

for public viewing in EDOCKET. For further information about the electronic docket, see EPA's notice describing the electronic docket at 67 FR 38102 (May 31, 2002), or go to www.epa.gov/edocket.

Title: NSPS for Flexible Vinyl and Urethane Coating and Printing (40 CFR part 60, subpart FFF), OMB Control Number 2060-0073, EPA ICR Number 1157.07. This is a request to renew an existing, approved collection that is scheduled to expire on June 30, 2003. Under OMB regulations, the Agency may continue to conduct or sponsor the collection of information while this submission is pending at OMB.

Abstract: The NSPS for Flexible Vinyl and Urethane Coating and Printing, published at 40 CFR part 60, subpart FFF, were proposed on January 18, 1983, and promulgated on June 29, 1984. These standards of performance for this category of new stationary sources of hazardous air pollutants are required by section 111 of the Clean Air Act. Facilities may meet the standards by using materials with a low concentration of Volatile Organic Compounds (VOCs), or by installing emission control devices. The information that is required to be submitted to the Agency or kept at the facility is needed to insure compliance with the regulation. These include initial one-time notifications, performance tests plans and reports and records of maintenance and shutdown, startup, and malfunctions. For facilities that install continuous monitoring system (CMS) there are performance tests, and maintenance reports. Excess emissions reports are submitted semiannually. Responses to the collection of information are mandatory (40 CFR part 60, subpart FFF).

An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB Control Number. The OMB Control Numbers for EPA's regulations are listed in 40 CFR part 9 and 48 CFR chapter 15, and are identified on the form and/or instrument, if applicable.

Burden Statement: The annual public reporting and recordkeeping burden for this collection of information are estimated to average 14 (rounded) hours per response. Burden means the total time, effort, or financial resources expended by persons to generate, maintain, retain, or disclose or provide information to or for a Federal agency. This includes the time needed to review instructions; develop, acquire, install, and utilize technology and systems for the purposes of collecting, validating, and verifying information, processing

and maintaining information, and disclosing and providing information; adjust the existing ways to comply with any previously applicable instructions and requirements; train personnel to be able to respond to a collection of information; search data sources; complete and review the collection of information; and transmit or otherwise disclose the information.

Respondents/Affected Entities: Flexible Vinyl and Urethane Coating and Printing Manufacturers.

Estimated Number of Respondents: 20.

Frequency of Response: Semiannually, initially.

Estimated Total Annual Hour Burden: 593.

Estimated Total Capital and Operations & Maintenance (O & M) Annual Costs: \$61,000.

Changes in the Estimates: There is an increase of 78 hours in the total estimated burden currently identified in the OMB Inventory of Approved ICR Burdens. This increase in burden from the most recently approved ICR is due to more accurate estimates of existing sources. We have assumed that there will be one new source (respondent) over the three years period of this ICR. The change in respondent cost is also due to a labor rate change where the rate has increased over the three years period of this ICR.

Dated: June 24, 2003.

Doreen Sterling,

Acting Director, Collection Strategies Division.

[FR Doc. 03-16734 Filed 7-1-03; 8:45 am]

BILLING CODE 6560-50-P

ENVIRONMENTAL PROTECTION AGENCY

[OPP-2003-0212; FRL-7312-9]

Indoxacarb; Notice of Filing a Pesticide Petition to Establish a Tolerance for a Certain Pesticide Chemical in or on Food

AGENCY: Environmental Protection Agency (EPA).

ACTION: Notice.

SUMMARY: This notice announces the initial filing of a pesticide petition proposing the establishment of regulations for residues of a certain pesticide chemical in or on various food commodities.

DATES: Comments, identified by docket ID number OPP-2003-0212, must be received on or before August 1, 2003.

ADDRESSES: Comments may be submitted electronically, by mail, or

through hand delivery/courier. Follow the detailed instructions as provided in Unit I. of the **SUPPLEMENTARY INFORMATION**.

FOR FURTHER INFORMATION CONTACT: Rita Kumar, Registration Division (7505C), Office of Pesticide Programs, Environmental Protection Agency, 1200 Pennsylvania Ave., NW., Washington, DC 20460-0001; telephone number: (703) 308-8291; e-mail address: kumar.rita@epa.gov.

SUPPLEMENTARY INFORMATION:

I. General Information

A. Does this Action Apply to Me?

You may be potentially affected by this action if you are an agricultural producer, food manufacturer, or pesticide manufacturer. Potentially affected entities may include, but are not limited to:

- Crop production (NAICS 111)
- Animal production (NAICS 112)
- Food manufacturing (NAICS 311)
- Pesticide manufacturing (NAICS 32532)

This listing is not intended to be exhaustive, but rather provides a guide for readers regarding entities likely to be affected by this action. Other types of entities not listed in this unit could also be affected. The North American Industrial Classification System (NAICS) codes have been provided to assist you and others in determining whether this action might apply to certain entities. If you have any questions regarding the applicability of this action to a particular entity, consult the person listed under **FOR FURTHER INFORMATION CONTACT**.

