of pesticides, only to provide a service of controlling pests without delivering any unapplied pesticide to any person so served, and section 9(a) of this Act shall apply to the establishment or other place where pesticides or devices are held for application by such persons with respect to pesticides or devices held for such application.

SEC. 27. [7 U.S.C. 136w–2] FAILURE BY THE STATE TO ASSURE ENFORCEMENT OF STATE PESTICIDE USE REGULATIONS.

(a) REFERRAL.—Upon receipt of any complaint or other information alleging or indicating a significant violation of the pesticide use provisions of this Act, the Administrator shall refer the matter to the appropriate State officials for their investigation of the matter consistent with the requirements of this Act. If, within thirty days, the State has not commenced appropriate enforcement action, the Administrator may act upon the complaint or information to the extent authorized under this Act.

(b) NOTICE.—Whenever the Administrator determines that a State having primary enforcement responsibility for pesticide use violations is not carrying out (or cannot carry out due to the lack of adequate legal authority) such responsibility, the Administrator shall notify the State. Such notice shall specify those aspects of the administration of the State program that are determined to be inadequate. The State shall have ninety days after receipt of the notice to correct any deficiencies. If after that time the Administrator determines that the State program remains inadequate, the Administrator may rescind, in whole or in part, the State’s primary enforcement responsibility for pesticide use violations.

(c) CONSTRUCTION.—Neither section 26 of this Act nor this section shall limit the authority of the Administrator to enforce this Act, where the Administrator determines that emergency conditions exist that require immediate action on the part of the Administrator and the State authority is unwilling or unable adequately to respond to the emergency.

(a) In General.—The Administrator, in coordination with the Secretary of Agriculture, shall identify those pests that must be brought under control. The Administrator shall also coordinate and cooperate with the Secretary of Agriculture's research and implementation programs to develop and improve the safe use and effectiveness of chemical, biological, and alternative methods to combat and control pests that reduce the quality and economical production and distribution of agricultural products to domestic and foreign consumers.

(b) Pest Control Availability.—
   (1) In General.—The Administrator, in cooperation with the Secretary of Agriculture, shall identify—
      (A) available methods of pest control by crop or animal;
      (B) minor pest control problems, both in minor crops and minor or localized problems in major crops; and
      (C) factors limiting the availability of specific pest control methods, such as resistance to control methods and regulatory actions limiting the availability of control methods.
   (2) Report.—The Secretary of Agriculture shall, not later than 180 days after the date of enactment of this subsection [November 28, 1990] and annually thereafter, prepare a report and send the report to the Administrator. The report shall—
      (A) contain the information described in paragraph (1) and the information required by section 1651 of the Food, Agriculture, Conservation, and Trade Act of 1990;
      (B) identify the crucial pest control needs where a shortage of control methods is indicated by the information described in paragraph (1); and
      (C) describe in detail research and extension efforts designed to address the needs identified in subparagraph (B).

(c) Integrated Pest Management.—The Administrator, in cooperation with the Secretary of Agriculture, shall develop approaches to the control of pests based on integrated pest management that respond to the needs of producers, with a special emphasis on minor pests.

(d) Public Health Pests.—The Administrator, in coordination with the Secretary of Agriculture and the Secretary of Health and Human Services, shall identify pests of significant public health importance and, in coordination with the Public Health Service, develop and implement programs to improve and facilitate the safe and necessary use of chemical, biological, and other methods to combat and control such pests of public health importance.

The Administrator shall submit an annual report to Congress before February 16 of each year and the first report shall be due February 15, 1979. The report shall include the total number of applications for conditional registration under sections 3(c)(7)(B) and 3(c)(7)(C) of this Act that were filed during the immediately pre-
ceding fiscal year, and, with respect to those applications approved, the Administrator shall report the Administrator's findings in each case, the conditions imposed and any modification of such conditions in each case, and the quantities produced of such pesticides.


Each State may establish minimum requirements for training of maintenance applicators and service technicians. Such training may include instruction in the safe and effective handling and use of pesticides in accordance with the Environmental Protection Agency approved labeling, and instruction in integrated pest management techniques. The authority of the Administrator with respect to minimum requirements for training of maintenance applicators and service technicians shall be limited to ensuring that each State understands the provisions of this section.


(a) The Administrator shall assure coordination of minor use issues through the establishment of a minor use program within the Office of Pesticide Programs. Such office shall be responsible for coordinating the development of minor use programs and policies and consulting with growers regarding minor use issues and registrations and amendments which are submitted to the Environmental Protection Agency.

(b) The Office of Pesticide Programs shall prepare a public report concerning the progress made on the registration of minor uses, including implementation of the exclusive use as an incentive for registering new minor uses, within 3 years of the passage of the Food Quality Protection Act of 1996.

[Aug. 3, 1996]


(a) In General.—The Secretary of Agriculture (hereinafter in this section referred to as the "Secretary") shall assure the coordination of the responsibilities of the Department of Agriculture related to minor uses of pesticides, including—

(1) carrying out the Inter-Regional Project Number 4 (IR–4) as described in section 2 of Public Law 89–106 (7 U.S.C. 450i(e)) and the national pesticide resistance monitoring program established under section 1651 of the Food, Agriculture, Conservation, and Trade Act of 1990 (7 U.S.C. 5882);

(2) supporting integrated pest management research;

(3) consulting with growers to develop data for minor uses; and

(4) providing assistance for minor use registrations, tolerances, and reregistrations with the Environmental Protection Agency.

(b)(1) Minor Use Pesticide Data.—

(A) Grant Authority.—The Secretary, in consultation with the Administrator, shall establish a program to make grants for the development of data to support minor use pesticide registrations and reregistrations. The amount of any
such grant shall not exceed \( \frac{1}{2} \) of the cost of the project for which the grant is made.

(B) APPLICANTS.—Any person who wants to develop data to support minor use pesticide registrations and reregistrations may apply for a grant under subparagraph (A). Priority shall be given to an applicant for such a grant who does not directly receive funds from the sale of pesticides registered for minor uses.

(C) DATA OWNERSHIP.—Any data that is developed under a grant under subparagraph (A) shall be jointly owned by the Department of Agriculture and the person who received the grant. Such a person shall enter into an agreement with the Secretary under which such person shall share any fee paid to such person under section 3(c)(1)(F).

(2) MINOR USE PESTICIDE DATA REVOLVING FUND.—

(A) ESTABLISHMENT.—There is established in the Treasury of the United States a revolving fund to be known as the Minor Use Pesticide Data Revolving Fund. The Fund shall be available without fiscal year limitation to carry out the authorized purposes of this subsection.

(B) CONTENTS OF THE FUND.—There shall be deposited in the Fund—

(i) such amounts as may be appropriated to support the purposes of this subsection; and

(ii) fees collected by the Secretary for any data developed under a grant under paragraph (1)(A).

(C) AUTHORIZATIONS OF APPROPRIATIONS.—There are authorized to be appropriated for each fiscal year to carry out the purposes of this subsection $10,000,000 to remain available until expended.


(a) DEFINITION OF COSTS.—In this section, the term “costs”, when used with respect to review and decisionmaking pertaining to an application for which registration service fees are paid under this section, means—

(1) costs to the extent that—

(A) officers and employees provide direct support for the review and decisionmaking for covered pesticide applications, associated tolerances, and corresponding risk and benefits information and analyses;

(B) persons and organizations under contract with the Administrator engage in the review of the applications, and corresponding risk and benefits information and assessments; and

(C) advisory committees and other accredited persons or organizations, on the request of the Administrator, engage in the peer review of risk or benefits information associated with covered pesticide applications;

(2) costs of management of information, and the acquisition, maintenance, and repair of computer and telecommunication resources (including software), used to support review of pesticide applications, associated tolerances, and corresponding risk and benefits information and analyses; and
(3) costs of collecting registration service fees under sub-
sections (b) and (c) and reporting, auditing, and accounting
under this section.

(b) FEES.—

(1) IN GENERAL.—Effective beginning on the effective date
of the Pesticide Registration Improvement Act of 2003, the Ad-
ministrator shall assess and collect covered pesticide regis-
tration service fees in accordance with this section.

(2) COVERED APPLICATIONS.—

(A) IN GENERAL.—An application for the registration of
a pesticide covered by this Act that is received by the Ad-
ministrator on or after the effective date of the Pesticide
Registration Improvement Act of 2003 or for any other ac-
tion covered by a table specified in paragraph (3) shall be
subject to a registration service fee under this section.

(B) EXISTING APPLICATIONS.—

(i) IN GENERAL.—Subject to clause (ii), an applica-
tion for the registration of a pesticide that was sub-
mitted to the Administrator before the effective date of
the Pesticide Registration Improvement Act of 2003 and
is pending on that effective date shall be subject
to a service fee under this section if the application is
for the registration of a new active ingredient that is
not listed in the Registration Division 2003 Work Plan
of the Office of Pesticide Programs of the Environ-
mental Protection Agency.

(ii) TOLERANCE OR EXEMPTION FEES.—The amount
of any fee otherwise payable for an application de-
scribed in clause (i) under this section shall be reduced
by the amount of any fees paid to support the related
petition for a pesticide tolerance or exemption under
301 et seq.).

(C) DOCUMENTATION.—An application subject to a reg-
istration service fee under this section shall be submitted
with documentation certifying—

(i) payment of the registration service fee; or

(ii) payment of at least 25 percent of the registra-
tion service fee and a request for a waiver from or re-
duction of the remaining amount of the registration
service fee.

(D) PAYMENT.—The registration service fee required
under this subsection shall be due upon submission of the
application.

(E) APPLICATIONS SUBJECT TO ADDITIONAL FEES.—An
application may be subject to additional fees if—

(i) the applicant identified the incorrect registration
service fee and decision review period;

(ii) after review of a waiver request, the Adminis-
trator denies the waiver request; or

(iii) after review of the application, the Adminis-
trator determines that a different registration service
fee and decision review period apply to the application.
(F) Effect of failure to pay fees.—The Administrator shall reject any application submitted without the required registration service fee.

(G) Non-refundable portion of fees.—
   (i) In general.—The Administrator shall retain 25 percent of the applicable registration service fee.
   (ii) Limitation.—Any waiver, refund, credit or other reduction in the registration service fee shall not exceed 75 percent of the registration service fee.

(H) Collection of unpaid fees.—In any case in which the Administrator does not receive payment of a registration service fee (or applicable portion of the registration service fee) by the date that is 30 days after the fee is due, the fee shall be treated as a claim of the United States Government subject to subchapter II of chapter 37 of title 31, United States Code.

(3) Schedule of covered applications and other actions and their registration service fees.—Subject to paragraph (6), the schedule of registration applications and other covered actions and their corresponding registration service fees shall be as follows:

**TABLE 1. — REGISTRATION DIVISION — NEW ACTIVE INGREDIENTS**

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R010</td>
<td>1</td>
<td>New Active Ingredient, Food use. (2)(3)</td>
<td>24</td>
<td>753,082</td>
</tr>
<tr>
<td>R020</td>
<td>2</td>
<td>New Active Ingredient, Food use; reduced risk. (2)(3)</td>
<td>18</td>
<td>627,568</td>
</tr>
<tr>
<td>R040</td>
<td>3</td>
<td>New Active Ingredient, Food use; Experimental Use Permit application; establish temporary tolerance; submitted before application for registration; credit 45% of fee toward new active ingredient application that follows. (3)</td>
<td>18</td>
<td>462,502</td>
</tr>
</tbody>
</table>
Table 1. — Registration Division — New Active Ingredients—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(1)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R060</td>
<td>4</td>
<td>New Active Ingredient, Non-food use; outdoor. (2)(3)</td>
<td>21</td>
<td>523,205</td>
</tr>
<tr>
<td>R070</td>
<td>5</td>
<td>New Active Ingredient, Non-food use; outdoor; reduced risk. (2)(3)</td>
<td>16</td>
<td>436,004</td>
</tr>
<tr>
<td>R090</td>
<td>6</td>
<td>New Active Ingredient, Non-food use; outdoor; Experimental Use Permit application; submitted before application for registration; credit 45% of fee toward new active ingredient application that follows. (3)</td>
<td>16</td>
<td>323,690</td>
</tr>
<tr>
<td>R110</td>
<td>7</td>
<td>New Active Ingredient, Non-food use; indoor. (2)(3)</td>
<td>20</td>
<td>290,994</td>
</tr>
<tr>
<td>R120</td>
<td>8</td>
<td>New Active Ingredient, Non-food use; indoor; reduced risk. (2)(3)</td>
<td>14</td>
<td>242,495</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)(1)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>--------</td>
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<td>----------------------------------------------------------------------</td>
<td>----------------------------------</td>
<td>-------------------------------</td>
</tr>
<tr>
<td>R121</td>
<td>9</td>
<td>New Active Ingredient, Non-food use; indoor; Experimental Use Permit application; submitted before application for registration; credit 45% of fee toward new active ingredient application that follows. (3)</td>
<td>18</td>
<td>182,327</td>
</tr>
<tr>
<td>R122</td>
<td>10</td>
<td>Enriched isomer(s) of registered mixed-isomer active ingredient. (2)(3)</td>
<td>18</td>
<td>317,128</td>
</tr>
<tr>
<td>R123</td>
<td>11</td>
<td>New Active Ingredient, Seed treatment only; includes agricultural and non-agricultural seeds; residues not expected in raw agricultural commodities. (2)(3)</td>
<td>18</td>
<td>471,861</td>
</tr>
</tbody>
</table>
TABLE 1. — REGISTRATION DIVISION — NEW ACTIVE INGREDIENTS—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
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</thead>
<tbody>
<tr>
<td>R125</td>
<td>12</td>
<td>New Active Ingredient, Seed treatment; Experimental Use Permit application; submitted before application for registration; credit 45% of fee toward new active ingredient application that follows. (3)</td>
<td>16</td>
<td>323,690</td>
</tr>
</tbody>
</table>

(1) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.

(2) All requests for new uses (food and/or nonfood) contained in any application for a new active ingredient or a first food use are covered by the base fee for that new active ingredient or first food use application and retain the same decision time review period as the new active ingredient or first food use application. The application must be received by the agency in one package. The base fee for the category covers a maximum of five new products. Each application for an additional new product registration and new inert approval that is submitted in the new active ingredient application package or first food use application package is subject to the registration service fee for a new product or a new inert approval. All such associated applications that are submitted together will be subject to the new active ingredient or first food use decision review time. In the case of a new active ingredient application, until that new active ingredient is approved, any subsequent application for another new product containing the same active ingredient or an amendment to the proposed labeling will be deemed a new active ingredient application, subject to the registration service fee and decision review time for a new active ingredient. In the case of a first food use application, until that first food use is approved, any subsequent application for an additional new food use or uses will be subject to the registration service fee and decision review time for a first food use. Any information that (a) was neither requested nor required by the Agency, and (b) is submitted by the applicant at the applicant’s initiative to support the application after completion of the technical deficiency screening, and (c) is not itself a covered registration application, must be assessed 25% of the full registration service fee for the new active ingredient or first food use application.
(3) Where the action involves approval of a new or amended label, on or before the end date of the decision review time, the Agency shall provide to the applicant a draft accepted label, including any changes made by the Agency that differ from the applicant-submitted label and relevant supporting data reviewed by the Agency. The applicant will notify the Agency that the applicant either (a) agrees to all of the terms associated with the draft accepted label as amended by the Agency and requests that it be issued as the accepted final Agency-stamped label; or (b) does not agree to one or more of the terms of the draft accepted label as amended by the Agency and requests additional time to resolve the difference(s); or (c) withdraws the application without prejudice for subsequent resubmission, but forfeits the associated registration service fee. For cases described in (b), the applicant shall have up to 30 calendar days to reach agreement with the Agency on the final terms of the Agency-accepted label. If the applicant agrees to all of the terms of the accepted label as in (a), including upon resolution of differences in (b), the Agency shall provide an accepted final Agency-stamped label to the registrant within 2 business days following the registrant’s written or electronic confirmation of agreement to the Agency.

