FEDERAL REGISTER

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Part XV

Environmental Protection Agency

Unified Agenda
I. Introduction

EPA is committed to a regulatory strategy that effectively achieves the Agency’s mission of protecting the environment and the health, welfare, and safety of Americans while also supporting economic growth, job creation, competitiveness, and innovation. EPA publishes the Semiannual Agenda of Regulatory and Deregulatory Actions to update the public about regulatory activity undertaken in support of this mission. In the Semiannual Agenda, EPA provides notice of our plans to review, propose, and issue regulations.

Additionally, EPA’s Semiannual Agenda includes information about rules that may have a significant economic impact on a substantial number of small entities, and review of those regulations under the Regulatory Flexibility Act, as amended.

In this document, EPA explains in greater detail the types of actions and information available in the Semiannual Agenda and actions that are currently undergoing review specifically for impacts on small entities.

A. EPA’s Regulatory Information

“E-Agenda,” “online regulatory agenda,” and “semiannual regulatory agenda” all refer to the same comprehensive collection of information that, until 2007, was published in the Federal Register. Currently, this information is only available through an online database, at both www.reginfo.gov and www.regulations.gov.

“Regulatory Flexibility Agenda” refers to a document that contains information about regulations that may have a significant impact on a substantial number of small entities. We continue to publish this document in the Federal Register pursuant to the Regulatory Flexibility Act of 1980. This document is available at https://www.govinfo.gov/app/collection/fr.

“Unified Regulatory Agenda” refers to the collection of all agencies’ agendas with an introduction prepared by the Regulatory Information Service Center facilitated by the General Service Administration.

“Regulatory Agenda Preamble” refers to the document you are reading now. It appears as part of the Regulatory Flexibility Agenda and introduces both EPA’s Regulatory Flexibility Agenda and the e-Agenda.

“610 Review” as required by the Regulatory Flexibility Act means a periodic, ten-year review of promulgating a final rule that has or may have a significant economic impact on a substantial number of small entities. EPA maintains a list of these actions at https://www.epa.gov/reg-flex/section-610-reviews. EPA is initiating one 610 review in spring 2019.

B. What key statutes and Executive Orders guide EPA’s rule and policymaking process?

A number of environmental laws authorize EPA’s actions, including but not limited to:
• Clean Air Act (CAA),
• Clean Water Act (CWA),
• Comprehensive Environmental Response, Compensation, and Liability Act (CERCLA, or Superfund),
• Emergency Planning and Community Right-to-Know Act (EPCRA),
• Federal Insecticide, Fungicide, and Rodenticide Act (FIFRA),
• Resource Conservation and Recovery Act (RCRA),
• Safe Drinking Water Act (SDWA), and
• Toxic Substances Control Act (TSCA).

Not only must EPA comply with environmental laws, but also administrative legal requirements that apply to the issuance of regulations, such as: The Administrative Procedure Act (APA), the Regulatory Flexibility Act (RFA) as amended by the Small Business Regulatory Enforcement Fairness Act (SBREFA), the Unfunded Mandates Reform Act (UMRA), the Paperwork Reduction Act (PRA), the National Technology Transfer and Advancement Act (NTTAA), and the Congressional Review Act (CRA).


C. How can you be involved in EPA’s rule and policymaking process?

You can make your voice heard by getting in touch with the contact person provided in each agenda entry. EPA encourages you to participate as early in the process as possible. You may also
participate by commenting on proposed rules published in the Federal Register
(FR).

Instructions on how to submit your comments through https://
www.regulations.gov are provided in each Notice of Proposed Rulemaking
(NPRM). To be most effective, comments should contain information
and data that support your position and you also should explain why EPA
should incorporate your suggestion in the rule or other type of action. You
can be particularly helpful and persuasive if you provide examples to illustrate your
concerns and offer specific alternative(s) to that proposed by EPA.

EPA believes its actions will be more
cost effective and protective if the
development process includes
stakeholders working with us to help
identify the most practical and effective
solutions to environmental problems.

EPA encourages you to become involved
in its rule and policymaking process.

For more information about EPA’s
efforts to increase transparency,
participation and collaboration in EPA
activities, please visit https://
www.epa.gov/open.

II. Semiannual Agenda of Regulatory
and Deregulatory Actions

A. What actions are included in the e-
Agenda and the Regulatory Flexibility
Agenda?

EPA includes regulations in the e-
Agenda. However, there is no legal
significance to the omission of an item
from the agenda, and EPA generally
does not include the following
categories of actions:

• Administrative actions such as
delegations of authority, changes of
address, or phone numbers;
• Under the CAA: Revisions to state
implementation plans; equivalent
methods for ambient air quality
monitoring; deletions from the
new source performance standards source
categories list; delegations of authority
to states; area designations for air
quality planning purposes;
• Under FIFRA: Registration-related
decisions, actions affecting the status
of currently registered pesticides, and data
–call-ins;
• Under the Federal Food, Drug, and
Cosmetic Act: Actions regarding
pesticide tolerances and food additive
regulations;
• Under TSCA: Actions involving
premanufacture notices and follow-up
activities for new chemical substances
and significant new uses, including
section 5(e) Prioritization and specific
exemptions under sections 5(b)(4) and
26(c); and actions related to

prioritization and risk evaluations for
individual or categories of existing
chemical substances under section 6;
• Under RCRA: Authorization of State
solid waste management plans;
• Under the CWA: State Water
Quality Standards; deletions from the
section 307(a) list of toxic pollutants;
suspensions of toxic testing
requirements under the National
Pollutant Discharge Elimination System
(NPDES); delegations of NPDES
authority to States;
• Under SDWA: Actions on State
underground injection control
programs.