B. How Can I Get Copies of This Document and Other Related Information?

1. **EPA Docket.** EPA has established an official public docket for this action under docket ID number OPP-2003-0212. The official public docket consists of the documents specifically referenced in this action, any public comments received, and other information related to this action. Although, a part of the official docket, the public docket does not include Confidential Business Information (CBI) or other information whose disclosure is restricted by statute. The official public docket is the collection of materials that is available for public viewing at the Public Information and Records Integrity Branch (PIRIB), Rm. 119, Crystal Mall #2, 1921 Jefferson Davis Hwy., Arlington, VA. This docket facility is open from 8:30 a.m. to 4 p.m., Monday through Friday, excluding legal

holidays. The docket telephone number is (703) 305-5805.

2. *Electronic access.* You may access this **Federal Register** document electronically through the EPA Internet under the "**Federal Register**" listings at <http://www.epa.gov/fedrgstr/>.

An electronic version of the public docket is available through EPA's electronic public docket and comment system, EPA dockets. You may use EPA Dockets at <http://www.epa.gov/edocket/> to submit or view public comments, access the index listing of the contents of the official public docket, and to access those documents in the public docket that are available electronically. Although not all docket materials may be available electronically, you may still access any of the publicly available docket materials through the docket facility identified in Unit I.B.1. Once in the system, select "search," then key in the appropriate docket ID number.

Certain types of information will not be placed in the EPA dockets. Information claimed as CBI and other information whose disclosure is restricted by statute, which is not included in the official public docket, will not be available for public viewing in EPA's electronic public docket. EPA's policy is that copyrighted material will not be placed in EPA's electronic public docket but will be available only in printed, paper form in the official public docket. To the extent feasible, publicly available docket materials will be made available in EPA's electronic public docket. When a document is selected from the index list in EPA dockets, the system will identify whether the document is available for viewing in EPA's electronic public docket. Although, not all docket materials may be available electronically, you may still access any of the publicly available docket materials through the docket facility identified in Unit I.B. EPA intends to work towards providing electronic access to all of the publicly available docket materials through EPA's electronic public docket.

For public commenters, it is important to note that EPA's policy is that public comments, whether submitted electronically or in paper, will be made available for public viewing in EPA's electronic public docket as EPA receives them and without change, unless the comment contains copyrighted material, CBI, or other information whose disclosure is restricted by statute. When EPA identifies a comment containing copyrighted material, EPA will provide a reference to that material in the version of the comment that is placed in EPA's electronic public docket. The

entire printed comment, including the copyrighted material, will be available in the public docket.

Public comments submitted on computer disks that are mailed or delivered to the docket will be transferred to EPA's electronic public docket. Public comments that are mailed or delivered to the docket will be scanned and placed in EPA's electronic public docket. Where practical, physical objects will be photographed, and the photograph will be placed in EPA's electronic public docket along with a brief description written by the docket staff.

C. How and To Whom Do I Submit Comments?

You may submit comments electronically, by mail, or through hand delivery/courier. To ensure proper receipt by EPA, identify the appropriate docket ID number in the subject line on the first page of your comment. Please ensure that your comments are submitted within the specified comment period. Comments received after the close of the comment period will be marked "late." EPA is not required to consider these late comments. If you wish to submit CBI or information that is otherwise protected by statute, please follow the instructions in Unit I.D. Do not use EPA Dockets or e-mail to submit CBI or information protected by statute.

1. *Electronically.* If you submit an electronic comment as prescribed in this unit, EPA recommends that you include your name, mailing address, and an e-mail address or other contact information in the body of your comment. Also include this contact information on the outside of any disk or CD ROM you submit, and in any cover letter accompanying the disk or CD ROM. This ensures that you can be identified as the submitter of the comment and allows EPA to contact you in case EPA cannot read your comment due to technical difficulties or needs further information on the substance of your comment. EPA's policy is that EPA will not edit your comment, and any identifying or contact information provided in the body of a comment will be included as part of the comment that is placed in the official public docket, and made available in EPA's electronic public docket. If EPA cannot read your comment due to technical difficulties and cannot contact you for clarification, EPA may not be able to consider your comment.

i. *EPA Dockets.* Your use of EPA's electronic public docket to submit comments to EPA electronically is EPA's preferred method for receiving comments. Go directly to EPA Dockets

at <http://www.epa.gov/edocket/>, and follow the online instructions for submitting comments. Once in the system, select "search," and then key in docket ID number OPP-2003-0212. The system is an "anonymous access" system, which means EPA will not know your identity, e-mail address, or other contact information unless you provide it in the body of your comment.

ii. *E-mail.* Comments may be sent by e-mail to opp-docket@epa.gov, Attention: Docket ID number OPP-2003-0212. In contrast to EPA's electronic public docket, EPA's e-mail system is not an "anonymous access" system. If you send an e-mail comment directly to the docket without going through EPA's electronic public docket, EPA's e-mail system automatically captures your e-mail address. E-mail addresses that are automatically captured by EPA's e-mail system are included as part of the comment that is placed in the official public docket, and made available in EPA's electronic public docket.

iii. *Disk or CD ROM.* You may submit comments on a disk or CD ROM that you mail to the mailing address identified in Unit I.C.2. These electronic submissions will be accepted in WordPerfect or ASCII file format. Avoid the use of special characters and any form of encryption.