### TABLE 2. — REGISTRATION DIVISION — NEW USES

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R130</td>
<td>13</td>
<td>First food use; indoor; food/food handling. (2) (3)</td>
<td>21</td>
<td>191,444</td>
</tr>
<tr>
<td>R140</td>
<td>14</td>
<td>Additional food use; Indoor; food/food handling. (3) (4)</td>
<td>15</td>
<td>44,672</td>
</tr>
<tr>
<td>R150</td>
<td>15</td>
<td>First food use. (2)(3)</td>
<td>21</td>
<td>317,104</td>
</tr>
<tr>
<td>R155</td>
<td>16 (new)</td>
<td>First food use, Experimental Use Permit application; a.i. registered for non-food outdoor use. (3)(4)</td>
<td>21</td>
<td>264,253</td>
</tr>
<tr>
<td>R160</td>
<td>17</td>
<td>First food use; reduced risk. (2)(3)</td>
<td>16</td>
<td>264,253</td>
</tr>
<tr>
<td>R170</td>
<td>18</td>
<td>Additional food use. (3) (4)</td>
<td>15</td>
<td>79,349</td>
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### Table 2. — Registration Division — New Uses—Continued

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<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(1)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R175</td>
<td>19</td>
<td>Additional food uses covered within a crop group resulting from the conversion of existing approved crop group(s) to one or more revised crop groups. (3)(4)</td>
<td>10</td>
<td>66,124</td>
</tr>
<tr>
<td>R180</td>
<td>20</td>
<td>Additional food use; reduced risk. (3)(4)</td>
<td>10</td>
<td>66,124</td>
</tr>
<tr>
<td>R190</td>
<td>21</td>
<td>Additional food uses; 6 or more submitted in one application. (3)(4)</td>
<td>15</td>
<td>476,090</td>
</tr>
<tr>
<td>R200</td>
<td>22</td>
<td>Additional Food Use; 6 or more submitted in one application; Reduced Risk. (3)(4)</td>
<td>10</td>
<td>396,742</td>
</tr>
<tr>
<td>R210</td>
<td>23</td>
<td>Additional food use; Experimental Use Permit application; establish temporary tolerance; no credit toward new use registration. (3)(4)</td>
<td>12</td>
<td>48,986</td>
</tr>
<tr>
<td>R220</td>
<td>24</td>
<td>Additional food use; Experimental Use Permit application; crop destruct basis; no credit toward new use registration. (3)(4)</td>
<td>6</td>
<td>19,838</td>
</tr>
</tbody>
</table>
### TABLE 2. — REGISTRATION DIVISION — NEW USES—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)&lt;sup&gt;(1)&lt;/sup&gt;</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R230</td>
<td>25</td>
<td>Additional use; non-food; outdoor. (3) (4)</td>
<td>15</td>
<td>31,713</td>
</tr>
<tr>
<td>R240</td>
<td>26</td>
<td>Additional use; non-food; outdoor; reduced risk. (3)(4)</td>
<td>10</td>
<td>26,427</td>
</tr>
<tr>
<td>R250</td>
<td>27</td>
<td>Additional use; non-food; outdoor; Experimental Use Permit application; no credit toward new use registration. (3)(4)</td>
<td>6</td>
<td>19,838</td>
</tr>
<tr>
<td>R251</td>
<td>28</td>
<td>Experimental Use Permit application which requires no changes to the tolerance(s); non-crop destruct basis. (3)</td>
<td>8</td>
<td>19,838</td>
</tr>
<tr>
<td>R260</td>
<td>29</td>
<td>New use; non-food; indoor. (3) (4)</td>
<td>12</td>
<td>15,317</td>
</tr>
<tr>
<td>R270</td>
<td>30</td>
<td>New use; non-food; indoor; reduced risk. (3)(4)</td>
<td>9</td>
<td>12,764</td>
</tr>
<tr>
<td>R271</td>
<td>31</td>
<td>New use; non-food; indoor; Experimental Use Permit application; no credit toward new use registration. (3)(4)</td>
<td>6</td>
<td>9,725</td>
</tr>
</tbody>
</table>
TABLE 2. — REGISTRATION DIVISION — NEW USES—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R273</td>
<td>32</td>
<td>Additional use; seed treatment; limited uptake into Raw Agricultural Commodities; includes crops with established tolerances (e.g., for soil or foliar application); includes food and/or non-food uses. (3)(4)</td>
<td>12</td>
<td>50,445</td>
</tr>
<tr>
<td>R274</td>
<td>33</td>
<td>Additional uses; seed treatment only; 6 or more submitted in one application; limited uptake into raw agricultural commodities; includes crops with established tolerances (e.g., for soil or foliar application); includes food and/or non-food uses. (3)(4)</td>
<td>12</td>
<td>302,663</td>
</tr>
</tbody>
</table>

(1) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.
(2) All requests for new uses (food and/or nonfood) contained in any application for a new active ingredient or a first food use are covered by the base fee for that new active ingredient or first food use application and retain the same decision time review period as the new active ingredient or first food use application. The application must be received by the agency in one package. The base fee for the category covers a maximum of five new products. Each application for an additional new product registration and new inert approval that is submitted in the new active ingredient application package or first food use application package is subject to the registration service fee for a new product or a new inert approval. All such associated applications that are submitted together will be subject to the new active ingredient or first food use decision review time. In the case of a new active ingredient application, until that new active ingredient is approved, any subsequent application for another new product containing the same active ingredient or an amendment to the proposed labeling will be deemed a new active ingredient application, subject to the registration service fee and decision review time for a new active ingredient. In the case of a first food use application, until that first food use is approved, any subsequent application for an additional new food use or uses will be subject to the registration service fee and decision review time for a first food use. Any information that (a) was neither requested nor required by the Agency, and (b) is submitted by the applicant at the applicant’s initiative to support the application after completion of the technical deficiency screening, and (c) is not itself a covered registration application, must be assessed 25% of the full registration service fee for the new active ingredient or first food use application.

(3) Where the action involves approval of a new or amended label, on or before the end date of the decision review time, the Agency shall provide to the applicant a draft accepted label, including any changes made by the Agency that differ from the applicant-submitted label and relevant supporting data reviewed by the Agency. The applicant will notify the Agency that the applicant either (a) agrees to all of the terms associated with the draft accepted label as amended by the Agency and requests that it be issued as the accepted final Agency-stamped label; or (b) does not agree to one or more of the terms of the draft accepted label as amended by the Agency and requests additional time to resolve the difference(s); or (c) withdraws the application without prejudice for subsequent resubmission, but forfeits the associated registration service fee. For cases described in (b), the applicant shall have up to 30 calendar days to reach agreement with the Agency on the final terms of the Agency-accepted label. If the applicant agrees to all of the terms of the accepted label as in (a), including upon resolution of differences in (b), the Agency shall provide an accepted final Agency-stamped label to the registrant within 2 business days following the registrant’s written or electronic confirmation of agreement to the Agency.
(4) Amendment applications to add the new use(s) to registered product labels are covered by the base fee for the new use(s). All items in the covered application must be submitted together in one package. Each application for an additional new product registration and new inert approval(s) that is submitted in the new use application package is subject to the registration service fee for a new product or a new inert approval. However, if a new use application only proposes to register the new use for a new product and there are no amendments in the application, then review of one new product application is covered by the new use fee. All such associated applications that are submitted together will be subject to the new use decision review time. Any application for a new product or an amendment to the proposed labeling (a) submitted subsequent to submission of the new use application and (b) prior to conclusion of its decision review time and (c) containing the same new uses, will be deemed a separate new-use application, subject to a separate registration service fee and new decision review time for a new use. If the new-use application includes non-food (indoor and/or outdoor), and food (outdoor and/or indoor) uses, the appropriate fee is due for each type of new use and the longest decision review time applies to all of the new uses requested in the application. Any information that (a) was neither requested nor required by the Agency, and (b) is submitted by the applicant at the applicant’s initiative to support the application after completion of the technical deficiency screen, and (c) is not itself a covered registration application, must be assessed 25% of the full registration service fee for the new use application.

### TABLE 3. — REGISTRATION DIVISION — IMPORT AND OTHER TOLERANCES

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R280</td>
<td>34</td>
<td>Establish import tolerance; new active ingredient or first food use. (2)</td>
<td>21</td>
<td>319,072</td>
</tr>
<tr>
<td>R290</td>
<td>35</td>
<td>Establish Import tolerance; Additional new food use.</td>
<td>15</td>
<td>63,816</td>
</tr>
<tr>
<td>R291</td>
<td>36</td>
<td>Establish import tolerances; additional food uses; 6 or more crops submitted in one petition.</td>
<td>15</td>
<td>382,886</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)(1)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>---------</td>
<td>------------</td>
<td>--------</td>
<td>--------------------------------</td>
<td>-----------------------------</td>
</tr>
<tr>
<td>R292</td>
<td>37</td>
<td>Amend an established tolerance (e.g., decrease or increase) and/or harmonize established tolerances with Codex MRLs; domestic or import; applicant-initiated.</td>
<td>11</td>
<td>45,341</td>
</tr>
<tr>
<td>R293</td>
<td>38</td>
<td>Establish tolerance(s) for inadvertent residues in one crop; applicant-initiated.</td>
<td>12</td>
<td>53,483</td>
</tr>
<tr>
<td>R294</td>
<td>39</td>
<td>Establish tolerances for inadvertent residues; 6 or more crops submitted in one application; applicant-initiated.</td>
<td>12</td>
<td>320,894</td>
</tr>
<tr>
<td>R295</td>
<td>40</td>
<td>Establish tolerance(s) for residues in one rotational crop in response to a specific rotational crop application; submission of corresponding label amendments which specify the necessary plant-back restrictions; applicant-initiated.</td>
<td>15</td>
<td>66,124</td>
</tr>
</tbody>
</table>
### TABLE 3. — REGISTRATION DIVISION — IMPORT AND OTHER TOLERANCES—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)&lt;sup&gt;(1)&lt;/sup&gt;</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R296</td>
<td>41</td>
<td>Establish tolerances for residues in rotational crops in response to a specific rotational crop petition; 6 or more crops submitted in one application; submission of corresponding label amendments which specify the necessary plant-back restrictions; applicant-initiated. (3) (4)</td>
<td>15</td>
<td>396,742</td>
</tr>
<tr>
<td>R297</td>
<td>42</td>
<td>Amend 6 or more established tolerances (e.g., decrease or increase) in one petition; domestic or import; applicant-initiated. (3) (4)</td>
<td>11</td>
<td>272,037</td>
</tr>
<tr>
<td>R298</td>
<td>43</td>
<td>Amend an established tolerance (e.g., decrease or increase); domestic or import; submission of corresponding amended labels (requiring science review). (3) (4)</td>
<td>13</td>
<td>58,565</td>
</tr>
</tbody>
</table>
TABLE 3. — REGISTRATION DIVISION — IMPORT AND OTHER TOLERANCES—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(^{(1)})</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R299</td>
<td>44</td>
<td>Amend 6 or more established tolerances (e.g., decrease or increase); domestic or import; submission of corresponding amended labels (requiring science review). (3) (4)</td>
<td>13</td>
<td>285,261</td>
</tr>
</tbody>
</table>

(1) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.

(2) All requests for new uses (food and/or nonfood) contained in any application for a new active ingredient or a first food use are covered by the base fee for that new active ingredient or first food use application and retain the same decision time review period as the new active ingredient or first food use application. The application must be received by the agency in one package. The base fee for the category covers a maximum of five new products. Each application for an additional new product registration and new inert approval that is submitted in the new active ingredient application package or first food use application package is subject to the registration service fee for a new product or a new inert approval. All such associated applications that are submitted together will be subject to the new active ingredient or first food use decision review time. In the case of a new active ingredient application, until that new active ingredient is approved, any subsequent application for another new product containing the same active ingredient or an amendment to the proposed labeling will be deemed a new active ingredient application, subject to the registration service fee and decision review time for a new active ingredient. In the case of a first food use application, until that first food use is approved, any subsequent application for an additional new food use or uses will be subject to the registration service fee and decision review time for a first food use. Any information that (a) was neither requested nor required by the Agency, and (b) is submitted by the applicant at the applicant’s initiative to support the application after completion of the technical deficiency screening, and (c) is not itself a covered registration application, must be assessed 25% of the full registration service fee for the new active ingredient or first food use application.
(3) Where the action involves approval of a new or amended label, on or before the end date of the decision review time, the Agency shall provide to the applicant a draft accepted label, including any changes made by the Agency that differ from the applicant-submitted label and relevant supporting data reviewed by the Agency. The applicant will notify the Agency that the applicant either (a) agrees to all of the terms associated with the draft accepted label as amended by the Agency and requests that it be issued as the accepted final Agency-stamped label; or (b) does not agree to one or more of the terms of the draft accepted label as amended by the Agency and requests additional time to resolve the difference(s); or (c) withdraws the application without prejudice for subsequent resubmission, but forfeits the associated registration service fee. For cases described in (b), the applicant shall have up to 30 calendar days to reach agreement with the Agency on the final terms of the Agency-accepted label. If the applicant agrees to all of the terms of the accepted label as in (a), including upon resolution of differences in (b), the Agency shall provide an accepted final Agency-stamped label to the registrant within 2 business days following the registrant’s written or electronic confirmation of agreement to the Agency.

(4) Amendment applications to add the revised use pattern(s) to registered product labels are covered by the base fee for the category. All items in the covered application must be submitted together in one package. Each application for an additional new product registration and new inert approval(s) that is submitted in the amendment application package is subject to the registration service fee for a new product or a new inert approval. However, if an amendment application only proposes to register the amendment for a new product and there are no amendments in the application, then review of one new product application is covered by the base fee. All such associated applications that are submitted together will be subject to the category decision review time.
### TABLE 4. — REGISTRATION DIVISION — NEW PRODUCTS

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(1)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R300</td>
<td>45</td>
<td>New product; or similar combination product (already registered) to an identical or substantially similar in composition and use to a registered product; registered source of active ingredient; no data review on acute toxicity, efficacy or CRP – only product chemistry data; cite-all data citation, or selective data citation where applicant owns all required data, or applicant submits specific authorization letter from data owner. Category also includes 100% repackage of registered end-use or manufacturing-use product that requires no data submission nor data matrix. (2)(3)</td>
<td>4</td>
<td>1,582</td>
</tr>
</tbody>
</table>
TABLE 4. — REGISTRATION DIVISION — NEW PRODUCTS— Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(1)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R301</td>
<td>46</td>
<td>New product; or similar combination product (already registered) to an identical or substantially similar in composition and use to a registered product; registered source of active ingredient; selective data citation only for data on product chemistry and/or acute toxicity and/or public health pest efficacy (identical data citation and claims to cited product(s)), where applicant does not own all required data and does not have a specific authorization letter from data owner. (2)(3)</td>
<td>4</td>
<td>1,897</td>
</tr>
</tbody>
</table>
TABLE 4. — REGISTRATION DIVISION — NEW PRODUCTS—
Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R310</td>
<td>47</td>
<td>New end-use or manufacturing-use product with registered source(s) of active ingredient(s); includes products containing two or more registered active ingredients previously combined in other registered products; excludes products requiring or citing an animal safety study; requires review of data package within RD only; includes data and/or waivers of data for only: • product chemistry and/or • acute toxicity and/or • child resistant packaging and/or • pest(s) requiring efficacy (4) - for up to 3 target pests. (2)(3)</td>
<td>7</td>
<td>7,301</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)(^{(1)})</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>---------</td>
<td>------------</td>
<td>------------------------------------------------------------------------</td>
<td>------------------------------------------</td>
<td>-----------------------------</td>
</tr>
<tr>
<td>R314</td>
<td>48</td>
<td>New end use product containing up to three registered active ingredients never before registered as this combination in a formulated product; new product label is identical or substantially similar to the labels of currently registered products which separately contain the respective component active ingredients; excludes products requiring or citing an animal safety study; requires review of data package within RD only; includes data and/or waivers of data for only: * product chemistry and/or * acute toxicity and/or * child resistant packaging and/or * pest(s) requiring efficacy (4) - for up to 3 target pests. (^{(2)})((^{(3)}))</td>
<td>8</td>
<td>8,626</td>
</tr>
</tbody>
</table>
### Table 4. — Registration Division — New Products— Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(^{(1)})</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R319</td>
<td>49</td>
<td>New end use product containing up to three registered active ingredients never before registered as this combination in a formulated product; new product label is identical or substantially similar to the labels of currently registered products which separately contain the respective component active ingredients; excludes products requiring or citing an animal safety study; requires review of data package within RD only; includes data and/or waivers of data for only: • product chemistry and/or • acute toxicity and/or • child resistant packaging and/or • pest(s) requiring efficacy (4) - for 4 to 7 target pests. (2)(3)</td>
<td>10</td>
<td>12,626</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>---------</td>
<td>------------</td>
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<td>-----------------------------</td>
<td>-----------------------------</td>
</tr>
<tr>
<td>R318</td>
<td>50 (new)</td>
<td>New end use product containing four or more registered active ingredients never before registered as this combination in a formulated product; new product label is identical or substantially similar to the labels of currently registered products which separately contain the respective component active ingredients; excludes products requiring or citing an animal safety study; requires review of data package within RD only; includes data and/or waivers of data for only: • product chemistry and/or • acute toxicity and/or • child resistant packaging and/or • pest(s) requiring efficacy (4) – for up to 3 target pests. (2)(3)</td>
<td>9</td>
<td>13,252</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)$^{(1)}$</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>--------</td>
<td>------------</td>
<td>--------</td>
<td>--------------------------------------</td>
<td>------------------------------</td>
</tr>
</tbody>
</table>
| R321   | 51 (new)   | New end use product containing four or more registered active ingredients never before registered as this combination in a formulated product; new product label is identical or substantially similar to the labels of currently registered products which separately contain the respective component active ingredients; excludes products requiring or citing an animal safety study; requires review of data package within RD only; includes data and/or waivers of data for only:  
  - product chemistry and/or  
  - acute toxicity and/or  
  - child resistant packaging and/or  
  - pest(s) requiring efficacy (4) - for 4 to 7 target pests. (2)(3) | 11 | 17,252 |
### TABLE 4. — REGISTRATION DIVISION — NEW PRODUCTS— Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R315</td>
<td>52</td>
<td>New end-use, on-animal product, registered source of active ingredient(s), with the submission of data and/or waivers for only: • animal safety and • pest(s) requiring efficacy (4) and/or • product chemistry and/or • acute toxicity and/or • child resistant packaging. (2) (3)</td>
<td>9</td>
<td>9,820</td>
</tr>
</tbody>
</table>
TABLE 4. — REGISTRATION DIVISION — NEW PRODUCTS—
Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(^{(1)})</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
</table>
| R316    | 53 (new)   | New end-use or manufacturing product with registered source(s) of active ingredient(s) including products containing two or more registered active ingredients previously combined in other registered products; excludes products requiring or citing an animal safety study; and requires review of data and/or waivers for only:
  • product chemistry and/or
  • acute toxicity and/or
  • child resistant packaging and/or
  • pest(s) requiring efficacy (4) - for greater than 3 and up to 7 target pests. (2)(3) | 9                                        | 11,301                                   |
<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(1)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R317</td>
<td>54 (new)</td>
<td>New end-use or manufacturing product with registered source(s) of active ingredient(s) including products containing 2 or more registered active ingredients previously combined in other registered products; excludes products requiring or citing an animal safety study; and requires review of data and/or waivers for only: • product chemistry and/or • acute toxicity and/or • child resistant packaging and/or • pest(s) requiring efficacy (4) - for greater than 7 target pests. (2)(3)</td>
<td>10</td>
<td>15,301</td>
</tr>
<tr>
<td>R320</td>
<td>55</td>
<td>New product; new physical form; requires data review in science divisions. (2)(3)</td>
<td>12</td>
<td>13,226</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>---------</td>
<td>------------</td>
<td>---------------------------------------------------------------------------------------------</td>
<td>-------------------------------</td>
<td>------------------------------</td>
</tr>
<tr>
<td>R331</td>
<td>56</td>
<td>New product; repack of identical registered end-use product as a manufacturing-use product, or identical registered manufacturing-use product as an end-use product; same registered uses only. (2)(3)</td>
<td>3</td>
<td>2,530</td>
</tr>
<tr>
<td>R332</td>
<td>57</td>
<td>New manufacturing-use product; registered active ingredient; unregistered source of active ingredient; submission of completely new generic data package; registered uses only; requires review in RD and science divisions. (2)(3)</td>
<td>24</td>
<td>283,215</td>
</tr>
</tbody>
</table>
Table 4. — Registration Division — New Products—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(1)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R333</td>
<td>58</td>
<td>New product; MUP or End use product with unregistered source of active ingredient; requires science data review; new physical form; etc. Cite-all or selective data citation where applicant owns all required data. (2)(3)</td>
<td>10</td>
<td>19,838</td>
</tr>
<tr>
<td>R334</td>
<td>59</td>
<td>New product; MUP or End use product with unregistered source of the active ingredient; requires science data review; new physical form; etc. Selective data citation. (2)(3)</td>
<td>11</td>
<td>23,100</td>
</tr>
</tbody>
</table>

(1) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.

(2) An application for a new end-use product using a source of active ingredient that (a) is not yet registered but (b) has an application pending with the Agency for review, will be considered an application for a new product with an unregistered source of active ingredient.
(3) Where the action involves approval of a new or amended label, on or before the end date of the decision review time, the Agency shall provide to the applicant a draft accepted label, including any changes made by the Agency that differ from the applicant-submitted label and relevant supporting data reviewed by the Agency. The applicant will notify the Agency that the applicant either (a) agrees to all of the terms associated with the draft accepted label as amended by the Agency and requests that it be issued as the accepted final Agency-stamped label; or (b) does not agree to one or more of the terms of the draft accepted label as amended by the Agency and requests additional time to resolve the difference(s); or (c) withdraws the application without prejudice for subsequent resubmission, but forfeits the associated registration service fee. For cases described in (b), the applicant shall have up to 30 calendar days to reach agreement with the Agency on the final terms of the Agency-accepted label. If the applicant agrees to all of the terms of the accepted label as in (a), including upon resolution of differences in (b), the Agency shall provide an accepted final Agency-stamped label to the registrant within 2 business days following the registrant’s written or electronic confirmation of agreement to the Agency.

(4) For the purposes of classifying proposed registration actions into PRIA categories, pest(s) requiring efficacy are: public health pests listed in PR Notice 2002–1, livestock pests (e.g. Horn flies, Stable flies), wood-destroying pests (e.g. termites, carpenter ants, wood-boring beetles) and certain invasive species (e.g. Asian Longhorned beetle, Emerald Ashborer). This list may be updated/refined as invasive pest needs arise. To determine the number of pests for the PRIA categories, pests have been placed into groups (general; e.g., cockroaches) and pest specific (specifically a test species). If seeking a label claim against a pest group (general), use the group listing below and each group will count as 1. The general pests groups are: mites, dust mites, chiggers, ticks, hard ticks, soft ticks, cattle ticks, scorpions, spiders, centipedes, lice, fleas, cockroaches, keds, bot flies, screwworms, filth flies, blow flies, house flies, flesh flies, mosquitoes, biting flies, horse flies, stable flies, deer flies, sand flies, biting midges, black flies, true bugs, bed bugs, stinging bees, wasps, yellow jackets, hornets, ants (excluding carpenter ants), fire and harvester ants, wood destroying beetles, carpenter ants, termites, subterranean termites, dry wood termites, arboreal termites, damp wood termites and invasive species. If seeking a claim against a specific pest without a general claim then each specific pest will count as 1.
TABLE 5. — REGISTRATION DIVISION — AMENDMENTS

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R340</td>
<td>60</td>
<td>Amendment requiring data review within RD (e.g., changes to precautionary label statements); includes adding/modifying pest(s) claims for up to 2 target pests, excludes products requiring or citing an animal safety study. (2)(3)(4)</td>
<td>4</td>
<td>4,988</td>
</tr>
<tr>
<td>R341</td>
<td>61</td>
<td>Amendment requiring data review within RD (e.g., changes to precautionary label statements); includes adding/modifying pest(s) claims for greater than 2 target pests, excludes products requiring or citing an animal safety study. (2)(3)(4)</td>
<td>6</td>
<td>5,988</td>
</tr>
<tr>
<td>R345</td>
<td>62</td>
<td>Amending on-animal products previously registered, with the submission of data and/or waivers for only: animal safety and pest(s) requiring efficacy (4) and/or product chemistry and/or acute toxicity and/or child resistant packaging. (2)(3)</td>
<td>7</td>
<td>8,820</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)^(1)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>--------</td>
<td>------------</td>
<td>------------------------------------------------------------------------</td>
<td>-----------------------------------</td>
<td>------------------------------</td>
</tr>
<tr>
<td>R350</td>
<td>63</td>
<td>Amendment requiring data review in science divisions (e.g., changes to REI, or PPE, or PHI, or use rate, or number of applications; or add aerial application; or modify GW/SW advisory statement). (2)(3)</td>
<td>9</td>
<td>13,226</td>
</tr>
<tr>
<td>R351</td>
<td>64</td>
<td>Amendment adding a new unregistered source of active ingredient. (2)(3)</td>
<td>8</td>
<td>13,226</td>
</tr>
<tr>
<td>R352</td>
<td>65</td>
<td>Amendment adding already approved uses; selective method of support; does not apply if the applicant owns all cited data. (2) (3)</td>
<td>8</td>
<td>13,226</td>
</tr>
<tr>
<td>R371</td>
<td>66</td>
<td>Amendment to Experimental Use Permit; (does not include extending a permit’s time period). (3)</td>
<td>6</td>
<td>10,090</td>
</tr>
</tbody>
</table>

^(1) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.

^(2) (a) EPA-initiated amendments shall not be charged registration service fees. (b) Registrant-initiated fast-track amendments are to be completed within the timelines specified in FIFRA Section 3(c)(3)(B) and are not subject to registration service fees. (c) Registrant-initiated fast-track amendments handled by the Antimicrobials Division are to be completed within the timelines specified in FIFRA Section 3(h) and are not subject to registration service fees. (d) Registrant initiated amendments submitted by notification under PR Notices, such as PR Notice 98–10, continue under PR Notice timelines and are not subject to registration service fees. (e) Submissions with data and requiring data review are subject to registration service fees.
(3) Where the action involves approval of a new or amended label, on or before the end date of the decision review time, the Agency shall provide to the applicant a draft accepted label, including any changes made by the Agency that differ from the applicant-submitted label and relevant supporting data reviewed by the Agency. The applicant will notify the Agency that the applicant either (a) agrees to all of the terms associated with the draft accepted label as amended by the Agency and requests that it be issued as the accepted final Agency-stamped label; or (b) does not agree to one or more of the terms of the draft accepted label as amended by the Agency and requests additional time to resolve the difference(s); or (c) withdraws the application without prejudice for subsequent resubmission, but forfeits the associated registration service fee. For cases described in (b), the applicant shall have up to 30 calendar days to reach agreement with the Agency on the final terms of the Agency-accepted label. If the applicant agrees to all of the terms of the accepted label as in (a), including upon resolution of differences in (b), the Agency shall provide an accepted final Agency-stamped label to the registrant within 2 business days following the registrant’s written or electronic confirmation of agreement to the Agency.

(4) For the purposes of classifying proposed registration actions into PRIA categories, “pest(s) requiring efficacy” are: public health pests listed in PR Notice 2002–1, livestock pests (e.g. Horn flies, Stable flies), wood-destroying pests (e.g. termites, carpenter ants, wood-boring beetles) and certain invasive species (e.g. Asian Longhorned beetle, Emerald Ashborer). This list may be updated/refined as invasive pest needs arise. To determine the number of pests for the PRIA categories, pests have been placed into groups (general; e.g., cockroaches) and pest specific (specifically a test species). If seeking a label claim against a pest group (general), use the group listing below and each group will count as 1. The general pests groups are: mites, dust mites, chiggers, ticks, hard ticks, soft ticks, cattle ticks, scorpions, spiders, centipedes, lice, fleas, cockroaches, keds, bot flies, screwworms, filth flies, blow flies, house flies, flesh flies, mosquitoes, biting flies, horse flies, stable flies, deer flies, sand flies, biting midges, black flies, true bugs, bed bugs, stinging bees, wasps, yellow jackets, hornets, ants (excluding carpenter ants), fire and harvester ants, wood destroying beetles, carpenter ants, termites, subterranean termites, dry wood termites, arboreal termites, damp wood termites and invasive species. If seeking a claim against a specific pest without a general claim then each specific pest will count as 1.

### TABLE 6. — REGISTRATION DIVISION — OTHER ACTIONS

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R124</td>
<td>67</td>
<td>Conditional Ruling on Pre-application Study Waivers; applicant-initiated.</td>
<td>6</td>
<td>2,530</td>
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</table>
### TABLE 6. — REGISTRATION DIVISION — OTHER ACTIONS— Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(^{(1)})</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R272</td>
<td>68</td>
<td>Review of Study Protocol applicant-initiated; excludes DART, pre-registra-tion conference, Rapid Response review, DNT protocol review, protocol needing HSRB review.</td>
<td>3</td>
<td>2,530</td>
</tr>
<tr>
<td>R275</td>
<td>69</td>
<td>Rebuttal of agency reviewed protocol, applicant initiated.</td>
<td>3</td>
<td>2,530</td>
</tr>
<tr>
<td>R370</td>
<td>70</td>
<td>Cancer reassessment; applicant-initiated.</td>
<td>18</td>
<td>198,250</td>
</tr>
</tbody>
</table>

\(^{(1)}\) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.

### TABLE 7. — ANTIMICROBIALS DIVISION — NEW ACTIVE INGREDIENTS

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(^{(1)})</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A380</td>
<td>71</td>
<td>New Active Ingredient; Indirect Food use; establish tolerance or tolerance exemption if required. (2)(3)</td>
<td>24</td>
<td>137,841</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>---------</td>
<td>------------</td>
<td>------------------------------------------------------------------------</td>
<td>-------------------------------</td>
<td>------------------------------</td>
</tr>
<tr>
<td>A390</td>
<td>72</td>
<td>New Active Ingredient; Direct Food use; establish tolerance or tolerance exemption if required. (2)(3)</td>
<td>24</td>
<td>229,733</td>
</tr>
<tr>
<td>A410</td>
<td>73</td>
<td>New Active Ingredient Non-food use.</td>
<td>21</td>
<td>229,733</td>
</tr>
<tr>
<td>A431</td>
<td>74</td>
<td>New Active Ingredient, Non-food use; low-risk. (2)(3)</td>
<td>12</td>
<td>80,225</td>
</tr>
</tbody>
</table>

(1) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.

(2) All requests for new uses (food and/or nonfood) contained in any application for a new active ingredient or a first food use are covered by the base fee for that new active ingredient or first food use application and retain the same decision time review period as the new active ingredient or first food use application. The application must be received by the agency in one package. The base fee for the category covers a maximum of five new products. Each application for an additional new product registration and new inert approval that is submitted in the new active ingredient application package or first food use application package is subject to the registration service fee for a new product or a new inert approval. All such associated applications that are submitted together will be subject to the new active ingredient or first food use decision review time. In the case of a new active ingredient application, until that new active ingredient is approved, any subsequent application for another new product containing the same active ingredient or an amendment to the proposed labeling will be deemed a new active ingredient application, subject to the registration service fee and decision review time for a new active ingredient. In the case of a first food use application, until that first food use is approved, any subsequent application for an additional new food use or uses will be subject to the registration service fee and decision review time for a first food use. Any information that (a) was neither requested nor required by the Agency, and (b) is submitted by the applicant at the applicant’s initiative to support the application after completion of the technical deficiency screening, and (c) is not itself a covered registration application, must be assessed 25% of the full registration service fee for the new active ingredient or first food use application.
(3) Where the action involves approval of a new or amended label, on or before the end date of the decision review time, the Agency shall provide to the applicant a draft accepted label, including any changes made by the Agency that differ from the applicant-submitted label and relevant supporting data reviewed by the Agency. The applicant will notify the Agency that the applicant either (a) agrees to all of the terms associated with the draft accepted label as amended by the Agency and requests that it be issued as the accepted final Agency-stamped label; or (b) does not agree to one or more of the terms of the draft accepted label as amended by the Agency and requests additional time to resolve the difference(s); or (c) withdraws the application without prejudice for subsequent resubmission, but forfeits the associated registration service fee. For cases described in (b), the applicant shall have up to 30 calendar days to reach agreement with the Agency on the final terms of the Agency-accepted label. If the applicant agrees to all of the terms of the accepted label as in (a), including upon resolution of differences in (b), the Agency shall provide an accepted final Agency-stamped label to the registrant within 2 business days following the registrant’s written or electronic confirmation of agreement to the Agency.

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(1)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A440</td>
<td>75</td>
<td>New Use, Indirect Food Use, establish tolerance or tolerance exemption. (2)(3)(4)</td>
<td>21</td>
<td>31,910</td>
</tr>
<tr>
<td>A441</td>
<td>76</td>
<td>Additional Indirect food uses; establish tolerances or tolerance exemptions if required; 6 or more submitted in one application. (3)(4)(5)</td>
<td>21</td>
<td>114,870</td>
</tr>
<tr>
<td>A450</td>
<td>77</td>
<td>New use, Direct food use, establish tolerance or tolerance exemption. (2)(3)(4)</td>
<td>21</td>
<td>95,724</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>--------</td>
<td>------------</td>
<td>--------</td>
<td>-------------------------------</td>
<td>------------------------------</td>
</tr>
<tr>
<td>A451</td>
<td>78</td>
<td>Additional Direct food uses; establish tolerances or tolerance exemptions if required; 6 or more submitted in one application. (3)(4)(5)</td>
<td>21</td>
<td>182,335</td>
</tr>
<tr>
<td>A500</td>
<td>79</td>
<td>New use, non-food. (4)(5)</td>
<td>12</td>
<td>31,910</td>
</tr>
<tr>
<td>A501</td>
<td>80</td>
<td>New use, non-food; 6 or more submitted in one application. (4)(5)</td>
<td>15</td>
<td>76,583</td>
</tr>
</tbody>
</table>

(1) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.

(2) All requests for new uses (food and/or nonfood) contained in any application for a new active ingredient or a first food use are covered by the base fee for that new active ingredient or first food use application and retain the same decision time review period as the new active ingredient or first food use application. The application must be received by the agency in one package. The base fee for the category covers a maximum of five new products. Each application for an additional new product registration and new inert approval that is submitted in the new active ingredient application package or first food use application package is subject to the registration service fee for a new product or a new inert approval. All such associated applications that are submitted together will be subject to the new active ingredient or first food use decision review time. In the case of a new active ingredient application, until that new active ingredient is approved, any subsequent application for another new product containing the same active ingredient or an amendment to the proposed labeling will be deemed a new active ingredient application, subject to the registration service fee and decision review time for a new active ingredient. In the case of a first food use application, until that first food use is approved, any subsequent application for an additional new food use or uses will be subject to the registration service fee and decision review time for a first food use. Any information that (a) was neither requested nor required by the Agency, and (b) is submitted by the applicant at the applicant’s initiative to support the application after completion of the technical deficiency screening, and (c) is not itself a covered registration application, must be assessed 25% of the full registration service fee for the new active ingredient or first food use application.
(3) If EPA data rules are amended to newly require clearance under section 408 of the FFDCA for an ingredient of an antimicrobial product where such ingredient was not previously subject to such a clearance, then review of the data for such clearance of such product is not subject to a registration service fee for the tolerance action for two years from the effective date of the rule.

(4) Where the action involves approval of a new or amended label, on or before the end date of the decision review time, the Agency shall provide to the applicant a draft accepted label, including any changes made by the Agency that differ from the applicant-submitted label and relevant supporting data reviewed by the Agency. The applicant will notify the Agency that the applicant either (a) agrees to all of the terms associated with the draft accepted label as amended by the Agency and requests that it be issued as the accepted final Agency-stamped label; or (b) does not agree to one or more of the terms of the draft accepted label as amended by the Agency and requests additional time to resolve the difference(s); or (c) withdraws the application without prejudice for subsequent resubmission, but forfeits the associated registration service fee. For cases described in (b), the applicant shall have up to 30 calendar days to reach agreement with the Agency on the final terms of the Agency-accepted label. If the applicant agrees to all of the terms of the accepted label as in (a), including upon resolution of differences in (b), the Agency shall provide an accepted final Agency-stamped label to the registrant within 2 business days following the registrant’s written or electronic confirmation of agreement to the Agency.