2. *By mail.* Send your comments to: Public Information and Records Integrity Branch (PIRIB) (7502C), Office of Pesticide Programs (OPP), Environmental Protection Agency, 1200 Pennsylvania Ave., NW., Washington, DC 20460-0001, Attention: Docket ID number OPP-2003-0212.

3. *By hand delivery or courier.* Deliver your comments to: Public Information and Records Integrity Branch (PIRIB), Office of Pesticide Programs (OPP), Environmental Protection Agency, Rm. 119, Crystal Mall #2, 1921 Jefferson Davis Hwy., Arlington, VA, Attention: Docket ID number OPP-2003-0212. Such deliveries are only accepted during the docket's normal hours of operation as identified in Unit I.B.1.

D. How Should I Submit CBI to The Agency?

Do not submit information that you consider to be CBI electronically through EPA's electronic public docket or by e-mail. You may claim information that you submit to EPA as CBI by marking any part or all of that information as CBI (if you submit CBI on disk or CD ROM, mark the outside of the disk or CD ROM as CBI and then identify electronically within the disk or CD ROM the specific information that is CBI). Information so marked will not be

disclosed except in accordance with procedures set forth in 40 CFR part 2.

In addition to one complete version of the comment that includes any information claimed as CBI, a copy of the comment that does not contain the information claimed as CBI must be submitted for inclusion in the public docket and EPA's electronic public docket. If you submit the copy that does not contain CBI on disk or CD ROM, mark the outside of the disk or CD ROM clearly that it does not contain CBI. Information not marked as CBI will be included in the public docket and EPA's electronic public docket without prior notice. If you have any questions about CBI or the procedures for claiming CBI, please consult the person listed under **FOR FURTHER INFORMATION CONTACT**.

E. What Should I Consider as I Prepare My Comments for EPA?

You may find the following suggestions helpful for preparing your comments:

1. Explain your views as clearly as possible.
2. Describe any assumptions that you used.
3. Provide copies of any technical information and/or data you used that support your views.
4. If you estimate potential burden or costs, explain how you arrived at the estimate that you provide.
5. Provide specific examples to illustrate your concerns.
6. Make sure to submit your comments by the deadline in this notice.
7. To ensure proper receipt by EPA, be sure to identify the docket ID number assigned to this action in the subject line on the first page of your response. You may also provide the name, date, and **Federal Register** citation.

II. What Action is the Agency Taking?

EPA has received a pesticide petition as follows proposing the establishment and/or amendment of regulations for residues of a certain pesticide chemical in or on various food commodities under section 408 of the Federal Food, Drug, and Cosmetic Act (FFDCA), 21 U.S.C. 346a. EPA has determined that this petition contains data or information regarding the elements set forth in FFDCA section 408(d)(2); however, EPA has not fully evaluated the sufficiency of the submitted data at this time or whether the data support granting of the petition. Additional data may be needed before EPA rules on the petition.

List of Subjects

Environmental protection, Agricultural commodities, Feed additives, Food additives, Pesticides and pests, Reporting and recordkeeping requirements.

Dated: June 19, 2003.

Peter Caulkins,

Director, Registration Division, Office of Pesticide Programs.

Summary of Petition

The petitioner's summary of the pesticide petition is printed below as required by FFDCA section 408(d)(3). The summary of the petition was prepared by E. I. Du Pont de Nemours and Company, Du Pont Crop Protection, and represents the view of the petitioner. The petition summary announces the availability of a description of the analytical methods available to EPA for the detection and measurement of the pesticide chemical residues or an explanation of why no such method is needed.

E. I. Du Pont de Nemours and Company Du Pont Crop Protection

PP 3F6576

EPA has received a pesticide petition (PP 3F6576) from E. I. Du Pont de Nemours and Company, Du Pont Crop Protection, Wilmington, Delaware, proposing pursuant to section 408(d) of the Federal Food, Drug and Cosmetic Act, 21 U.S.C. 346a(d), to amend 40 CFR part 180, by establishing a tolerance for combined residues of Indoxacarb, [(S)-methyl 7-chloro-2,5-dihydro-2-[(methoxycarbonyl)[4-(trifluoromethoxy)phenyl]amino]carbonyl]indeno[1,2-e] [1,3,4]oxadiazine-4a(3H)-carboxylate] and its R-enantiomer [(R)-methyl 7-chloro-2,5-dihydro-2-[(methoxycarbonyl)[4-(trifluoromethoxy)phenyl]amino]carbonyl]indeno[1,2-e] [1,3,4]oxadiazine-4a(3H)-carboxylate] in a 75:25 mixture (DPX MP062), respectively, in or on the raw agricultural commodity as follows: grape, 2 ppm and raisin, 6 ppm. An analytical enforcement method (LC-UV) is available for determining plant residues. EPA has determined that the petition contains data or information regarding the elements set forth in section 408(d)(2) of the FFDCA; however, EPA has not fully evaluated the sufficiency of the submitted data at this time or whether the data support granting of the petition. Additional data may be needed before EPA rules on the petition.

A. Residue Chemistry

The active ingredient in the end-use formulation, Avaunt[®], is a 75:25 mixture of two isomers, indoxacarb (DPX-KN128) and IN-KN127. Only one of the isomers, indoxacarb (DPX-KN128), has insecticidal activity. Since the insecticidal efficacy is based on the concentration of indoxacarb (DPX-KN128), the application rates have been normalized on an indoxacarb (DPX-KN128) basis. The proposed tolerance expression includes both indoxacarb (DPX-KN128) and IN-KN127 and the residue method does not distinguish between the enantiomers, therefore, residues are reported as the sum of indoxacarb (DPX-KN128) combined with IN-KN127. Residues of indoxacarb (DPX-KN128) combined with IN-KN127 will be referred to as "KN128/KN127".

1. *Plant metabolism* The metabolism of indoxacarb in plants is adequately understood to support these tolerances. Plant metabolism studies in cotton, lettuce, and tomatoes showed no significant metabolites. The only significant residue was parent compound.

2. *Analytical method.* The plant residue enforcement method detects and quantitates indoxacarb in various matrices including sweet corn, lettuce, tomato, broccoli, apple, grape, cottonseed, tomato, peanut and soybean commodity samples by HPLC-UV. The limit of quantitation in the method allows monitoring of crops with indoxacarb residues at or above the levels proposed in these tolerances.

3. *Magnitude of residues—*a. Grapes.** Residue studies were conducted at a total of 13 field sites. All studies were done using Avaunt[®] Insecticide containing 30% active ingredient (300 grams (g) DPX-KN128 per kilogram (kg), weight/weight (w/w). Four broadcast applications of Avaunt[®] Insecticide were made at each test site at a maximum rate of 0.11 lb. active ingredient (a.i.) DPX-KN128/acre/application (maximum seasonal use rate of 0.44 lb DPX-KN128/acre). Applications were made approximately 5 days apart. Residues were measured as the combination of DPX-KN128 and IN-KN127 (enantiomers not resolved by the analytical method). Maximum residues of KN128/KN127 in individual duplicate samples were 1.72 parts per million (ppm) at a pre-harvest interval (PHI) of 7 days (range 0.066 to 1.72 ppm).

b. *Grape processing.* A grape processing study was also performed in California. *Grapes* received four applications of an exaggerated rate of

0.555 lb. of indoxacarb per acre, 5X the labeled rate. Samples were collected from control and treated plots 7 days after the last application. The *grape* samples were then processed using standard *grape* processing procedures.

Samples collected at the processing facility were whole fruit, raisins and *grape* juice. The mean KN128/KN127 residues in whole fruit, raisins and *grape* juice treated with the exaggerated

rate were 1.34 ppm, 3.66 ppm and <0.01 ppm, respectively.

B. Toxicological Profile

1. *Acute toxicity*. Based on EPA criteria, indoxacarb is classified as follows for toxicity categories:

Guideline	Title	Results	Category
81-1	Acute oral toxicity	LD ₅₀ : 1,730 milligrams/kilogram (mg/kg) (M Rat) LD ₅₀ : 268 mg/kg (F Rat)	Category II
81-2	Acute dermal toxicity	LD ₅₀ : >5,000 mg/kg (Rat)	Category IV
81-3	Acute inhalation toxicity	LC ₅₀ : >5.5 (milligrams/per Liter (mg/L) (M Rat) (70% MUP)	Category IV
81-4	Primary eye irritation	Effects reversed within 72 hours (Rabbit)	Category III
81-5	Primary dermal irritation	No irritation (Rabbit)	Category IV
81-6	Skin sensitization	Sensitizer (Guinea Pig)	

Formulated products are slightly less acutely toxic than indoxacarb.

In an acute neurotoxicity study, indoxacarb exhibited decreased forelimb grip strength, decreased foot splay, and some evidence of slightly reduced motor activity, but only at the highest doses tested. The no adverse effects level (NOAEL) was 100 mg/kg for males and 12.5 mg/kg for females based on body weight effects in females >50 mg/kg.

2. *Genotoxicity*. Indoxacarb has shown no genotoxic activity in the following listed *in-vitro* and *in-vivo* tests:

- i. Ames—Negative
- ii. *In-vitro* mammalian gene mutation (CHO/HGPRT)—Negative
- iii. *In-vitro* unscheduled DNA synthesis—Negative
- iv. *In-vitro* chromosomal aberration—Negative
- v. *In-vivo* mouse micronucleus—Negative

3. *Reproductive and developmental toxicity*. The results of a series of studies indicated that there were no reproductive, developmental or teratogenic hazards associated with the use of indoxacarb. In a 2-generation rat reproduction study, the parental NOAEL was 1.5 mg/kg/day. The parental NOAEL was based on observations of reduced weight gain and food consumption for the higher concentration groups of the F0 generation, and potential treatment-related changes in spleen weights for the higher groups of the F1 generation. There was no effect on mating or fertility. The NOAEL for fertility and reproduction was 6.4 mg/kg/day. The offspring NOAEL was 1.5 mg/kg/day, and was based on the reduced mean

pup weights noted for the F1 litters of the higher concentration groups. The effects on pup weights occurred only at a maternal effect level and may have been due to altered growth and nutrition in the dams. In studies conducted to evaluate developmental toxicity potential, indoxacarb was neither teratogenic nor uniquely toxic to the conceptus (i.e., not considered a developmental toxin). Developmental studies conducted in rats and rabbits demonstrated that the rat was more susceptible than the rabbit to the maternal and fetal effects of DPX-MP062. Developmental toxicity was observed only in the presence of maternal toxicity. The NOAEL for maternal and fetal effects in rats was 2 mg/kg/day based on body weight effects and decreased food consumption at 4 mg/kg/day. The NOAEL for developmental effects in fetuses was >4 mg/kg/day. In rabbits, the maternal and fetal NOAELs were 500 mg/kg/day based on body weight effects, decreased food consumption in dams and decreased weight and delayed ossification in fetuses at 1,000 mg/kg/day.