(5) Amendment applications to add the new use(s) to registered product labels are covered by the base fee for the new use(s). All items in the covered application must be submitted together in one package. Each application for an additional new product registration and new inert approval(s) that is submitted in the new use application package is subject to the registration service fee for a new product or a new inert approval. However, if a new use application only proposes to register the new use for a new product and there are no amendments in the application, then review of one new product application is covered by the new use fee. All such associated applications that are submitted together will be subject to the new use decision review time. Any application for a new product or an amendment to the proposed labeling (a) submitted subsequent to submission of the new use application and (b) prior to conclusion of its decision review time and (c) containing the same new uses, will be deemed a separate new-use application, subject to a separate registration service fee and new decision review time for a new use. If the new-use application includes non-food (indoor and/or outdoor), and food (outdoor and/or indoor) uses, the appropriate fee is due for each type of new use and the longest decision review time applies to all of the new uses requested in the application. Any information that (a) was neither requested nor required by the Agency, and (b) is submitted by the applicant at the applicant’s initiative to support the application after completion of the technical deficiency screen, and (c) is not itself a covered registration application, must be assessed 25% of the full registration service fee for the new use application.
### TABLE 9. — ANTIMICROBIALS DIVISION — NEW PRODUCTS AND AMENDMENTS

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(1)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A530</td>
<td>81</td>
<td>New product, identical or substantially similar in composition and use to a registered product; no data review or only product chemistry data; cite all data citation or selective data citation where applicant owns all required data; or applicant submits specific authorization letter from data owner. Category also includes 100% re-package of registered end-use or manufacturing use product that requires no data submission nor data matrix. (2)(3)</td>
<td>4</td>
<td>1,278</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>--------</td>
<td>------------</td>
<td>--------</td>
<td>------------------------------</td>
<td>-------------------------------</td>
</tr>
<tr>
<td>A531</td>
<td>82</td>
<td>New product; identical or substantially similar in composition and use to a registered product; registered source of active ingredient; selective data citation only for data on product chemistry and/or acute toxicity and/or public health pest efficacy, where applicant does not own all required data and does not have a specific authorization letter from data owner. (2)(3)</td>
<td>4</td>
<td>1,824</td>
</tr>
</tbody>
</table>

TABLE 9. — ANTIMICROBIALS DIVISION — NEW PRODUCTS AND AMENDMENTS—Continued
TABLE 9. — ANTIMICROBIALS DIVISION — NEW PRODUCTS AND AMENDMENTS—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A532</td>
<td>83</td>
<td>New product; identical or substantially similar in composition and use to a registered product; registered active ingredient; unregistered source of active ingredient; cite-all data citation except for product chemistry; product chemistry data submitted. (2)(3)</td>
<td>5</td>
<td>5,107</td>
</tr>
<tr>
<td>A540</td>
<td>84</td>
<td>New end use product; FIFRA § 2(mm) uses only; up to 25 public health organisms. (2)(3)(5)(6)</td>
<td>5</td>
<td>5,107</td>
</tr>
<tr>
<td>A541</td>
<td>85 (new)</td>
<td>New end use product; FIFRA § 2(mm) uses only; 26–50 public health organisms. (2)(3)(5)(6)</td>
<td>7</td>
<td>8,500</td>
</tr>
<tr>
<td>A542</td>
<td>86 (new)</td>
<td>New end use product; FIFRA § 2(mm) uses only; ≥ 51 public health organisms. (2)(3)(5)</td>
<td>10</td>
<td>15,000</td>
</tr>
</tbody>
</table>
### Table 9. — Antimicrobials Division — New Products and Amendments—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(^{(1)})</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A550</td>
<td>87</td>
<td>New end-use product; uses other than FIFRA § 2(mm); non-FQPA product. (^{(2)(3)(5)})</td>
<td>9</td>
<td>13,226</td>
</tr>
<tr>
<td>A560</td>
<td>88</td>
<td>New manufacturing use product; registered active ingredient; selective data citation. (^{(2)(3)})</td>
<td>6</td>
<td>12,596</td>
</tr>
<tr>
<td>A565</td>
<td>89 (new)</td>
<td>New manufacturing-use product; registered active ingredient; unregistered source of active ingredient; submission of new generic data package; registered uses only; requires science review. (^{(2)(3)})</td>
<td>12</td>
<td>18,234</td>
</tr>
<tr>
<td>A570</td>
<td>90</td>
<td>Label amendment requiring data review; up to 25 public health organisms. (^{(3)(4)(5)(6)})</td>
<td>4</td>
<td>3,831</td>
</tr>
<tr>
<td>A573</td>
<td>91 (new)</td>
<td>Label amendment requiring data review; 26–50 public health organisms. (^{(2)(3)(5)(7)})</td>
<td>6</td>
<td>6,350</td>
</tr>
</tbody>
</table>
TABLE 9. — ANTIMICROBIALS DIVISION — NEW PRODUCTS AND AMENDMENTS—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)$^{(1)}$</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A574</td>
<td>92 (new)</td>
<td>Label amendment requiring data review; ≥ 51 public health organisms. (2)(3)(5)(7)</td>
<td>9</td>
<td>11,000</td>
</tr>
<tr>
<td>A572</td>
<td>93</td>
<td>New Product or amendment requiring data review for risk assessment by Science Branch (e.g., changes to REI, or PPE, or use rate). (2)(3)(4)</td>
<td>9</td>
<td>13,226</td>
</tr>
</tbody>
</table>

(1) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.

(2) An application for a new end-use product using a source of active ingredient that (a) is not yet registered but (b) has an application pending with the Agency for review, will be considered an application for a new product with an unregistered source of active ingredient.

(3) Where the action involves approval of a new or amended label, on or before the end date of the decision review time, the Agency shall provide to the applicant a draft accepted label, including any changes made by the Agency that differ from the applicant-submitted label and relevant supporting data reviewed by the Agency. The applicant will notify the Agency that the applicant either (a) agrees to all of the terms associated with the draft accepted label as amended by the Agency and requests that it be issued as the accepted final Agency-stamped label; or (b) does not agree to one or more of the terms of the draft accepted label as amended by the Agency and requests additional time to resolve the difference(s); or (c) withdraws the application without prejudice for subsequent resubmission, but forfeits the associated registration service fee. For cases described in (b), the applicant shall have up to 30 calendar days to reach agreement with the Agency on the final terms of the Agency-accepted label. If the applicant agrees to all of the terms of the accepted label as in (a), including upon resolution of differences in (b), the Agency shall provide an accepted final Agency-stamped label to the registrant within 2 business days following the registrant’s written or electronic confirmation of agreement to the Agency.
(4)(a) EPA-initiated amendments shall not be charged registration service fees. (b) Registrant-initiated fast-track amendments are to be completed within the timelines specified in FIFRA Section 3(c)(3)(B) and are not subject to registration service fees. (c) Registrant-initiated fast-track amendments handled by the Antimicrobials Division are to be completed within the timelines specified in FIFRA Section 3(h) and are not subject to registration service fees. (d) Registrant initiated amendments submitted by notification under PR Notices, such as PR Notice 98–10, continue under PR Notice timelines and are not subject to registration service fees. (e) Submissions with data and requiring data review are subject to registration service fees.

(5) The applicant must identify the substantially similar product if opting to use cite-all or the selective method to support acute toxicity data requirements.

(6) Once a submission for a new product with public health organisms has been submitted and classified in either A540 or A541, additional organisms submitted for the same product before expiration of the first submission’s original decision review time period will result in reclassification of both the original and subsequent submission into the appropriate new category based on the sum of the number of organisms in both submissions. A reclassification would result in a new PRIA start date and require additional fees to meet the fee of the new category.

(7) Once a submission for a label amendment with public health organisms has been submitted and classified in either A570 or A573, additional organisms submitted for the same product before expiration of the first submission’s original decision review time period will result in reclassification of both the original and subsequent submission into the appropriate new category based on the sum of the number of organisms in both submissions. A reclassification would result in a new PRIA start date and require additional fees to meet the fee of the new category.

TABLE 10. — ANTIMICROBIALS DIVISION — EXPERIMENTAL USE PERMITS AND OTHER ACTIONS

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A520</td>
<td>94</td>
<td>Experimental Use Permit application, non-food use. (2)</td>
<td>9</td>
<td>6,383</td>
</tr>
</tbody>
</table>
TABLE 10. — ANTIMICROBIALS DIVISION — EXPERIMENTAL USE PERMITS AND OTHER ACTIONS—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)$^1$</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A521</td>
<td>95</td>
<td>Review of public health efficacy study protocol within AD, per AD Internal Guidance for the Efficacy Protocol Review Process; Code will also include review of public health efficacy study protocol and data review for devices making pesticidal claims; applicant-initiated; Tier 1.</td>
<td>4</td>
<td>4,726</td>
</tr>
<tr>
<td>A522</td>
<td>96</td>
<td>Review of public health efficacy study protocol outside AD by members of AD Efficacy Protocol Review Expert Panel; Code will also include review of public health efficacy study protocol and data review for devices making pesticidal claims; applicant-initiated; Tier 2.</td>
<td>12</td>
<td>12,156</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)(1)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>--------</td>
<td>------------</td>
<td>----------------------------------------------------------------------</td>
<td>-----------------------------------</td>
<td>-----------------------------</td>
</tr>
<tr>
<td>A537</td>
<td>97 (new)</td>
<td>New Active Ingredient/New Use, Experimental Use Permit application; Direct food use; Establish tolerance or tolerance exemption if required. Credit 45% of fee toward new active ingredient/new use application that follows.</td>
<td>18</td>
<td>153,156</td>
</tr>
<tr>
<td>A538</td>
<td>98 (new)</td>
<td>New Active Ingredient/New Use, Experimental Use Permit application; Indirect food use; Establish tolerance or tolerance exemption if required. Credit 45% of fee toward new active ingredient/new use application that follows.</td>
<td>18</td>
<td>95,724</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)(1)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>---------</td>
<td>------------</td>
<td>-----------------------------------------------------------------------</td>
<td>----------------------------------</td>
<td>------------------------------</td>
</tr>
<tr>
<td>A539</td>
<td>99 (new)</td>
<td>New Active Ingredient/New Use, Experimental Use Permit application; Nonfood use. Credit 45% of fee toward new active ingredient/new use application that follows.</td>
<td>15</td>
<td>92,163</td>
</tr>
<tr>
<td>A529</td>
<td>100</td>
<td>Amendment to Experimental Use Permit; requires data review or risk assessment. (2)</td>
<td>9</td>
<td>11,429</td>
</tr>
<tr>
<td>A523</td>
<td>101</td>
<td>Review of protocol other than a public health efficacy study (i.e., Toxicology or Exposure Protocols).</td>
<td>9</td>
<td>12,156</td>
</tr>
<tr>
<td>A571</td>
<td>102</td>
<td>Science reassessment: Cancer risk, refined ecological risk, and/or endangered species; applicant-initiated.</td>
<td>18</td>
<td>95,724</td>
</tr>
<tr>
<td>A533</td>
<td>103 (new)</td>
<td>Exemption from the requirement of an Experimental Use Permit. (2)</td>
<td>4</td>
<td>2,482</td>
</tr>
</tbody>
</table>
### TABLE 10. — ANTIMICROBIALS DIVISION — EXPERIMENTAL USE PERMITS AND OTHER ACTIONS—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(^{(1)})</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A534</td>
<td>104</td>
<td>Rebuttal of agency reviewed protocol, applicant initiated.</td>
<td>4</td>
<td>4,726</td>
</tr>
<tr>
<td>A535</td>
<td>105</td>
<td>Conditional Ruling on Pre-application Study Waiver or Data Bridging Argument; applicant-initiated.</td>
<td>6</td>
<td>2,409</td>
</tr>
<tr>
<td>A536</td>
<td>106</td>
<td>Conditional Ruling on Pre-application Direct Food, Indirect Food, Nonfood use determination; applicant-initiated.</td>
<td>4</td>
<td>2,482</td>
</tr>
</tbody>
</table>

(1) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.

(2) Where the action involves approval of a new or amended label, on or before the end date of the decision review time, the Agency shall provide to the applicant a draft accepted label, including any changes made by the Agency that differ from the applicant-submitted label and relevant supporting data reviewed by the Agency. The applicant will notify the Agency that the applicant either (a) agrees to all of the terms associated with the draft accepted label as amended by the Agency and requests that it be issued as the accepted final Agency-stamped label; or (b) does not agree to one or more of the terms of the draft accepted label as amended by the Agency and requests additional time to resolve the difference(s); or (c) withdraws the application without prejudice for subsequent resubmission, but forfeits the associated registration service fee. For cases described in (b), the applicant shall have up to 30 calendar days to reach agreement with the Agency on the final terms of the Agency-accepted label. If the applicant agrees to all of the terms of the accepted label as in (a), including upon resolution of differences in (b), the Agency shall provide an accepted final Agency-stamped label to the registrant within 2 business days following the registrant’s written or electronic confirmation of agreement to the Agency.
## TABLE 11. — BIOPESTICIDES DIVISION — NEW ACTIVE INGREDIENTS

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(_1)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B580</td>
<td>107</td>
<td>New active ingredient; food use; petition to establish a tolerance. (2)(3)</td>
<td>20</td>
<td>51,053</td>
</tr>
<tr>
<td>B590</td>
<td>108</td>
<td>New active ingredient; food use; petition to establish a tolerance exemption. (2)(3)</td>
<td>18</td>
<td>31,910</td>
</tr>
<tr>
<td>B600</td>
<td>109</td>
<td>New active ingredient; non-food use. (2)(3)</td>
<td>13</td>
<td>19,146</td>
</tr>
<tr>
<td>B610</td>
<td>110</td>
<td>New active ingredient; Experimental Use Permit application; petition to establish a temporary tolerance or temporary tolerance exemption. (3)</td>
<td>10</td>
<td>12,764</td>
</tr>
<tr>
<td>B611</td>
<td>111</td>
<td>New active ingredient; Experimental Use Permit application; petition to establish permanent tolerance exemption. (3)</td>
<td>12</td>
<td>12,764</td>
</tr>
<tr>
<td>B612</td>
<td>112</td>
<td>New active ingredient; no change to a permanent tolerance exemption. (2)(3)</td>
<td>10</td>
<td>17,550</td>
</tr>
</tbody>
</table>

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\(_1\) As Amended Through P.L. 116-8, Enacted March 08, 2019
### TABLE 11. — BIOPESTICIDES DIVISION — NEW ACTIVE INGREDIENTS—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(1)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B613</td>
<td>113</td>
<td>New active ingredient; petition to convert a temporary tolerance or a temporary tolerance exemption to a permanent tolerance or tolerance exemption. (2)(3)</td>
<td>11</td>
<td>17,550</td>
</tr>
<tr>
<td>B620</td>
<td>114</td>
<td>New active ingredient; Experimental Use Permit application; non-food use including crop destruct. (3)</td>
<td>7</td>
<td>6,383</td>
</tr>
</tbody>
</table>

(1) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.

(2) All requests for new uses (food and/or nonfood) contained in any application for a new active ingredient or a first food use are covered by the base fee for that new active ingredient or first food use application and retain the same decision time review period as the new active ingredient or first food use application. The application must be received by the agency in one package. The base fee for the category covers a maximum of five new products. Each application for an additional new product registration and new inert approval that is submitted in the new active ingredient application package or first food use application package is subject to the registration service fee for a new product or a new inert approval. All such associated applications that are submitted together will be subject to the new active ingredient or first food use decision review time. In the case of a new active ingredient application, until that new active ingredient is approved, any subsequent application for another new product containing the same active ingredient or an amendment to the proposed labeling will be deemed a new active ingredient application, subject to the registration service fee and decision review time for a new active ingredient. In the case of a first food use application, until that first food use is approved, any subsequent application for an additional new food use or uses will be subject to the registration service fee and decision review time for a first food use. Any information that (a) was neither requested nor required by the Agency, and (b) is submitted by the applicant at the applicant’s initiative to support the application after completion of the technical deficiency screening, and (c) is not itself a covered registration application, must be assessed 25% of the full registration service fee for the new active ingredient or first food use application.
(3) Where the action involves approval of a new or amended label, on or before the end date of the decision review time, the Agency shall provide to the applicant a draft accepted label, including any changes made by the Agency that differ from the applicant-submitted label and relevant supporting data reviewed by the Agency. The applicant will notify the Agency that the applicant either (a) agrees to all of the terms associated with the draft accepted label as amended by the Agency and requests that it be issued as the accepted final Agency-stamped label; or (b) does not agree to one or more of the terms of the draft accepted label as amended by the Agency and requests additional time to resolve the difference(s); or (c) withdraws the application without prejudice for subsequent resubmission, but forfeits the associated registration service fee. For cases described in (b), the applicant shall have up to 30 calendar days to reach agreement with the Agency on the final terms of the Agency-accepted label. If the applicant agrees to all of the terms of the accepted label as in (a), including upon resolution of differences in (b), the Agency shall provide an accepted final Agency-stamped label to the registrant within 2 business days following the registrant’s written or electronic confirmation of agreement to the Agency.

TABLE 12. — BIOPESTICIDES DIVISION — NEW USES

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B630</td>
<td>115</td>
<td>First food use; petition to establish a tolerance exemption. (2)(4)</td>
<td>13</td>
<td>12,764</td>
</tr>
<tr>
<td>B631</td>
<td>116</td>
<td>New food use; petition to amend an established tolerance. (3)(4)</td>
<td>12</td>
<td>12,764</td>
</tr>
<tr>
<td>B640</td>
<td>117</td>
<td>First food use; petition to establish a tolerance. (3)(4)</td>
<td>19</td>
<td>19,146</td>
</tr>
<tr>
<td>B643</td>
<td>118</td>
<td>New Food use; petition to amend an established tolerance exemption. (3)(4)</td>
<td>10</td>
<td>12,764</td>
</tr>
<tr>
<td>B642</td>
<td>119</td>
<td>First food use; indoor; food/food handling. (2)(4)</td>
<td>12</td>
<td>31,910</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)(1)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>---------</td>
<td>------------</td>
<td>------------------------------------------------------------------------</td>
<td>----------------------------------</td>
<td>------------------------------</td>
</tr>
<tr>
<td>B644</td>
<td>120</td>
<td>New use, no change to an established tolerance or tolerance exemption.  (3)(4)</td>
<td>8</td>
<td>12,764</td>
</tr>
<tr>
<td>B650</td>
<td>121</td>
<td>New use; non-food. (3)(4)</td>
<td>7</td>
<td>6,383</td>
</tr>
<tr>
<td>B645</td>
<td>122 (new)</td>
<td>New food use; Experimental Use Permit application; petition to amend or add a tolerance exemption. (4)</td>
<td>12</td>
<td>12,764</td>
</tr>
<tr>
<td>B646</td>
<td>123 (new)</td>
<td>New use; non-food use including crop destruct; Experimental Use Permit application. (4)</td>
<td>7</td>
<td>6,383</td>
</tr>
</tbody>
</table>

(1) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.
(2) All requests for new uses (food and/or nonfood) contained in any application for a new active ingredient or a first food use are covered by the base fee for that new active ingredient or first food use application and retain the same decision time review period as the new active ingredient or first food use application. The application must be received by the agency in one package. The base fee for the category covers a maximum of five new products. Each application for an additional new product registration and new inert approval that is submitted in the new active ingredient application package or first food use application package is subject to the registration service fee for a new product or a new inert approval. All such associated applications that are submitted together will be subject to the new active ingredient or first food use decision review time. In the case of a new active ingredient application, until that new active ingredient is approved, any subsequent application for another new product containing the same active ingredient or an amendment to the proposed labeling will be deemed a new active ingredient application, subject to the registration service fee and decision review time for a new active ingredient. In the case of a first food use application, until that first food use is approved, any subsequent application for an additional new food use or uses will be subject to the registration service fee and decision review time for a first food use. Any information that (a) was neither requested nor required by the Agency, and (b) is submitted by the applicant at the applicant’s initiative to support the application after completion of the technical deficiency screening, and (c) is not itself a covered registration application, must be assessed 25% of the full registration service fee for the new active ingredient or first food use application.