4. *Subchronic toxicity*. Subchronic (90-day) feeding studies were conducted with rats, mice, and dogs. In a 90-day feeding study in rats, the NOAEL was 3.1 and 2.1 mg/kg/day for males and females, respectively. In male rats, the NOAEL was based on decreased body weight and nutritional parameters, mild hemolytic anemia and decreased total protein and globulin concentration. In female rats, the NOAEL was based on decreased body weight and food efficiency.

In a subchronic neurotoxicity study in rats, there was no evidence of neurotoxicity at 11.9 and 6.09 mg/kg/day, the highest dose tested for males and females, respectively. The subchronic NOAEL in dogs (5.0 mg/kg/day, M/F) was based on hemolytic anemia. Erythrocyte values for most dogs were within a range that would be considered normal for dogs in a clinical setting. Mice were less sensitive to indoxacarb than the rats or dogs. NOAELs (23 mg/kg/day, males, 16 mg/kg/day, females) were based on mortality (males only); increased reticulocytes and Heinz bodies and decreased body weight, weight gain, food consumption, food efficiency; and increased clinical signs (leaning to one side and/or with abnormal gait or mobility) (females only). In a 28-day repeated dose dermal study, the NOAEL was 50 mg/kg/day based on decreased body weights, body weight gains, food consumption, and food efficiency in females, and changes in hematology parameters, the spleen and clinical signs of toxicity in both sexes in rats.

5. *Chronic toxicity*. Chronic studies with indoxacarb were conducted on rats, mice, and dogs to determine oncogenic potential and/or chronic toxicity of the compound. Effects generally similar to those observed in the 90-day studies were seen in the chronic studies. Indoxacarb was not oncogenic in rats or mice. The chronic NOAEL in male rats was 5 mg/kg/day based on body weight and nutritional effects. In females, the NOAEL of 2.1 mg/kg/day was based on body weight and nutritional changes, as well as biologically significant hematologic changes at 3.6 mg/kg/day and above.

Hemolytic effects were present only through the 6-month evaluation and only in females. The regenerative nature of indoxacarb-induced hemolytic anemia was demonstrated by the absence of significant changes in indicators of circulating erythrocyte mass at later evaluations. In mice, the chronic NOAEL of 2.6 mg/kg/day for males was based on deceased body weight and weight gain effects and food efficiency at 13.8 mg/kg/day and above. The NOAEL for females was 4.0 mg/kg/day based on body weight nutritional effects, neurotoxicity, and clinical signs at 20 mg/kg/day. In dogs, the chronic NOAEL was about 2.3 and 2.4 mg/kg/day in males and females, respectively based on hemolytic effects similar to those seen in the subchronic dog study.

6. *Animal metabolism—i. Livestock animal metabolism.* Animal metabolism has been studied in the rat, hen, and cow and is well understood. In contrast to crops, indoxacarb is extensively metabolized in animals.

ii. *Poultry.* In poultry, hens were fed at 10 ppm/day for 5 days, 87–88% of the total administered dose was excreted; parent comprised 51–54% of the total dose in excreta. Concentration of residues in eggs were low, 0.3–0.4 of the total dose, as was the concentration of residues in muscle, 0.2% of the total dose. Parent and metabolite IN-JT333 were not detected in egg whites; only insecticidally inactive metabolites were identified. Parent and IN-JT333 were found in egg yolks; however, their concentrations were very low—0.01–0.02 ppm. Concentrations of parent and IN-JT333 in muscle were at or below the limit of quantitation, (LOQ) (0.01 ppm).

iii. *Cattle. For the cow study,* the cattle were fed at 10 ppm/day for 5 days; approximately 20% of the total administered dose was excreted in urine and 53–60% was excreted in feces in 5 days. Four-tenths to 1.2% of the total dose in urine was parent indicating extensive metabolism; parent represented 46–68% of the fecal activity. Thus, most residues were not absorbed; those residues that were absorbed were extensively metabolized. Less than 1% of the total administered dose was in milk, most of which was parent compound. The insecticidally active metabolite IN-JT333 was not found in milk. Residues in muscle represented less than 0.01% of the total administered dose most of which was parent. IN-JT333 was not detected in muscle. No other metabolites were seen above 10% of the dose, thus only parent and IN-JT333 were monitored in the cattle feeding study.

iv. *Cattle feeding study.* A cattle feeding study was conducted with

indoxacarb at doses of 7.5 ppm, 22.5 and 75 ppm. The mean KN128/KN127 concentrations were proportional to the dosing level in whole milk, skim milk, cream, muscle, fat, liver and kidney. Based on final residue values for the respective commodities contributing to the cattle diet, the anticipated dietary burden in dairy cattle is 51.7 ppm and the anticipated dietary burden in beef cattle is 49.1 ppm. The proposed grape use will not increase the animal dietary burden. Based on standard curves constructed from data in the cattle feeding study, KN128/KN127 concentrations at the 51.7 ppm feeding level are 0.123 ppm for whole milk, 0.033 ppm for skim milk and 1.46 ppm for cream. The KN128/KN127 concentrations at the 49.1 ppm feeding level are 0.046 ppm for muscle, 1.37 ppm for fat, 0.012 ppm for liver and 0.026 ppm for kidney. Tolerances have been established at 1.5 ppm in fat (cattle, goat, horse, sheep and hog), 0.05 ppm in meat, 0.03 ppm in meat by-products, 0.15 ppm in milk and 4.0 ppm in milk fat.

7. *Metabolite toxicology.* In rats, indoxacarb was readily absorbed at low dose (5 mg/kg), but saturated at the high dose (150 mg/kg). Indoxacarb was metabolized extensively, based on very low excretion of parent compound in bile and extensive excretion of metabolized dose in the urine and feces. Some parent compound remained unabsorbed and was excreted in the feces. No parent compound was excreted in the urine. The retention and elimination of the metabolite IN-JT333 from fat appeared to be the overall rate determining process for elimination of radioactive residues from the body. Metabolites in urine were cleaved products (containing only one radiolabel), while the major metabolites in the feces retained both radiolabels. Major metabolic reactions included hydroxylation of the indanone ring, hydrolysis of the carboxylmethyl group from the amino nitrogen and the opening of the oxadiazine ring, which gave rise to cleaved products. Metabolites were identified by mass spectral analysis, NMR, ultraviolet (UV), and/or by comparison to standards chemically synthesized or produced by microsomal enzymes.

8. *Endocrine disruption. Lifespan, and multigenerational bioassays in mammals, and acute and subchronic studies* on aquatic organisms and wildlife did not reveal endocrine effects. Any endocrine related effects would have been detected in this definitive array of required tests. The probability of any such effect due to agricultural uses of indoxacarb is negligible.

C. Aggregate Exposure

Tolerances for indoxacarb are proposed to support agricultural use on grapes. There are residential uses of indoxacarb pending (fire ant bait), however, the risk from that use has been found to be negligible.

1. *Dietary exposure.* The chronic reference dose (RfD) of 0.02 mg/kg bwt/day is based on a NOAEL of 2.0 mg/kg bwt/day from the subchronic rat feeding study, the subchronic rat neurotoxicity study, and the chronic/carcinogenicity study, using an uncertainty factor of 100. The acute RfD for the general population is 0.12 mg/kg/day, based on the NOAEL of 12.5 mg/kg in the acute neurotoxicity study and an uncertainty factor of 100. The acute RfD for females 13–50 years of age is 0.02 mg/kg/day, based on the NOAEL of 2 mg/kg/day observed in the developmental rat toxicity study and using an uncertainty factor of 100.

i. *Food. Chronic dietary exposure assessment.* Chronic dietary exposure resulting from the currently approved use of indoxacarb on apples, Crop group 5 (brassica vegetables), cotton, pears, peppers, sweet corn, tomatoes, eggplant, alfalfa, head and leaf lettuce, peanuts, potatoes, soybeans, cranberries (current Section 18 use) and the proposed use on grapes are well within acceptable limits for all sectors of the population. The Chronic Module of the Dietary Exposure Evaluation Model (DEEM™, Exponent, Inc., formerly Novigen Sciences, Inc., Version 7.76) was used to conduct the assessment with the reference dose (RfD) of 0.02 mg/kg/day. The analysis used overall mean field trial values, processing factors and projected peak percent crop treated values. Secondary residues in milk, meat and poultry products were also included in the analysis. The chronic dietary exposure to indoxacarb is 0.000089 mg/kg/day, and utilizes 0.4% of the RfD for the overall U.S. population. The exposure of the most highly exposed subgroup in the population, children age 1–6 years, is 0.000238 mg/kg/day, and utilizes 1.2% of the RfD. The table below lists the results of this analysis, which indicate large margins of safety for each population subgroup and very low probability of effects resulting from chronic exposure to indoxacarb.

Subgroup	Maximum Dietary Exposure (mg/kg/day)	% RfD
U.S. Population	0.000089	0.4

Subgroup	Maximum Dietary Exposure (mg/kg/day)	% RfD
Non-Nursing Infants (<1 year old)	0.000063	0.3
Children (1–6 years)	0.000238	1.2
Children (7–12 years)	0.000126	0.6
Females (13+, nursing)	0.000073	0.4
Males (13–19 years)	0.000090	0.5