(3) Amendment applications to add the new use(s) to registered product labels are covered by the base fee for the new use(s). All items in the covered application must be submitted together in one package. Each application for an additional new product registration and new inert approval(s) that is submitted in the new use application package is subject to the registration service fee for a new product or a new inert approval. However, if a new use application only proposes to register the new use for a new product and there are no amendments in the application, then review of one new product application is covered by the new use fee. All such associated applications that are submitted together will be subject to the new use decision review time. Any application for a new product or an amendment to the proposed labeling (a) submitted subsequent to submission of the new use application and (b) prior to conclusion of its decision review time and (c) containing the same new uses, will be deemed a separate new-use application, subject to a separate registration service fee and new decision review time for a new use. If the new-use application includes non-food (indoor and/or outdoor), and food (outdoor and/or indoor) uses, the appropriate fee is due for each type of new use and the longest decision review time applies to all of the new uses requested in the application. Any information that (a) was neither requested nor required by the Agency, and (b) is submitted by the applicant at the applicant’s initiative to support the application after completion of the technical deficiency screen, and (c) is not itself a covered registration application, must be assessed 25% of the full registration service fee for the new use application.
Sec. 33 FEDERAL INSECTICIDE, FUNGICIDE, AND RODENTICIDE Act...

(4) Where the action involves approval of a new or amended label, on or before the end date of the decision review time, the Agency shall provide to the applicant a draft accepted label, including any changes made by the Agency that differ from the applicant-submitted label and relevant supporting data reviewed by the Agency. The applicant will notify the Agency that the applicant either (a) agrees to all of the terms associated with the draft accepted label as amended by the Agency and requests that it be issued as the accepted final Agency-stamped label; or (b) does not agree to one or more of the terms of the draft accepted label as amended by the Agency and requests additional time to resolve the difference(s); or (c) withdraws the application without prejudice for subsequent resubmission, but forfeits the associated registration service fee. For cases described in (b), the applicant shall have up to 30 calendar days to reach agreement with the Agency on the final terms of the Agency-accepted label. If the applicant agrees to all of the terms of the accepted label as in (a), including upon resolution of differences in (b), the Agency shall provide an accepted final Agency-stamped label to the registrant within 2 business days following the registrant’s written or electronic confirmation of agreement to the Agency.
TABLE 13. — BIOPESTICIDES DIVISION — NEW PRODUCTS

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B652</td>
<td>124</td>
<td>New product; registered source of active ingredient; requires petition to amend established tolerance or tolerance exemption; requires 1) submission of product specific data; or 2) citation of previously reviewed and accepted data; or 3) submission or citation of data generated at government expense; or 4) submission or citation of scientifically-sound rationale based on publicly available literature or other relevant information that addresses the data requirement; or 5) submission of a request for a data requirement to be waived supported by a scientifically-sound rationale explaining why the data requirement does not apply. (2)(3)</td>
<td>13</td>
<td>12,764</td>
</tr>
</tbody>
</table>
TABLE 13. — BIOPESTICIDES DIVISION — NEW PRODUCTS—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B660</td>
<td>125</td>
<td>New product; registered source of active ingredient(s); identical or substantially similar in composition and use to a registered product. No data review, or only product chemistry data; cite-all data citation, or selective data citation where applicant owns all required data or authorization from data owner is demonstrated. Category includes 100% re-pack of registered end-use or manufacturing-use product that requires no data submission or data matrix. For microbial pesticides, the active ingredient(s) must not be re-isolated. (2)(3)</td>
<td>4</td>
<td>1,278</td>
</tr>
</tbody>
</table>
TABLE 13. — BIOPESTICIDES DIVISION — NEW PRODUCTS— Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(1)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B670</td>
<td>126</td>
<td>New product; registered source of active ingredient(s); requires: 1) submission of product specific data; or 2) citation of previously reviewed and accepted data; or 3) submission or citation of data generated at government expense; or 4) submission or citation of a scientifically-sound rationale based on publicly available literature or other relevant information that addresses the data requirement; or 5) submission of a request for a data requirement to be waived supported by a scientifically-sound rationale explaining why the data requirement does not apply. (2)(3)</td>
<td>7</td>
<td>5,107</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>--------</td>
<td>------------</td>
<td>--------</td>
<td>-------------------------------</td>
<td>-----------------------------</td>
</tr>
<tr>
<td>B671</td>
<td>127</td>
<td>New product; unregistered source of active ingredient(s); requires a petition to amend an established tolerance or tolerance exemption; requires: 1) submission of product specific data; or 2) citation of previously reviewed and accepted data; or 3) submission or citation of data generated at government expense; or 4) submission or citation of a scientifically sound rationale based on publicly available literature or other relevant information that addresses the data requirement; or 5) submission of a request for a data requirement to be waived supported by a scientifically sound rationale explaining why the data requirement does not apply. (2)(3)</td>
<td>17</td>
<td>12,764</td>
</tr>
</tbody>
</table>
### TABLE 13. — BIOPESTICIDES DIVISION — NEW PRODUCTS—

**Continued**

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(^{(1)} )</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B672</td>
<td>128</td>
<td>New product; unregistered source of active ingredient(s); non-food use or food use requires: 1) submission of product specific data; or 2) citation of previously reviewed and accepted data; or 3) submission or citation of data generated at government expense; or 4) submission or citation of a scientifically sound rationale based on publicly available literature or other relevant information that addresses the data requirement; or 5) submission of a request for a data requirement to be waived supported by a scientifically sound rationale explaining why the data requirement does not apply. (2)(3)</td>
<td>13</td>
<td>9,118</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)(^{(1)})</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>---------</td>
<td>------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>------------------------------------------</td>
<td>----------------------------</td>
</tr>
<tr>
<td>B673</td>
<td>129</td>
<td>New product MUP/EP; unregistered source of active ingredient(s); citation of Technical Grade Active Ingredient (TGAI) data previously reviewed and accepted by the Agency. Requires an Agency determination that the cited data supports the new product. (2)(3)</td>
<td>10</td>
<td>5,107</td>
</tr>
<tr>
<td>B674</td>
<td>130</td>
<td>New product MUP; Repack of identical registered end-use product as a manufacturing-use product; same registered uses only. (2)(3)</td>
<td>4</td>
<td>1,278</td>
</tr>
<tr>
<td>B675</td>
<td>131</td>
<td>New Product MUP; registered source of active ingredient; submission of completely new generic data package; registered uses only. (2)(3)</td>
<td>10</td>
<td>9,118</td>
</tr>
</tbody>
</table>
New product; more than one active ingredient where one active ingredient is an unregistered source; product chemistry data must be submitted; requires: 1) submission of product specific data, and 2) citation of previously reviewed and accepted data; or 3) submission or citation of data generated at government expense; or 4) submission or citation of a scientifically-sound rationale based on publicly available literature or other relevant information that addresses the data requirement; or 5) submission of a request for a data requirement to be waived supported by a scientifically-sound rationale explaining why the data requirement does not apply. (2)(3)
New end-use non-food animal product with submission of two or more target animal safety studies; includes data and/or waivers of data for only:

- product chemistry and/or
- acute toxicity and/or
- public health pest efficacy and/or
- animal safety studies and/or
- child resistant packaging. (2)(3)

(1) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.

(2) An application for a new end-use product using a source of active ingredient that (a) is not yet registered but (b) has an application pending with the Agency for review, will be considered an application for a new product with an unregistered source of active ingredient.

(3) Where the action involves approval of a new or amended label, on or before the end date of the decision review time, the Agency shall provide to the applicant a draft accepted label, including any changes made by the Agency that differ from the applicant-submitted label and relevant supporting data reviewed by the Agency. The applicant will notify the Agency that the applicant either (a) agrees to all of the terms associated with the draft accepted label as amended by the Agency and requests that it be issued as the accepted final Agency-stamped label; or (b) does not agree to one or more of the terms of the draft accepted label as amended by the Agency and requests additional time to resolve the difference(s); or (c) withdraws the application without prejudice for subsequent resubmission, but forfeits the associated registration service fee. For cases described in (b), the applicant shall have up to 30 calendar days to reach agreement with the Agency on the final terms of the Agency-accepted label. If the applicant agrees to all of the terms of the accepted label as in (a), including upon resolution of differences in (b), the Agency shall provide an accepted final Agency-stamped label to the registrant within 2 business days following the registrant’s written or electronic confirmation of agreement to the Agency.
TABLE 14. — BIOPESTICIDES DIVISION — AMENDMENTS

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B621</td>
<td>134</td>
<td>Amendment; Experimental Use Permit; no change to an established temporary tolerance or tolerance exemption. (3)</td>
<td>7</td>
<td>5,107</td>
</tr>
<tr>
<td>B622</td>
<td>135</td>
<td>Amendment; Experimental Use Permit; petition to amend an established or temporary tolerance or tolerance exemption. (3)</td>
<td>11</td>
<td>12,764</td>
</tr>
<tr>
<td>B641</td>
<td>136</td>
<td>Amendment of an established tolerance or tolerance exemption. (3)</td>
<td>13</td>
<td>12,764</td>
</tr>
<tr>
<td>B680</td>
<td>137</td>
<td>Amendment; registered sources of active ingredient(s); no new use(s); no changes to an established tolerance or tolerance exemption. Requires data submission. (2)(3)</td>
<td>5</td>
<td>5,107</td>
</tr>
<tr>
<td>B681</td>
<td>138</td>
<td>Amendment; unregistered source of active ingredient(s). Requires data submission. (2)(3)</td>
<td>7</td>
<td>6,079</td>
</tr>
<tr>
<td>B683</td>
<td>139</td>
<td>Label amendment; requires review/update of previous risk assessment(s) without data submission (e.g., labeling changes to REI, PPE, PHI). (2)(3)</td>
<td>6</td>
<td>5,107</td>
</tr>
<tr>
<td>B684</td>
<td>140</td>
<td>Amending non-food animal product that includes submission of target animal safety data; previously registered. (2)(3)</td>
<td>8</td>
<td>8,820</td>
</tr>
</tbody>
</table>
TABLE 14. — BIOPESTICIDES DIVISION — AMENDMENTS—
Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B685</td>
<td>141 (new)</td>
<td>Amendment; add a new biochemical unregistered source of active ingredient or a new microbial production site. Requires submission of analysis of samples data and source/production site-specific manufacturing process description. (3)</td>
<td>5</td>
<td>5,107</td>
</tr>
</tbody>
</table>

(1) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.

(2) (a) EPA-initiated amendments shall not be charged registration service fees. (b) Registrant-initiated fast-track amendments are to be completed within the timelines specified in FIFRA Section 3(c)(3)(B) and are not subject to registration service fees. (c) Registrant-initiated fast-track amendments handled by the Antimicrobials Division are to be completed within the timelines specified in FIFRA Section 3(h) and are not subject to registration service fees. (d) Registrant initiated amendments submitted by notification under PR Notices, such as PR Notice 98–10, continue under PR Notice timelines and are not subject to registration service fees. (e) Submissions with data and requiring data review are subject to registration service fees.

(3) Where the action involves approval of a new or amended label, on or before the end date of the decision review time, the Agency shall provide to the applicant a draft accepted label, including any changes made by the Agency that differ from the applicant-submitted label and relevant supporting data reviewed by the Agency. The applicant will notify the Agency that the applicant either (a) agrees to all of the terms associated with the draft accepted label as amended by the Agency and requests that it be issued as the accepted final Agency-stamped label; or (b) does not agree to one or more of the terms of the draft accepted label as amended by the Agency and requests additional time to resolve the difference(s); or (c) withdraws the application without prejudice for subsequent resubmission, but forfeits the associated registration service fee. For cases described in (b), the applicant shall have up to 30 calendar days to reach agreement with the Agency on the final terms of the Agency-accepted label. If the applicant agrees to all of the terms of the accepted label as in (a), including upon resolution of differences in (b), the Agency shall provide an accepted final Agency-stamped label to the registrant within 2 business days following the registrant’s written or electronic confirmation of agreement to the Agency.
<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)((1))</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B690</td>
<td>142</td>
<td>New active ingredient; food or non-food use. (2)(6)</td>
<td>7</td>
<td>2,554</td>
</tr>
<tr>
<td>B700</td>
<td>143</td>
<td>Experimental Use Permit application; new active ingredient or new use. (6)</td>
<td>7</td>
<td>1,278</td>
</tr>
<tr>
<td>B701</td>
<td>144</td>
<td>Extend or amend Experimental Use Permit. (6)</td>
<td>4</td>
<td>1,278</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>--------</td>
<td>-----------</td>
<td>----------------------------------------------------------------------</td>
<td>-------------------------------</td>
<td>------------------------------</td>
</tr>
<tr>
<td>B710</td>
<td>145</td>
<td>New product; registered source of active ingredient(s); identical or substantially similar in composition and use to a registered product; no change in an established tolerance or tolerance exemption. No data review, or only product chemistry data; cite-all data citation, or selective data citation where applicant owns all required data or authorization from data owner is demonstrated. Category includes 100% re-package of registered end-use or manufacturing-use product that requires no data submission or data matrix. (3)(6)</td>
<td>4</td>
<td>1,278</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>--------</td>
<td>------------</td>
<td>--------</td>
<td>------------------------------</td>
<td>------------------------------</td>
</tr>
<tr>
<td>B720</td>
<td>146</td>
<td>New product; registered source of active ingredient(s); requires: 1) submission of product specific data; or 2) citation of previously reviewed and accepted data; or 3) submission or citation of data generated at government expense; or 4) submission or citation of a scientifically sound rationale based on publicly available literature or other relevant information that addresses the data requirement; or 5) submission of a request for a data requirement to be waived supported by a scientifically sound rationale explaining why the data requirement does not apply. (3)(6)</td>
<td>5</td>
<td>1,278</td>
</tr>
</tbody>
</table>
### TABLE 15. — BIOPESTICIDES DIVISION — SCLP—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B721</td>
<td>147</td>
<td>New product; unregistered source of active ingredient. (3)(6)</td>
<td>7</td>
<td>2,676</td>
</tr>
<tr>
<td>B722</td>
<td>148</td>
<td>New use and/or amendment; petition to establish a tolerance or tolerance exemption. (4)(5)(6)</td>
<td>7</td>
<td>2,477</td>
</tr>
<tr>
<td>B730</td>
<td>149</td>
<td>Label amendment requiring data submission. (4)(6)</td>
<td>5</td>
<td>1,278</td>
</tr>
</tbody>
</table>

(1) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.

(2) All requests for new uses (food and/or nonfood) contained in any application for a new active ingredient or a first food use are covered by the base fee for that new active ingredient or first food use application and retain the same decision time review period as the new active ingredient or first food use application. The application must be received by the agency in one package. The base fee for the category covers a maximum of five new products. Each application for an additional new product registration and new inert approval that is submitted in the new active ingredient application package or first food use application package is subject to the registration service fee for a new product or a new inert approval. All such associated applications that are submitted together will be subject to the new active ingredient or first food use decision review time. In the case of a new active ingredient application, until that new active ingredient is approved, any subsequent application for another new product containing the same active ingredient or an amendment to the proposed labeling will be deemed a new active ingredient application, subject to the registration service fee and decision review time for a new active ingredient. In the case of a first food use application, until that first food use is approved, any subsequent application for an additional new food use or uses will be subject to the registration service fee and decision review time for a first food use. Any information that (a) was neither requested nor required by the Agency, and (b) is submitted by the applicant at the applicant’s initiative to support the application after completion of the technical deficiency screening, and (c) is not itself a covered registration application, must be assessed 25% of the full registration service fee for the new active ingredient or first food use application.

(3) An application for a new end-use product using a source of active ingredient that (a) is not yet registered but (b) has an application pending with the Agency for review, will be considered an application for a new product with an unregistered source of active ingredient.
(4) (a) EPA-initiated amendments shall not be charged registration service fees. (b) Registrant-initiated fast-track amendments are to be completed within the timelines specified in FIFRA Section 3(c)(3)(B) and are not subject to registration service fees. (c) Registrant-initiated fast-track amendments handled by the Antimicrobials Division are to be completed within the timelines specified in FIFRA Section 3(h) and are not subject to registration service fees. (d) Registrant initiated amendments submitted by notification under PR Notices, such as PR Notice 98–10, continue under PR Notice timelines and are not subject to registration service fees. (e) Submissions with data and requiring data review are subject to registration service fees.

(5) Amendment applications to add the new use(s) to registered product labels are covered by the base fee for the new use(s). All items in the covered application must be submitted together in one package. Each application for an additional new product registration and new inert approval(s) that is submitted in the new use application package is subject to the registration service fee for a new product or a new inert approval. However, if a new use application only proposes to register the new use for a new product and there are no amendments in the application, then review of one new product application is covered by the new use fee. All such associated applications that are submitted together will be subject to the new use decision review time. Any application for a new product or an amendment to the proposed labeling (a) submitted subsequent to submission of the new use application and (b) prior to conclusion of its decision review time and (c) containing the same new uses, will be deemed a separate new-use application, subject to a separate registration service fee and new decision review time for a new use. If the new-use application includes non-food (indoor and/or outdoor), and food (outdoor and/or indoor) uses, the appropriate fee is due for each type of new use and the longest decision review time applies to all of the new uses requested in the application. Any information that (a) was neither requested nor required by the Agency, and (b) is submitted by the applicant at the applicant’s initiative to support the application after completion of the technical deficiency screen, and (c) is not itself a covered registration application, must be assessed 25% of the full registration service fee for the new use application.