ii. *Acute dietary exposure.* Acute dietary exposure resulting from the currently approved use of indoxacarb on apples, Crop Group 5 (brassica vegetables), cotton, pears, peppers, sweet corn, tomatoes, eggplant, alfalfa, head and leaf lettuce, peanuts, soybeans, potatoes, cranberries (current Section 18 use) and the proposed use on grapes are well within acceptable limits for all sectors of the population. The Dietary Exposure Evaluation Model (DEEM™, Exponent, Inc., formerly Novigen Sciences, Inc., Version 7.76) was used to conduct the assessment. Margins of exposure (MOE) were calculated based on an acute NOAEL of 2 mg/kg/day for women of childbearing age and a NOAEL of 12 mg/kg/day for children and the general population (Pesticide Fact Sheet for Indoxacarb). The Tier 3 analysis used distributions of field trial residue data adjusted for projected peak percent crop treated. Secondary residues in milk, meat and poultry products were also included in the analysis. The results of this analysis are given in the table below. The percent of the acute population adjusted dose (a PAD) for all population subgroups shows that an adequate margin of safety exists in each case. Thus, the acute dietary safety of indoxacarb for established and the follow-on use clearly meets the Food Quality Protection Act (FQPA) standard of reasonable certainty of no harm and presents acceptable acute dietary risk.

Subgroup	99.9 th Percentile of exposure	
	Exposure (mg/kg/day)	% Acute population adjusted dose (aPAD)
U.S. population	0.008795	7.3

Subgroup	99.9 th Percentile of exposure	
	Exposure (mg/kg/day)	% Acute population adjusted dose (aPAD)
All infants	0.024729	20.6
Non-nursing (<1 year)	0.026036	21.7
Children (1–6 years)	0.013973	11.6
Children (7–12 years)	0.006882	5.7
Females (13–19 years)	0.005119	25.6
Females (20+, not pregnant or nursing)	0.005358	26.8
Females (13–50 years)	0.005307	26.5

iii. *Drinking water.* Indoxacarb is highly unlikely to contaminate groundwater resources due to its immobility in soil, low water solubility, high soil sorption, and moderate soil half-life. Based on the PRZM/EXAMS and SCI-GROW models the estimated environmental concentrations (EECs) of indoxacarb and its R-enantiomer for acute exposures are estimated to be 6.84 parts per billion (ppb) for surface water and 0.0025 ppb for ground water. The EECs for chronic exposures are estimated to be 0.316 ppb for surface water and 0.0025 ppb for ground water. Drinking water levels of comparisons (DWLOCs), theoretical upper allowable limits on the pesticide's concentration in drinking water, were calculated to be much higher than the EEC's. The chronic DWLOC's ranged from 198 to 697 ppb. The acute DWLOC's ranged from 440 to 3,890 ppb. Thus, exposure via drinking water is acceptable.

2. *Non-dietary exposure.* Indoxacarb product registrations for residential non-food uses are pending. Non-occupational, non-dietary exposure for DPX-MP062 has been estimated to be extremely small. Therefore, the potential for non-dietary exposure is insignificant.

D. Cumulative Effects

EPA's consideration of a common mechanism of toxicity is not necessary at this time because there is no indication that toxic effects of indoxacarb would be cumulative with those of any other chemical compounds. Oxadiazine chemistry is new, and

indoxacarb has a novel mode of action compared to currently registered active ingredients.

E. Safety Determination

1. *U.S. population.* Dietary and occupational exposure will be the major routes of exposure to the U.S. population, and ample margins of safety have been demonstrated for both situations. The chronic dietary exposure to indoxacarb is 0.000089 mg/kg/day, which utilizes 0.4% of the RfD for the overall U.S. population, using mean field trial values, processing factors and projected peak percent crop treated values. The percent of the acute population adjusted dose (7.3% aPAD) for the overall U.S. population shows that an adequate margin of safety exists. Using only pesticide handlers exposure data base (PHED) data levels A and B (those with a high level of confidence), margin of exposure (MOEs) for occupational exposure are 650 for mixer/loaders and 1,351 for airblast applicators (worst-case). Based on the completeness and reliability of the toxicity data and the conservative exposure assessments, there is a reasonable certainty that no harm will result from the aggregate exposure of residues of indoxacarb including all anticipated dietary exposure and all other non-occupational exposures.

2. *Infants and children.* Chronic dietary exposure of the most highly exposed subgroup in the population, children age 1–6 years, is 0.000238 mg/kg/day or 1.2% of the RfD. For infants (non-nursing, <1 yr.), the exposure accounts for 0.3% of the RfD. For acute exposure at the 99.9th percentile (based on a Tier 3 assessment) the exposure was 0.013973 mg/kg/day (11.6% a PAD) for children 1–6 and 0.026036 mg/kg/day (21.7% a PAD) for non-nursing infants. There are residential uses of indoxa carb pending, but exposure is calculated to be extremely minimal. The estimated levels of indoxa carb in drinking water are well below the DWLOC. Based on the completeness and reliability of the toxicity data, the lack of toxicological endpoints of special concern, the lack of any indication that children are more sensitive than adults to indoxa carb, and the conservative exposure assessment, there is a reasonable certainty that no harm will result to infants and children from the aggregate exposure of residues of indoxa carb, including all anticipated dietary exposure and all other non-occupational exposures. Accordingly, there is no need to apply an additional safety factor for infants and children.

F. International Tolerances

To date, no international tolerances exist for indoxacarb.

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ENVIRONMENTAL PROTECTION AGENCY

[OPP-2003-0211; FRL-7312-8]

Dinotefuran; Notice of Filing a Pesticide Petition to Establish a Tolerance for a Certain Pesticide Chemical in or on Food

AGENCY: Environmental Protection Agency (EPA).