(6) Where the action involves approval of a new or amended label, on or before the end date of the decision review time, the Agency shall provide to the applicant a draft accepted label, including any changes made by the Agency that differ from the applicant-submitted label and relevant supporting data reviewed by the Agency. The applicant will notify the Agency that the applicant either (a) agrees to all of the terms associated with the draft accepted label as amended by the Agency and requests that it be issued as the accepted final Agency-stamped label; or (b) does not agree to one or more of the terms of the draft accepted label as amended by the Agency and requests additional time to resolve the difference(s); or (c) withdraws the application without prejudice for subsequent resubmission, but forfeits the associated registration service fee. For cases described in (b), the applicant shall have up to 30 calendar days to reach agreement with the Agency on the final terms of the Agency-accepted label. If the applicant agrees to all of the terms of the accepted label as in (a), including upon resolution of differences in (b), the Agency shall provide an accepted final Agency-stamped label to the registrant within 2 business days following the registrant’s written or electronic confirmation of agreement to the Agency.
TABLE 16. — BIOPESTICIDES DIVISION — OTHER ACTIONS

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(^{(1)})</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B614</td>
<td>150</td>
<td>Pre-application; Conditional Ruling on rationales for addressing a data requirement in lieu of data; applicant-initiated; applies to one rationale at a time.</td>
<td>3</td>
<td>2,530</td>
</tr>
<tr>
<td>B615</td>
<td>151</td>
<td>Rebuttal of agency reviewed protocol, applicant initiated.</td>
<td>3</td>
<td>2,530</td>
</tr>
<tr>
<td>B682</td>
<td>152</td>
<td>Protocol review; applicant initiated; excludes time for HSRB review.</td>
<td>3</td>
<td>2,432</td>
</tr>
</tbody>
</table>

\(^{(1)}\) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.
TABLE 17. — BIOPESTICIDES DIVISION — PIP

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(^{(1)})</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
</table>
| B740    | 153        | Experimental Use Permit application; no petition for tolerance/tolerance exemption. Includes:  
1. non-food/feed use(s) for a new (2) or registered (3) PIP (12);  
2. food/feed use(s) for a new or registered PIP with crop destruct (12);  
3. food/feed use(s) for a new or registered PIP in which an established tolerance/tolerance exemption exists for the intended use(s). (4)(12) | 6                                         | 95,724                       |
### Table 17. — Biopesticides Division — PIP—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B741</td>
<td>154</td>
<td>Experimental Use Permit application; no petition for tolerance/tolerance exemption. Includes: 1. non-food/feed use(s) for a new (2) or registered (3) PIP; 2. food/feed use(s) for a new or registered PIP with crop destruct; 3. food/feed use(s) for a new or registered PIP in which an established tolerance/tolerance exemption exists for the intended use(s); SAP Review. (12)</td>
<td>12</td>
<td>159,538</td>
</tr>
<tr>
<td>B750</td>
<td>155</td>
<td>Experimental Use Permit application; with a petition to establish a temporary or permanent tolerance/tolerance exemption for the active ingredient. Includes new food/feed use for a registered (3) PIP. (4)(12)</td>
<td>9</td>
<td>127,630</td>
</tr>
</tbody>
</table>
TABLE 17. — BIOPESTICIDES DIVISION — PIP—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B770</td>
<td>156</td>
<td>Experimental Use Permit application; new (2) PIP; with petition to establish a temporary tolerance/tolerance exemption for the active ingredient; credit 75% of B771 fee toward registration application for a new active ingredient that follows; SAP review. (5)(12)</td>
<td>15</td>
<td>191,444</td>
</tr>
<tr>
<td>B771</td>
<td>157</td>
<td>Experimental Use Permit application; new (2) PIP; with petition to establish a temporary tolerance/tolerance exemption for the active ingredient; credit 75% of B771 fee toward registration application for a new active ingredient that follows. (12)</td>
<td>10</td>
<td>127,630</td>
</tr>
<tr>
<td>B772</td>
<td>158</td>
<td>Application to amend or extend an Experimental Use Permit; no petition since the established tolerance/tolerance exemption for the active ingredient is unaffected. (12)</td>
<td>3</td>
<td>12,764</td>
</tr>
</tbody>
</table>
TABLE 17. — BIOPESTICIDES DIVISION — PIP—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B773</td>
<td>159</td>
<td>Application to amend or extend an Experimental Use Permit; with petition to extend a temporary tolerance/tolerance exemption for the active ingredient. (12)</td>
<td>5</td>
<td>31,910</td>
</tr>
<tr>
<td>B780</td>
<td>160</td>
<td>Registration application; new (2) PIP; non-food/feed. (12)</td>
<td>12</td>
<td>159,537</td>
</tr>
<tr>
<td>B790</td>
<td>161</td>
<td>Registration application; new (2) PIP; non-food/feed; SAP review. (5)(12)</td>
<td>18</td>
<td>223,351</td>
</tr>
<tr>
<td>B800</td>
<td>162</td>
<td>Registration application; new (2) PIP; with petition to establish permanent tolerance/tolerance exemption for the active ingredient based on an existing temporary tolerance/tolerance exemption. (12)</td>
<td>13</td>
<td>172,300</td>
</tr>
<tr>
<td>B810</td>
<td>163</td>
<td>Registration application; new (2) PIP; with petition to establish permanent tolerance/tolerance exemption for the active ingredient based on an existing temporary tolerance/tolerance exemption. SAP review. (5)(12)</td>
<td>19</td>
<td>236,114</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>--------</td>
<td>------------</td>
<td>------------------------------------------------------------------------</td>
<td>------------------------------</td>
<td>------------------------------</td>
</tr>
<tr>
<td>B820</td>
<td>164</td>
<td>Registration application; new (2) PIP; with petition to establish or amend a permanent tolerance/exemption of an active ingredient. (12)</td>
<td>15</td>
<td>204,208</td>
</tr>
<tr>
<td>B840</td>
<td>165</td>
<td>Registration application; new (2) PIP; with petition to establish or amend a permanent tolerance/exemption of an active ingredient. SAP review. (5)(12)</td>
<td>21</td>
<td>268,022</td>
</tr>
<tr>
<td>B851</td>
<td>166</td>
<td>Registration application; new event of a previously registered PIP active ingredient(s); no petition since permanent tolerance/exemption is already established for the active ingredient(s). (12)</td>
<td>9</td>
<td>127,630</td>
</tr>
<tr>
<td>B870</td>
<td>167</td>
<td>Registration application; registered (3) PIP; new product; new use; no petition since a permanent tolerance/exemption is already established for the active ingredient(s). (4) (12)</td>
<td>9</td>
<td>38,290</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)(^{(1)})</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>--------</td>
<td>------------</td>
<td>------------------------------------------------------------------------</td>
<td>------------------------------------------</td>
<td>------------------------------</td>
</tr>
<tr>
<td>B880</td>
<td>168</td>
<td>Registration application; registered (3) PIP; new product or new terms of registration; additional data submitted; no petition since a permanent tolerance/tolerance exemption is already established for the active ingredient(s). (6) (7) (12)</td>
<td>9</td>
<td>31,910</td>
</tr>
<tr>
<td>B881</td>
<td>169</td>
<td>Registration application; registered (3) PIP; new product or new terms of registration; additional data submitted; no petition since a permanent tolerance/tolerance exemption is already established for the active ingredient(s). SAP review. (5)(6)(7)(12)</td>
<td>15</td>
<td>95,724</td>
</tr>
</tbody>
</table>
TABLE 17. — BIOPESTICIDES DIVISION — PIP—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(1)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B882</td>
<td>170</td>
<td>Registration application; new (2) PIP, seed increase with negotiated acreage cap and time-limited registration; with petition to establish a permanent tolerance/tolerance exemption for the active ingredient based on an existing temporary tolerance/tolerance exemption; SAP Review. (8)(12)</td>
<td>15</td>
<td>191,444</td>
</tr>
<tr>
<td>B883</td>
<td>171</td>
<td>Registration application; new (2) PIP, seed increase with negotiated acreage cap and time-limited registration; with petition to establish a permanent tolerance/tolerance exemption for the active ingredient based on an existing temporary tolerance/tolerance exemption. (8) (12)</td>
<td>9</td>
<td>127,630</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)(^{(1)})</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>--------</td>
<td>------------</td>
<td>----------------------------------------------------------------------</td>
<td>----------------------------------------</td>
<td>------------------------------</td>
</tr>
<tr>
<td>B884</td>
<td>172</td>
<td>Registration application; new (2) PIP, seed increase with negotiated acreage cap and time-limited registration; with petition to establish a permanent tolerance/tolerance exemption for the active ingredient. (8)(12)</td>
<td>12</td>
<td>159,537</td>
</tr>
<tr>
<td>B885</td>
<td>173</td>
<td>Registration application; registered (3) PIP, seed increase; breeding stack of previously approved PIPs, same crop; no petition since a permanent tolerance/tolerance exemption is already established for the active ingredient(s). (9)(12)</td>
<td>6</td>
<td>31,910</td>
</tr>
<tr>
<td>B886</td>
<td>174 (new)</td>
<td>Registration application; new (2) PIP, seed increase with negotiated acreage cap and time-limited registration; with petition to establish a permanent tolerance/tolerance exemption for the active ingredient. SAP Review. (8) (12)</td>
<td>18</td>
<td>223,351</td>
</tr>
</tbody>
</table>
TABLE 17. — BIOPESTICIDES DIVISION — PIP—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(1)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B890</td>
<td>175</td>
<td>Application to amend a seed increase registration; converts registration to commercial registration; no petition since permanent tolerance/tolerance exemption is already established for the active ingredient(s). (12)</td>
<td>9</td>
<td>63,816</td>
</tr>
<tr>
<td>B891</td>
<td>176</td>
<td>Application to amend a seed increase registration; converts registration to a commercial registration; no petition since a permanent tolerance/tolerance exemption already established for the active ingredient(s); SAP review. (5)(12)</td>
<td>15</td>
<td>127,630</td>
</tr>
<tr>
<td>B900</td>
<td>177</td>
<td>Application to amend a registration, including actions such as extending an expiration date, modifying an IRM plan, or adding an insect to be controlled. (10)(11)(12)</td>
<td>6</td>
<td>12,764</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)&lt;sup&gt;(1)&lt;/sup&gt;</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>---------</td>
<td>------------</td>
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<td>------------------------------------------</td>
<td>-----------------------------</td>
</tr>
<tr>
<td>B901</td>
<td>178</td>
<td>Application to amend a registration, including actions such as extending an expiration date, modifying an IRM plan, or adding an insect to be controlled. SAP review. (10) (11) (12)</td>
<td>12</td>
<td>76,578</td>
</tr>
<tr>
<td>B902</td>
<td>179</td>
<td>PIP Protocol review.</td>
<td>3</td>
<td>6,383</td>
</tr>
<tr>
<td>B903</td>
<td>180</td>
<td>Inert ingredient tolerance exemption; e.g., a marker such as NPT II; reviewed in BPPD.</td>
<td>6</td>
<td>63,816</td>
</tr>
<tr>
<td>B904</td>
<td>181</td>
<td>Import tolerance or tolerance exemption; processed commodities/food only (inert or active ingredient).</td>
<td>9</td>
<td>127,630</td>
</tr>
<tr>
<td>B905</td>
<td>182</td>
<td>SAP Review.</td>
<td>6</td>
<td>63,816</td>
</tr>
<tr>
<td>B905</td>
<td>182 (new)</td>
<td>Petition to establish a temporary tolerance/tolerance exemption for one or more active ingredients.</td>
<td>3</td>
<td>31,907</td>
</tr>
<tr>
<td>B907</td>
<td>184 (new)</td>
<td>Petition to establish a temporary tolerance/tolerance exemption for one or more active ingredients based on an existing temporary tolerance/tolerance exemption.</td>
<td>3</td>
<td>12,764</td>
</tr>
</tbody>
</table>
TABE 17. — BIOPESTICIDES DIVISION — PIP—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B908</td>
<td>185</td>
<td>Petition to establish a temporary tolerance/tolerance exemption for one or more active ingredients or inert ingredients.</td>
<td>3</td>
<td>44,671</td>
</tr>
</tbody>
</table>

(1) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.
(2) New PIP = a PIP with an active ingredient that has not been registered.
(3) Registered PIP = a PIP with an active ingredient that is currently registered.
(4) Transfer registered PIP through conventional breeding for new food/feed use, such as from field corn to sweet corn.
(5) The scientific data involved in this category are complex. EPA often seeks technical advice from the Scientific Advisory Panel on risks that pesticides pose to wildlife, farm workers, pesticide applicators, non-target species, as well as insect resistance, and novel scientific issues surrounding new technologies. The scientists of the SAP neither make nor recommend policy decisions. They provide advice on the science used to make these decisions. Their advice is invaluable to the EPA as it strives to protect humans and the environment from risks posed by pesticides. Due to the time it takes to schedule and prepare for meetings with the SAP, additional time and costs are needed.
(6) Registered PIPs stacked through conventional breeding.
(7) Deployment of a registered PIP with a different IRM plan (e.g., seed blend).
(8) The negotiated acreage cap will depend upon EPA’s determination of the potential environmental exposure, risk(s) to non-target organisms, and the risk of targeted pest developing resistance to the pesticidal substance. The uncertainty of these risks may reduce the allowable acreage, based upon the quantity and type of non-target organism data submitted and the lack of insect resistance management data, which is usually not required for seed-increase registrations. Registrants are encouraged to consult with EPA prior to submission of a registration application in this category.
(9) Application can be submitted prior to or concurrently with an application for commercial registration.
(10) For example, IRM plan modifications that are applicant-initiated.
(11) EPA-initiated amendments shall not be charged fees.
(12) Where the action involves approval of a new or amended label, on or before the end date of the decision review time, the Agency shall provide to the applicant a draft accepted label, including any changes made by the Agency that differ from the applicant-submitted label and relevant supporting data reviewed by the Agency. The applicant will notify the Agency that the applicant either (a) agrees to all of the terms associated with the draft accepted label as amended by the Agency and requests that it be issued as the accepted final Agency-stamped label; or (b) does not agree to one or more of the terms of the draft accepted label as amended by the Agency and requests additional time to resolve the difference(s); or (c) withdraws the application without prejudice for subsequent resubmission, but forfeits the associated registration service fee. For cases described in (b), the applicant shall have up to 30 calendar days to reach agreement with the Agency on the final terms of the Agency-accepted label. If the applicant agrees to all of the terms of the accepted label as in (a), including upon resolution of differences in (b), the Agency shall provide an accepted final Agency-stamped label to the registrant within 2 business days following the registrant’s written or electronic confirmation of agreement to the Agency.

TABLE 18. — INERT INGREDIENTS

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months) (1)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>I001</td>
<td>186</td>
<td>Approval of new food use inert ingredient. (2)(3)</td>
<td>13</td>
<td>27,000</td>
</tr>
<tr>
<td>I002</td>
<td>187</td>
<td>Amend currently approved inert ingredient tolerance or exemption from tolerance; new data. (2)</td>
<td>11</td>
<td>7,500</td>
</tr>
<tr>
<td>I003</td>
<td>188</td>
<td>Amend currently approved inert ingredient tolerance or exemption from tolerance; no new data. (2)</td>
<td>9</td>
<td>3,308</td>
</tr>
<tr>
<td>I004</td>
<td>189</td>
<td>Approval of new non-food use inert ingredient. (2)</td>
<td>6</td>
<td>11,025</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>---------</td>
<td>------------</td>
<td>--------</td>
<td>------------------------------</td>
<td>------------------------------</td>
</tr>
<tr>
<td>1005</td>
<td>190</td>
<td>Amend currently approved non-food use inert ingredient with new use pattern; new data. (2)</td>
<td>6</td>
<td>5,513</td>
</tr>
<tr>
<td>1006</td>
<td>191</td>
<td>Amend currently approved non-food use inert ingredient with new use pattern; no new data. (2)</td>
<td>3</td>
<td>3,308</td>
</tr>
<tr>
<td>1007</td>
<td>192</td>
<td>Approval of substantially similar non-food use inert ingredients when original inert is compositionally similar with similar use pattern. (2)</td>
<td>4</td>
<td>1,654</td>
</tr>
<tr>
<td>1008</td>
<td>193</td>
<td>Approval of new or amended polymer inert ingredient, food use. (2)</td>
<td>5</td>
<td>3,749</td>
</tr>
<tr>
<td>1009</td>
<td>194</td>
<td>Approval of new or amended polymer inert ingredient, non-food use. (2)</td>
<td>4</td>
<td>3,087</td>
</tr>
<tr>
<td>1010</td>
<td>195</td>
<td>Petition to amend a single tolerance exemption descriptor, or single non-food use descriptor, to add ( \leq 10 ) CASRN; no new data. (2)</td>
<td>6</td>
<td>1,654</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>---------</td>
<td>------------</td>
<td>------------------------------------------------------------------------</td>
<td>------------------------------</td>
<td>------------------------------</td>
</tr>
<tr>
<td>1011</td>
<td>196 (new)</td>
<td>Approval of new food use safener with tolerance or exemption from tolerance. (2)(8)</td>
<td>24</td>
<td>597,683</td>
</tr>
<tr>
<td>1012</td>
<td>197 (new)</td>
<td>Approval of new non-food use safener. (2)(8)</td>
<td>21</td>
<td>415,241</td>
</tr>
<tr>
<td>1013</td>
<td>198 (new)</td>
<td>Approval of additional food use for previously approved safener with tolerance or exemption from tolerance. (2)</td>
<td>15</td>
<td>62,975</td>
</tr>
<tr>
<td>1014</td>
<td>199 (new)</td>
<td>Approval of additional non-food use for previously approved safener. (2)</td>
<td>15</td>
<td>25,168</td>
</tr>
<tr>
<td>1015</td>
<td>200 (new)</td>
<td>Approval of new generic data for previously approved food use safener. (2)</td>
<td>24</td>
<td>269,728</td>
</tr>
<tr>
<td>1016</td>
<td>201 (new)</td>
<td>Approval of amendment(s) to tolerance and label for previously approved safener. (2)</td>
<td>13</td>
<td>55,776</td>
</tr>
</tbody>
</table>

(1) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.
(2) If another covered application is submitted that depends upon an application to approve an inert ingredient, each application will be subject to its respective registration service fee. The decision review time line for both submissions will be the longest of the associated applications. If the application covers multiple ingredients grouped by EPA into one chemical class, a single registration service fee will be assessed for approval of those ingredients.
(3) If EPA data rules are amended to newly require clearance under section 408 of the FFDCA for an ingredient of an antimicrobial product where such ingredient was not previously subject to such a clearance, then review of the data for such clearance of such product is not subject to a registration service fee for the tolerance action for two years from the effective date of the rule.

(4) Any other covered application that is associated with and dependent on the HSRB review will be subject to its separate registration service fee. The decision review times for the associated actions run concurrently, but will end at the date of the latest review time.