ACTION: Notice.

SUMMARY: This notice announces the initial filing of a pesticide petition proposing the establishment of regulations for residues of a certain pesticide chemical in or on various food commodities.

DATES: Comments, identified by docket ID number OPP-2003-0211, must be received on or before August 1, 2003.

ADDRESSES: Comments may be submitted electronically, by mail, or through hand delivery/courier. Follow the detailed instructions as provided in Unit I. of the **SUPPLEMENTARY INFORMATION**.

FOR FURTHER INFORMATION CONTACT: Rita Kumar, Registration Division (7505C), Office of Pesticide Programs, Environmental Protection Agency, 1200 Pennsylvania Ave., NW., Washington, DC 20460-0001; telephone number: (703) 308-8291; e-mail address: kumar.rita@epa.gov.

SUPPLEMENTARY INFORMATION:

I. General Information

A. Does this Action Apply to Me?

You may be potentially affected by this action if you are an agricultural producer, food manufacturer, or pesticide manufacturer. Potentially affected entities may include, but are not limited to:

- Crop production (NAICS 111)
- Animal production (NAICS 112)
- Food manufacturing (NAICS 311)
- Pesticide manufacturing (NAICS 28522)

This listing is not intended to be exhaustive, but rather provides a guide for readers regarding entities likely to be affected by this action. Other types of entities not listed in this unit could also be affected. The North American Industrial Classification System (NAICS) codes have been provided to assist you and others in determining

whether this action might apply to certain entities. If you have any questions regarding the applicability of this action to a particular entity, consult the person listed under **FOR FURTHER INFORMATION CONTACT**.

B. How Can I Get Copies of This Document and Other Related Information?

1. *Docket.* EPA has established an official public docket for this action under docket identification (ID) number OPP-2003-0211. The official public docket consists of the documents specifically referenced in this action, any public comments received, and other information related to this action. Although a part of the official docket, the public docket does not include Confidential Business Information (CBI) or other information whose disclosure is restricted by statute. The official public docket is the collection of materials that is available for public viewing at the Public Information and Records Integrity Branch (PIRIB), Rm. 119, Crystal Mall #2, 1921 Jefferson Davis Hwy., Arlington, VA. This docket facility is open from 8:30 a.m. to 4 p.m., Monday through Friday, excluding legal holidays. The docket telephone number is (703) 305-5805.

2. *Electronic access.* You may access this **Federal Register** document electronically through the EPA Internet under the “**Federal Register**” listings at <http://www.epa.gov/fedrgstr/>.

An electronic version of the public docket is available through EPA’s electronic public docket and comment system, EPA Dockets. You may use EPA Dockets at <http://www.epa.gov/edocket/> to submit or view public comments, access the index listing of the contents of the official public docket, and to access those documents in the public docket that are available electronically. Although not all docket materials may be available electronically, you may still access any of the publicly available docket materials through the docket facility identified in Unit I.B.1. Once in the system, select “search,” then key in the appropriate docket ID number.

Certain types of information will not be placed in EPA’s Dockets. Information claimed as CBI and other information whose disclosure is restricted by statute, which is not included in the official public docket, will not be available for public viewing in EPA’s electronic public docket. EPA’s policy is that copyrighted material will not be placed in EPA’s electronic public docket but will be available only in printed, paper form in the official public docket. To the extent feasible, publicly available docket materials will be made available

in EPA’s electronic public docket. When a document is selected from the index list in EPA Dockets, the system will identify whether the document is available for viewing in EPA’s electronic public docket. Although not all docket materials may be available electronically, you may still access any of the publicly available docket materials through the docket facility identified in Unit I.B. EPA intends to work towards providing electronic access to all of the publicly available docket materials through EPA’s electronic public docket.

For public commenters, it is important to note that EPA’s policy is that public comments, whether submitted electronically or in paper, will be made available for public viewing in EPA’s electronic public docket as EPA receives them and without change, unless the comment contains copyrighted material, CBI, or other information whose disclosure is restricted by statute. When EPA identifies a comment containing copyrighted material, EPA will provide a reference to that material in the version of the comment that is placed in EPA’s electronic public docket. The entire printed comment, including the copyrighted material, will be available in the public docket.

Public comments submitted on computer disks that are mailed or delivered to the docket will be transferred to EPA’s electronic public docket. Public comments that are mailed or delivered to the docket will be scanned and placed in EPA’s electronic public docket. Where practical, physical objects will be photographed, and the photograph will be placed in EPA’s electronic public docket along with a brief description written by the docket staff.

C. How and to Whom Do I Submit Comments?

You may submit comments electronically, by mail, or through hand delivery/courier. To ensure proper receipt by EPA, identify the appropriate docket ID number in the subject line on the first page of your comment. Please ensure that your comments are submitted within the specified comment period. Comments received after the close of the comment period will be marked “late.” EPA is not required to consider these late comments. If you wish to submit CBI or information that is otherwise protected by statute, please follow the instructions in Unit I.D. Do not use EPA Dockets or e-mail to submit CBI or information protected by statute.

1. *Electronically.* If you submit an electronic comment as prescribed in this