(5) Any other covered application that is associated with and dependent on the SAP review will be subject to its separate registration service fee. The decision review time for the associated action will be extended by the decision review time for the SAP review.

(6) An application for a new end-use product using a source of active ingredient that (a) is not yet registered but (b) has an application pending with the Agency for review, will be considered an application for a new product with an unregistered source of active ingredient.

(7) Where the action involves approval of a new or amended label, on or before the end date of the decision review time, the Agency shall provide to the applicant a draft accepted label, including any changes made by the Agency that differ from the applicant-submitted label and relevant supporting data reviewed by the Agency. The applicant will notify the Agency that the applicant either (a) agrees to all of the terms associated with the draft accepted label as amended by the Agency and requests that it be issued as the accepted final Agency-stamped label; or (b) does not agree to one or more of the terms of the draft accepted label as amended by the Agency and requests additional time to resolve the difference(s); or (c) withdraws the application without prejudice for subsequent resubmission, but forfeits the associated registration service fee. For cases described in (b), the applicant shall have up to 30 calendar days to reach agreement with the Agency on the final terms of the Agency-accepted label. If the applicant agrees to all of the terms of the accepted label as in (a), including upon resolution of differences in (b), the Agency shall provide an accepted final Agency-stamped label to the registrant within 2 business days following the registrant’s written or electronic confirmation of agreement to the Agency.

(8) If a new safener is submitted in the same package as a new active ingredient, and that new active ingredient is determined to be reduced risk, then the safener would get the same reduced timeframe as the new active ingredient.
### TABLE 19. — EXTERNAL REVIEW AND MISCELLANEOUS ACTIONS

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>M001</td>
<td>202</td>
<td>Study protocol requiring Human Studies Review Board review as defined in 40 CFR Part 26 in support of an active ingredient. (4)</td>
<td>9</td>
<td>7,938</td>
</tr>
<tr>
<td>M002</td>
<td>203</td>
<td>Completed study requiring Human Studies Review Board review as defined in 40 CFR Part 26 in support of an active ingredient. (4)</td>
<td>9</td>
<td>7,938</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)(^{(1)})</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>---------</td>
<td>------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------</td>
<td>------------------------------------------</td>
<td>------------------------------</td>
</tr>
<tr>
<td>M003</td>
<td>204</td>
<td>External technical peer review of new active ingredient, product, or amendment (e.g., consultation with FIFRA Scientific Advisory Panel) for an action with a decision timeframe of less than 12 months. Applicant initiated request based on a requirement of the Administrator, as defined by FIFRA § 25(d), in support of a novel active ingredient, or unique use pattern or application technology. Excludes PIP active ingredients. (5)</td>
<td>12</td>
<td>63,945</td>
</tr>
</tbody>
</table>
### Table 19. — External Review and Miscellaneous Actions—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)(^{(1)})</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>M004</td>
<td>205</td>
<td>External technical peer review of new active ingredient, product, or amendment (e.g., consultation with FIFRA Scientific Advisory Panel) for an action with a decision timeframe of greater than 12 months. Applicant initiated request based on a requirement of the Administrator, as defined by FIFRA § 25(d), in support of a novel active ingredient, or unique use pattern or application technology. Excludes PIP active ingredients. (5)</td>
<td>18</td>
<td>63,945</td>
</tr>
</tbody>
</table>
### TABLE 19. — EXTERNAL REVIEW AND MISCELLANEOUS ACTIONS—Continued

<table>
<thead>
<tr>
<th>EPA No.</th>
<th>New CR No.</th>
<th>Action</th>
<th>Decision Review Time (Months)&lt;sup&gt;(1)&lt;/sup&gt;</th>
<th>Registration Service Fee ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>M005</td>
<td>206</td>
<td>New Product: Combination, Contains a combination of active ingredients from a registered and/or unregistered source; conventional, antimicrobial and/or biopesticide. Requires coordination with other regulatory divisions to conduct review of data, label and/or verify the validity of existing data as cited. Only existing uses for each active ingredient in the combination product. (6)(7)</td>
<td>9</td>
<td>22,050</td>
</tr>
<tr>
<td>M006</td>
<td>207</td>
<td>Request for up to 5 letters of certification (Gold Seal) for one actively registered product (excludes distributor products). (8)</td>
<td>1</td>
<td>277</td>
</tr>
<tr>
<td>M007</td>
<td>208</td>
<td>Request to extend Exclusive Use of data as provided by FIFRA Section 3(c)(1)(F')(ii).</td>
<td>12</td>
<td>5,513</td>
</tr>
<tr>
<td>EPA No.</td>
<td>New CR No.</td>
<td>Action</td>
<td>Decision Review Time (Months)(1)</td>
<td>Registration Service Fee ($)</td>
</tr>
<tr>
<td>--------</td>
<td>------------</td>
<td>----------------------------------------------------------------------</td>
<td>----------------------------------</td>
<td>-----------------------------</td>
</tr>
<tr>
<td>M008</td>
<td>209</td>
<td>Request to grant Exclusive Use of data as provided by FIFRA Section 3(c)(1)(F)(vi) for a minor use, when a FIFRA Section 2(ll)(2) determination is required.</td>
<td>15</td>
<td>1,654</td>
</tr>
<tr>
<td>M009</td>
<td>210 (new)</td>
<td>Non-FIFRA Regulated Determination: Applicant initiated, per product.</td>
<td>4</td>
<td>2,363</td>
</tr>
<tr>
<td>M010</td>
<td>211 (new)</td>
<td>Conditional ruling on pre-application, product substantial similarity.</td>
<td>4</td>
<td>2,363</td>
</tr>
<tr>
<td>M011</td>
<td>212 (new)</td>
<td>Label amendment to add the DiE logo; requires data review; no other label changes. (9)</td>
<td>4</td>
<td>3,648</td>
</tr>
</tbody>
</table>

(1) A decision review time that would otherwise end on a Saturday, Sunday, or federal holiday, will be extended to end on the next business day.

(2) If another covered application is submitted that depends upon an application to approve an inert ingredient, each application will be subject to its respective registration service fee. The decision review time line for both submissions will be the longest of the associated applications. If the application covers multiple ingredients grouped by EPA into one chemical class, a single registration service fee will be assessed for approval of those ingredients.

(3) If EPA data rules are amended to newly require clearance under section 408 of the FFDCA for an ingredient of an antimicrobial product where such ingredient was not previously subject to such a clearance, then review of the data for such clearance of such product is not subject to a registration service fee for the tolerance action for two years from the effective date of the rule.
(4) Any other covered application that is associated with and dependent on the HSRB review will be subject to its separate registration service fee. The decision review times for the associated actions run concurrently, but will end at the date of the latest review time.

(5) Any other covered application that is associated with and dependent on the SAP review will be subject to its separate registration service fee. The decision review time for the associated action will be extended by the decision review time for the SAP review.

(6) An application for a new end-use product using a source of active ingredient that (a) is not yet registered but (b) has an application pending with the Agency for review, will be considered an application for a new product with an unregistered source of active ingredient.

(7) Where the action involves approval of a new or amended label, on or before the end date of the decision review time, the Agency shall provide to the applicant a draft accepted label, including any changes made by the Agency that differ from the applicant-submitted label and relevant supporting data reviewed by the Agency. The applicant will notify the Agency that the applicant either (a) agrees to all of the terms associated with the draft accepted label as amended by the Agency and requests that it be issued as the accepted final Agency-stamped label; or (b) does not agree to one or more of the terms of the draft accepted label as amended by the Agency and requests additional time to resolve the difference(s); or (c) withdraws the application without prejudice for subsequent resubmission, but forfeits the associated registration service fee. For cases described in (b), the applicant shall have up to 30 calendar days to reach agreement with the Agency on the final terms of the Agency-accepted label. If the applicant agrees to all of the terms of the accepted label as in (a), including upon resolution of differences in (b), the Agency shall provide an accepted final Agency-stamped label to the registrant within 2 business days following the registrant’s written or electronic confirmation of agreement to the Agency.

(8) Due to low fee and short time frame this category is not eligible for small business waivers. Gold seal applies to one registered product.

(9) This category includes amendments the sole purpose of which is to add DfE (or equivalent terms that do not use “safe” or derivatives of “safe”) logos to a label. DfE is a voluntary program. A label bearing a DfE logo is not considered an Agency endorsement because the ingredients in the qualifying product must meet objective, scientific criteria established and widely publicized by EPA.

(4) PENDING PESTICIDE REGISTRATION APPLICATIONS.—

(A) IN GENERAL.—An applicant that submitted a registration application to the Administrator before the effective date of the Pesticide Registration Improvement Act of 2003, but that is not required to pay a registration service fee under paragraph (2)(B), may, on a voluntary basis, pay a registration service fee in accordance with paragraph (2)(B).

(B) VOLUNTARY FEE.—The Administrator may not compel payment of a registration service fee for an application described in subparagraph (A).

(C) DOCUMENTATION.—An application for which a voluntary registration service fee is paid under this paragraph shall be submitted with documentation certifying—

(i) payment of the registration service fee; or

(ii) a request for a waiver from or reduction of the registration service fee.
(5) Resubmission of Covered Applications.—If a covered application is submitted by a person that paid the fee for the application under paragraph (2), is determined by the Administrator to be complete, and is not approved or is withdrawn (without a waiver or refund), the submission of the same covered application by the same person (or a licensee, assignee, or successor of the person) shall not be subject to a fee under paragraph (2).

(6) Fee Adjustment.—

(A) In General.—Effective for a covered application received during the period beginning on October 1, 2019, and ending on September 30, 2021, the Administrator shall increase by 5 percent the registration service fee payable for the application under paragraph (3).

(B) Additional Adjustment.—Effective for a covered application received on or after October 1, 2021, the Administrator shall increase by an additional 5 percent the registration service fee in effect as of September 30, 2021.

(C) Publication.—The Administrator shall publish in the Federal Register the service fee schedules revised pursuant to this paragraph.

(7) Waivers and Reductions.—

(A) In General.—An applicant for a covered application may request the Administrator to waive or reduce the amount of a registration service fee payable under this section under the circumstances described in subparagraphs (D) through (G), except that no waiver or fee reduction shall be provided in connection with a request for a letter of certification (commonly referred to as a Gold Seal letter).

(B) Documentation.—

(i) In General.—A request for a waiver from or reduction of the registration service fee shall be accompanied by appropriate documentation demonstrating the basis for the waiver or reduction.

(ii) Certification.—The applicant shall provide to the Administrator a written certification, signed by a responsible officer, that the documentation submitted to support the waiver or reduction request is accurate.

(iii) Inaccurate Documentation.—An application shall be subject to the applicable registration service fee payable under paragraph (3) if, at any time, the Administrator determines that—

(I) the documentation supporting the waiver or reduction request is not accurate; or

(II) based on the documentation or any other information, the waiver or reduction should not have been granted or should not be granted.

(C) Determination to Grant or Deny Request.—As soon as practicable, but not later than 60 days, after the date on which the Administrator receives a request for a waiver or reduction of a registration service fee under this paragraph, the Administrator shall—
Sec. 33 FEDERAL INSECTICIDE, FUNGICIDE, AND RODENTICIDE A...

(i) determine whether to grant or deny the request; and
(ii) notify the applicant of the determination.

(D) MINOR USES.—
(i) IN GENERAL.—The Administrator may exempt from, or waive a portion of, the registration service fee for an application for minor uses for a pesticide.
(ii) SUPPORTING DOCUMENTATION.—An applicant requesting a waiver or exemption under this subparagraph shall provide supporting documentation that demonstrates, to the satisfaction of the Administrator, that anticipated revenues from the uses that are the subject of the application would be insufficient to justify imposition of the full application fee.

(E) IR–4 EXEMPTION.—The Administrator shall exempt an application from the registration service fee if the Administrator determines that—
(i) the application is solely associated with a tolerance petition submitted in connection with the Inter-Regional Project Number 4 (IR–4) as described in section 2 of Public Law 89–106 (7 U.S.C. 450i(e)); and
(ii) the exemption is in the public interest.

(F) SMALL BUSINESSES.—
(i) IN GENERAL.—The Administrator shall waive 50 percent of the registration service fees payable by an entity for a covered application under this section if the entity is a small business (as defined in section 4(i)(1)(E)(ii)) at the time of application.
(ii) WAIVER OF FEES.—The Administrator shall waive 75 percent of the registration service fees payable by an entity under this section if the entity—
(I) is a small business (as defined in section 4(i)(1)(E)(ii)) at the time of application; and
(II) has average annual global gross revenues described in section 4(i)(1)(E)(ii)(I)(bb) that do not exceed $10,000,000, at the time of application.
(iii) FORMATION FOR WAIVER.—The Administrator shall not grant a waiver under this subparagraph if the Administrator determines that the entity submitting the application has been formed or manipulated primarily for the purpose of qualifying for the waiver.
(iv) DOCUMENTATION.—An entity requesting a waiver under this subparagraph shall provide to the Administrator—
(I) documentation demonstrating that the entity is a small business (as defined in section 4(i)(1)(E)(ii)) at the time of application; and
(II) if the entity is requesting a waiver of 75 percent of the applicable registration service fees payable under this section, documentation demonstrating that the entity has an average annual global gross revenue described in section 4(i)(1)(E)(ii)(I)(bb) that does not exceed $10,000,000, at the time of application.
(G) **FEDERAL AND STATE AGENCY EXEMPTIONS.**—An agency of the Federal Government or a State government shall be exempt from covered registration service fees under this section.

(8) **REFUNDS.**—

(A) **EARLY WITHDRAWALS.**—If, during the first 60 days after the beginning of the applicable decision time review period under subsection (f)(3), a covered application is withdrawn by the applicant, the Administrator shall refund all but 25 percent.¹⁷ of the total registration service fee payable under paragraph (3) for the application.

(B) **WITHDRAWALS AFTER THE FIRST 60 DAYS OF DECISION REVIEW TIME PERIOD.**—

(i) **IN GENERAL.**—If a covered application is withdrawn after the first 60 days of the applicable decision time review period, the Administrator shall determine what portion, if any, of the total registration service fee payable under paragraph (3) for the application may be refunded based on the proportion of the work completed at the time of withdrawal.

(ii) **TIMING.**—The Administrator shall—

(I) make the determination described in clause (i) not later than 90 days after the date the application is withdrawn; and

(II) provide any refund as soon as practicable after the determination.

(C) **DISCRETIONARY REFUNDS.**—

(i) **IN GENERAL.**—In the case of a covered application that has been filed with the Administrator and has not been withdrawn by the applicant, but for which the Administrator has not yet made a final determination, the Administrator may refund a portion of a covered registration service fee if the Administrator determines that the refund is justified.

(ii) **BASIS.**—The Administrator may provide a refund for an application under this subparagraph—

(I) on the basis that, in reviewing the application, the Administrator has considered data submitted in support of another covered application;

(II) on the basis that the Administrator completed portions of the review of the application before the effective date of this section; or

(III) on the basis that the Administrator rejected the application under subsection (f)(4)(B).

(D) **CREDITED FEES.**—In determining whether to grant a refund under this paragraph, the Administrator shall take into account any portion of the registration service fees credited under paragraph (2) or (4).

(c) **PESTICIDE REGISTRATION FUND.**—

(1) **ESTABLISHMENT.**—There is established in the Treasury of the United States a Pesticide Registration Fund to be used

¹⁷So in original. The period probably should not appear.
in carrying out this section (referred to in this section as the “Fund”), consisting of—

(A) such amounts as are deposited in the Fund under paragraph (2);
(B) any interest earned on investment of amounts in the Fund under paragraph (5); and
(C) any proceeds from the sale or redemption of investments held in the Fund.

(2) Deposits in Fund.—Subject to paragraph (4), the Administrator shall deposit fees collected under this section in the Fund.

(3) Expenditures from Fund.—

(A) In General.—Subject to subparagraphs (B) and (C) and paragraph (4), the Administrator may make expenditures from the Fund—

(i) to cover the costs associated with the review and decisionmaking pertaining to all applications for which registration service fees have been paid under this section; and
(ii) to otherwise carry out this section.

(B) Worker Protection, Partnership Grants, and Pesticide Safety Education.—

(i) In General.—For each of fiscal years 2013 through 2023, the Administrator shall use approximately 1/17 of the amount in the Fund (but not less than $1,000,000) to enhance scientific and regulatory activities relating to worker protection, with an emphasis on field-worker populations in the United States.

(ii) Partnership Grants.—Of the amounts in the Fund, the Administrator shall use for partnership grants, for each of fiscal years 2013 through 2023, $500,000.

(iii) Pesticide Safety Education Program.—Of the amounts in the Fund, the Administrator shall use $500,000 for each of fiscal years 2013 through 2023 to carry out the pesticide safety education program.

(4) Collections and Appropriations Acts.—The fees authorized by this section and amounts deposited in the Fund—

(A) shall be collected and made available for obligation only to the extent provided in advance in appropriations Acts; and

(B) shall be available without fiscal year limitation.

(5) Unused Funds.—

(A) In General.—Amounts in the Fund not currently needed to carry out this section shall be—

(i) maintained readily available or on deposit;

(ii) invested in obligations of the United States or guaranteed by the United States; or

\[10^{16}\]Indention of clauses (i) through (iii) is so in original (as amended by sec. 5(e)(3)(A) of Public Law 110–94). Clauses probably should be indented.
(iii) invested in obligations, participations, or other instruments that are lawful investments for fiduciary, trust, or public funds.

(B) USE OF INVESTMENT INCOME.—After consultation with the Secretary of the Treasury, the Administrator may use income from investments described in clauses (ii) and (iii) of subparagraph (A) to carry out this section.

d) ASSESSMENT OF FEES.—

(1) DEFINITION OF COVERED FUNCTIONS.—In this subsection, the term “covered functions” means functions of the Office of Pesticide Programs of the Environmental Protection Agency, as identified in key programs and projects of the final operating plan for the Environmental Protection Agency submitted as part of the budget process for fiscal year 2002, regardless of any subsequent transfer of 1 or more of the functions to another office or agency or the subsequent transfer of a new function to the Office of Pesticide Programs.

(2) MINIMUM AMOUNT OF APPROPRIATIONS.—Registration service fees may not be assessed for a fiscal year under this section unless the amount of appropriations for salaries, contracts, and expenses for the functions (as in existence in fiscal year 2012) of the Office of Pesticide Programs of the Environmental Protection Agency for the fiscal year (excluding the amount of any fees appropriated for the fiscal year) are equal to or greater than the amount of appropriations for covered functions for fiscal year 2012 (excluding the amount of any fees appropriated for the fiscal year).

(3) USE OF FEES.—Registration service fees authorized by this section shall be available, in the aggregate, only to defray increases in the costs associated with the review and decision-making for the review of pesticide registration applications and associated tolerances (including increases in the number of full-time equivalent positions in the Environmental Protection Agency engaged in those activities) over the costs for fiscal year 2002, excluding costs paid from fees appropriated for the fiscal year.

(4) SUBSEQUENT AUTHORITY.—If the Administrator does not assess registration service fees under subsection (b) during any portion of a fiscal year as the result of paragraph (2) and is subsequently permitted to assess the fees under subsection (b) during the fiscal year, the Administrator shall assess and collect the fees, without any modification in rate, at any time during the fiscal year, notwithstanding any provisions of subsection (b) relating to the date fees are to be paid.

(e) REFORMS TO REDUCE DECISION TIME REVIEW PERIODS.—To the maximum extent practicable consistent with the degrees of risk presented by pesticides and the type of review appropriate to evaluate risks, the Administrator shall identify and evaluate reforms to the pesticide registration process under this Act with the goal of reducing decision review periods in effect on the effective date of the Pesticide Registration Improvement Extension Act of 2018 for pesticide registration actions for covered pesticide registration applications (including reduced risk applications). Such reforms shall include identifying opportunities for streamlining review processes.
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for applications for a new active ingredient or a new use and providing prompt feedback to applicants during such review process.

(f) Decision Time Review Periods.—

(1) In General.—Not later than 30 days after the effective date of the Pesticide Registration Improvement Extension Act of 2018, the Administrator shall make publicly available a schedule of decision review periods for covered pesticide registration actions or for any other action covered by a table specified in subsection (b)(3) and corresponding registration service fees under this Act.

(2) Report.—The schedule shall be the same as the applicable schedule provided under subsection (b)(3).

(3) Applications Subject to Decision Time Review Periods.—The decision time review periods specified in paragraph (1) shall apply to—

(A) covered pesticide registration applications subject to registration service fees under subsection (b)(2);
(B) covered pesticide registration applications for which an applicant has voluntarily paid registration service fees under subsection (b)(4); and
(C) applications for any other action covered by a table specified in subsection (b)(3).

(4) Start of Decision Time Review Period.—

(A) In General.—Except as provided in subparagraphs (C), (D), and (E), in the case of a covered application accompanied by the registration service fee required under this section, the decision time review period begins 21 days after the date on which the Administrator receives the covered application and fee.

(B) Initial Content and Preliminary Technical Screenings.—

(i) Screenings.—

(I) Initial Content.—Not later than 21 days after receiving an application and the required registration service fee, the Administrator shall conduct an initial screening of the contents of the application in accordance with clause (iii).

(II) Preliminary Technical Screening.—After conducting the initial content screening described in subclause (I) and in accordance with clause (iv), the Administrator shall conduct a preliminary technical screening—

(aa) not later than 45 days after the date on which the decision time review period begins (for applications with decision time review periods of not more than 180 days); and
(bb) not later than 90 days after the date on which the decision time review period begins (for applications with decision time review periods greater than 180 days).

(ii) Rejection.—

(I) In General.—If the Administrator determines at any time before the Administrator completes the preliminary technical screening under
clause (i)(II) that the application failed the initial content or preliminary technical screening and the applicant does not correct the failure before the date that is 10 business days after the applicant receives a notification of the failure, the Administrator shall reject the application.

(II) WRITTEN NOTIFICATION.—The Administrator shall make every effort to provide a written notification of a rejection under subclause (I) during the 10-day period that begins on the date the Administrator completes the preliminary technical screening.

(iii) REQUIREMENTS OF INITIAL CONTENT SCREENING.—In conducting an initial content screening of an application, the Administrator shall determine whether—

(I)(aa) the applicable registration service fee has been paid; or

(bb) at least 25 percent of the applicable registration service fee has been paid and the application contains a waiver or refund request for the outstanding amount and documentation establishing the basis for the waiver request; and

(II) the application appears to contain all the necessary forms, data, and draft labeling, formatted in accordance with guidance published by the Administrator.

(iv) REQUIREMENTS OF PRELIMINARY TECHNICAL SCREENING.—In conducting a preliminary technical screening of an application, the Administrator shall determine if—

(I) the application and the data and information submitted with the application are accurate and complete; and

(II) the application, data, and information are consistent with the proposed labeling and any proposal for a tolerance or exemption from the requirement for a tolerance under section 408 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 346a), and are such that, subject to full review under the standards of this Act, could result in the granting of the application.

(C) APPLICATIONS WITH WAIVER OR REDUCTION REQUESTS.—

(i) IN GENERAL.—In the case of an application submitted with a request for a waiver or reduction of registration service fees under subsection (b)(7), the decision time review period shall be determined in accordance with this subparagraph.

(ii) REQUEST GRANTED WITH NO ADDITIONAL FEES REQUIRED.—If the Administrator grants the waiver or reduction request and no additional fee is required, the decision time review period begins on the earlier of—
(I) the date on which the Administrator grants the request; or
(II) the date that is 60 days after the date of receipt of the application.

(iii) REQUEST GRANTED WITH ADDITIONAL FEES REQUIRED.—If the Administrator grants the waiver or reduction request, in whole or in part, but an additional registration service fee is required, the decision time review period begins on the date on which the Administrator receives certification of payment of the applicable registration service fee.

(iv) REQUEST DENIED.—If the Administrator denies the waiver or reduction request, the decision time review period begins on the date on which the Administrator receives certification of payment of the applicable registration service fee.

(D) PENDING APPLICATIONS.—
(i) IN GENERAL.—The start of the decision time review period for applications described in clause (ii) shall be the date on which the Administrator receives certification of payment of the applicable registration service fee.

(ii) APPLICATIONS.—Clause (i) applies to—
(I) covered pesticide registration applications for which voluntary fees have been paid under subsection (b)(4); and
(II) covered pesticide registration applications received on or after the effective date of the Pesticide Registration Improvement Act of 2003 but submitted without the applicable registration service fee required under this section due to the inability of the Administrator to assess fees under subsection (d)(1).

(E) 2003 WORK PLAN.—In the case of a covered pesticide registration application listed in the Registration Division 2003 Work Plan of the Office of Pesticide Programs of the Environmental Protection Agency, the decision time review period begins on the date that is 30 days after the effective date of the Pesticide Registration Improvement Act of 2003.

(5) EXTENSION OF DECISION TIME REVIEW PERIOD.—The Administrator and the applicant may mutually agree in writing to extend a decision time review period under this subsection.

(g) JUDICIAL REVIEW.—
(1) IN GENERAL.—Any applicant adversely affected by the failure of the Administrator to make a determination on the application of the applicant for registration of a new active ingredient or new use for which a registration service fee is paid under this section may obtain judicial review of the failure solely under this section.

(2) SCOPE.—
(A) IN GENERAL.—In an action brought under this subsection, the only issue on review is whether the Administrator failed to make a determination on the application
specified in paragraph (1) by the end of the applicable decision time review period required under subsection (f) for the application.

(B) OTHER ACTIONS.—No other action authorized or required under this section shall be judicially reviewable by a Federal or State court.

(3) TIMING.—

(A) IN GENERAL.—A person may not obtain judicial review of the failure of the Administrator to make a determination on the application specified in paragraph (1) before the expiration of the 2-year period that begins on the date on which the decision time review period for the application ends.

(B) MEETING WITH ADMINISTRATOR.—To be eligible to seek judicial review under this subsection, a person seeking the review shall first request in writing, at least 120 days before filing the complaint for judicial review, a decision review meeting with the Administrator.

(4) REMEDIES.—The Administrator may not be required or permitted to refund any portion of a registration service fee paid in response to a complaint that the Administrator has failed to make a determination on the covered pesticide registration application specified in paragraph (1) by the end of the applicable decision review period.

(h) ACCOUNTING.—The Administrator shall—

(1) provide an annual accounting of the registration service fees paid to the Administrator and disbursed from the Fund, by providing financial statements in accordance with—

(A) the Chief Financial Officers Act of 1990 (Public Law 101–576; 104 Stat. 2838) and amendments made by that Act; and

(B) the Government Management Reform Act of 1994 (Public Law 103–356; 108 Stat. 3410) and amendments made by that Act;

(2) provide an accounting describing expenditures from the Fund authorized under subsection (c); and

(3) provide an annual accounting describing collections and expenditures authorized under subsection (d).

(i) AUDITING.—

(1) FINANCIAL STATEMENTS OF AGENCIES.—For the purpose of section 3515(c) of title 31, United States Code, the Fund shall be considered a component of an executive agency.

(2) COMPONENTS.—The annual audit required under sections 3515(b) and 3521 of that title of the financial statements of activities under this section shall include an analysis of—

(A) the fees collected under subsection (b) and disbursed;

(B) compliance with subsection (f);

(C) the amount appropriated to meet the requirements of subsection (d)(1); and

(D) the reasonableness of the allocation of the overhead allocation of costs associated with the review and decisionmaking pertaining to applications under this section.
(3) **INSPECTOR GENERAL.**—The Inspector General of the Environmental Protection Agency shall—

(A) conduct the annual audit required under this subsection; and

(B) report the findings and recommendations of the audit to the Administrator and to the appropriate committees of Congress.

(j) **PERSONNEL LEVELS.**—All full-time equivalent positions supported by fees authorized and collected under this section shall not be counted against the agency-wide personnel level goals of the Environmental Protection Agency.

(k) **REPORTS.**—

(1) **IN GENERAL.**—Not later than March 1, 2005, and each March 1 thereafter through March 1, 2023, the Administrator shall publish an annual report describing actions taken under this section.

(2) **CONTENTS.**—The report shall include—

(A) a review of the progress made in carrying out each requirement of subsections (e) and (f), including—

(i) the number of applications reviewed, including the decision times for each application specified in subsection (f);

(ii) the number of label amendments that have been reviewed using electronic means;

(iii) the amount of money from the Reregistration and Expedited Processing Fund used to carry out inert ingredient review and review of similar applications under section 4(k)(3);

(iv) the number of applications completed for identical or substantially similar applications under section 3(c)(3)(B), including the number of such applications completed within 90 days pursuant to that section;

(v) the number of actions pending in each category of actions described in subsection (f)(3), as well as the number of inert ingredients;

(vi) to the extent determined appropriate by the Administrator and consistent with the authorities of the Administrator and limitations on delegation of functions by the Administrator, recommendations for—

(I) expanding the use of self-certification in all appropriate areas of the registration process;

(II) providing for accreditation of outside reviewers and the use of outside reviewers to conduct the review of major portions of applications;

(III) reviewing the scope of use of the notification process to cover broader categories of registration actions;

(IV) providing for electronic submission and review of labels, including process improvements to further enhance the procedures used in electronic label review; and
(V) the allowance and use of summaries of acute toxicity studies;
(vii) the use of performance-based contracts, other contracts, and procurement to ensure that—
(I) the goals of this Act for the timely review of applications for registration are met; and
(II) the registration program is administered in the most productive and cost-effective manner practicable; and
(viii) the number of extensions of decision time review periods agreed to under subsection (f)(5) along with a description of the reason that the Administrator was unable to make a decision within the initial decision time review period;

(B) a description of the staffing and resources relating to the costs associated with the review and decisionmaking pertaining to applications;

(C) a review of the progress in meeting the timeline requirements of section 4(g);

(D) a review of the progress in carrying out section 3(g), including—

(i) the number of pesticides or pesticide cases reviewed and the number of registration review decisions completed, including—
(I) the number of cases cancelled;
(II) the number of cases requiring risk mitigation measures;
(III) the number of cases removing risk mitigation measures;
(IV) the number of cases with no risk mitigation needed; and
(V) the number of cases in which risk mitigation has been fully implemented;

(ii) a description of the staffing and resources relating to the costs associated with the review and decisionmaking relating to reregistration and registration review for compliance with the deadlines specified in this Act;

(iii) to the extent determined appropriate by the Administrator and consistent with the authorities of the Administrator and limitations on delegation of functions by the Administrator, recommendations for—

(I) process improvements in the handling of registration review under section 3(g);

(II) providing for accreditation of outside reviewers and the use of outside reviewers in the registration review process; and

(III) streamlining the registration review process, consistent with section 3(g);

(E) a review of the progress in meeting the timeline requirements for the review of antimicrobial pesticide products under section 3(h);
(F) a review of the progress in carrying out the review of inert ingredients, including the number of applications pending, the number of new applications, the number of applications reviewed, staffing, and resources devoted to the review of inert ingredients and recommendations to improve the timeliness of review of inert ingredients;

(G) a review of the progress made toward—
   (i) carrying out paragraphs (4) and (5) of section 4(k) and the amounts from the Reregistration and Expedited Processing Fund used for the purposes described in such paragraphs;
   (ii) implementing enhancements to—
      (I) the electronic tracking of covered applications;
      (II) the electronic tracking of conditional registrations;
      (III) the endangered species database;
      (IV) the electronic review of labels submitted with covered applications; and
      (V) the electronic review and assessment of confidential statements of formula submitted with covered applications; and
   (iii) facilitating public participation in certain registration actions and the registration review process by providing electronic notification to interested parties of additions to the public docket;

(H) the number of applications rejected by the Administrator under the initial content and preliminary technical screening conducted under subsection (f)(4);

(I) a review of the progress made in updating the Pesticide Incident Data System, including progress toward making the information contained in the System available to the public (as the Administrator determines is appropriate);

(J) an assessment of the public availability of summary pesticide usage data;

(K) a review of the progress made in developing, updating, and implementing product performance test guidelines for pesticide products that are intended to control invertebrate pests of significant public health importance and, by regulation, prescribing product performance data requirements for such pesticide products registered under section 3;

(L) a review of the progress made in the priority review and approval of new pesticides to control invertebrate public health pests that may transmit vector-borne disease for use in the United States, including each territory or possession of the United States, and United States military installations globally;

(M) a review of the progress made in implementing enhancements to the good laboratory practices standards compliance monitoring program established under part 160 of title 40 of the Code of Federal Regulations (or successor regulations);
(N) the number of approvals for active ingredients, new uses, and pesticide end use products granted in connection with the Design for the Environment program (or any successor program) of the Environmental Protection Agency; and

(O) with respect to funds in the Pesticide Registration Fund reserved under subsection (c)(3), a review that includes—

(i) a description of the amount and use of such funds—

(I) to carry out activities relating to worker protection under clause (i) of subsection (c)(3)(B);

(II) to award partnership grants under clause (ii) of such subsection; and

(III) to carry out the pesticide safety education program under clause (iii) of such subsection;

(ii) an evaluation of the appropriateness and effectiveness of the activities, grants, and program described in clause (i);

(iii) a description of how stakeholders are engaged in the decision to fund such activities, grants, and program; and

(iv) with respect to activities relating to worker protection carried out under subparagraph (B)(i) of such subsection, a summary of the analyses from stakeholders, including from worker community-based organizations, on the appropriateness and effectiveness of such activities.

(3) METHOD.—The Administrator shall publish a report required by this subsection by such method as the Administrator determines to be the most effective for efficiently disseminating the report, including publication of the report on the Internet site of the Environmental Protection Agency.

(4) OTHER REPORT.—

(A) SCOPE.—In addition to the annual report described in paragraph (1), not later than October 1, 2016, the Administrator shall submit to the Committee on Agriculture of the House of Representatives and the Committee on Agriculture, Nutrition, and Forestry of the Senate a report that includes an analysis of the impact of maintenance fees on small businesses that have—

(i) 10 or fewer employees; and

(ii) annual global gross revenue that does not exceed $2,000,000.

(B) INFORMATION REQUIRED.—In conducting the analysis described in subparagraph (A), the Administrator shall collect, and include in the report under that subparagraph, information on—

(i) the number of small businesses described in subparagraph (A) that are paying maintenance fees; and

(ii) the number of registrations each company holds.
(l) **Savings Clause.**—Nothing in this section affects any other duties, obligations, or authorities established by any other section of this Act, including the right to judicial review of duties, obligations, or authorities established by any other section of this Act.

(m) **Termination of Effectiveness.**—

(1) **In General.**—Except as provided in paragraph (2), the authority provided by this section terminates on September 30, 2023.

(2) **Phase Out.**—

(A) **Fiscal Year 2024.**—During fiscal year 2024, the requirement to pay and collect registration service fees applies, except that the level of registration service fees payable under this section shall be reduced 40 percent below the level in effect on September 30, 2023.

(B) **Fiscal Year 2025.**—During fiscal year 2025, the requirement to pay and collect registration service fees applies, except that the level of registration service fees payable under this section shall be reduced 70 percent below the level in effect on September 30, 2023.

(C) **September 30, 2025.**—Effective September 30, 2025, the requirement to pay and collect registration service fees terminates.

(D) **Decision Review Periods.**—

(i) **Pending Applications.**—In the case of an application received under this section before September 30, 2023, the application shall be reviewed in accordance with subsection (f).

(ii) **New Applications.**—In the case of an application received under this section on or after September 30, 2023, subsection (f) shall not apply to the application.

**SEC. 34.** [7 U.S.C. 136x] **SEVERABILITY.**

If any provision of this Act or the application thereof to any person or circumstance is held invalid, the invalidity shall not affect other provisions or applications of this Act which can be given effect without regard to the invalid provision or application, and to this end the provisions of this Act are severable.

**SEC. 35.** [7 U.S.C. 136y] **AUTHORIZATION FOR APPROPRIATIONS.**

There is authorized to be appropriated to carry out this Act (other than section 23(a))—

(1) $83,000,000 for fiscal year 1989, of which not more than $13,735,500 shall be available for research under this Act;

(2) $95,000,000 for fiscal year 1990, of which not more than $14,343,600 shall be available for research under this Act; and

(3) $95,000,000 for fiscal year 1991, of which not more than $14,978,200 shall be available for research under this Act.