Meanwhile, the Regulatory Flexibility
Agenda includes:
• Actions likely to have a significant
economic impact on a substantial
number of small entities.
• Rules the Agency has identified for
periodic review under section 610 of the
RFA.
• EPA is initiating one 610 review in this
Agenda.

B. How is the e-Agenda organized?

Online, you can choose how to sort
the agenda entries by specifying the
characteristics of the entries of interest
in the desired individual data fields for
both the www.reginfo.gov and
www.regulations.gov versions of the e-
Agenda. You can sort based on the
following characteristics: EPA
subagency (such as Office of Water);
stage of rulemaking as described in the
following paragraphs; alphabetically by
title; or the Regulation Identifier
Number (RIN), which is assigned
sequentially when an action is added to
the agenda.

Each entry in the Agenda is associated
with one of five rulemaking stages. The
rulemaking stages are:

1. Prerule Stage—EPA’s prerule
actions generally are intended to
determine whether the agency should
initiate rulemaking. Prerulemakings
may include anything that influences or
leads to rulemaking; this would include
Advance Notices of Proposed
Rulemaking (ANPRMs), studies or
analyses of the possible need for
regulatory action.

2. Proposed Rule Stage—Proposed
rulemaking actions include EPA’s
Notice of Proposed Rulemakings
(NPRMs); these proposals are scheduled
to publish in the Federal Register
within the next year.

3. Final Rule Stage—Final rulemaking
actions are those actions that EPA is
scheduled to finalize and publish in the
Federal Register within the next year.

4. Long-Term Actions—This section
includes rulemakings for which the next
scheduled regulatory action (such as
publication of a NPRM or final rule) is
under twelve or more months into the future.

We urge you to explore becoming
involved even if an action is listed in the
Long-Term category.

5. Completed Actions—EPA’s
completed actions are those that have
been promulgated and published in the
Federal Register since publication of
the fall 2018 Agenda. The term
completed actions also includes actions
that EPA is no longer considering and
has elected to “withdraw” and also the
results of any RFA section 610 reviews.

C. What information is in the Regulatory
Flexibility Agenda and the e-Agenda?

The Regulatory Flexibility Agenda
entries include only the nine categories
of information that are required by the
Regulatory Flexibility Act of 1980 and
by Federal Register Agenda printing
requirements: Sequence Number, RIN,
Title, Description, Statutory Authority,
Section 610 Review, if applicable,
Regulatory Flexibility Analysis
Required, Schedule and Contact Person.

Note that the electronic version of the
Agenda (E-Agenda) replicates each of
these actions with more extensive
information, described below.

E-Agenda entries include:

Title: A brief description of the
subject of the regulation. The notation
“Section 610 Review” follows the title
if we are reviewing the rule as part of
our periodic review of existing rules
under section 610 of the RFA (5 U.S.C.
610).

Priority: Each entry is placed into one of
the five following categories:

a. Economically Significant: Under
Executive Order 12866, a rulemaking
that may have an annual effect on the
economy of $100 million or more, or
adversely affect in a material way the
economy, a sector of the economy,
productivity, competition, jobs, the
environment, public health or safety, or
State, local, or tribal governments or
communities.

b. Other Significant: A rulemaking
that is not economically significant but
is considered significant for other
reasons. This category includes rules
that may:

1. Create a serious inconsistency or
otherwise interfere with an action taken
or planned by another agency;

2. Materially alter the budgetary
impact of entitlements, grants, user fees,
or loan programs, or the rights and
obligations of recipients; or

3. Raise novel legal or policy issues
arising out of legal mandates, the
President’s priorities, or the principles
in Executive Order 12866.
Executive Order 13771 Designation:

Each entry is placed into one of the following categories:

- **a. Deregulatory**: When finalized, an action is expected to have total costs less than zero.
- **b. Regulatory**: The action is either
  - (i) a significant regulatory action as defined in section 3(f) of Executive Order 12866, or
  - (ii) a significant guidance document (e.g., significant interpretive guidance) reviewed by OMB’s Office of Information and Regulatory Affairs (OIRA) under the procedures of Executive Order 12866 that, when finalized, is expected to impose total costs greater than zero;
- **c. Fully or Partially Exempt**: The action has been granted, or is expected to be granted, a full or partial waiver under one or more of the following circumstances:
  - (i) It is expressly exempt by Executive Order 13771 (issued with respect to a “military, national security, or foreign affairs function of the United States”; or related to “agency organization, management, or personnel”), or
  - (ii) it addresses an emergency such as critical health, safety, financial, or non-exempt national security matters (offset requirements may be exempted or delayed), or
  - (iii) it is required to meet a statutory or judicial deadline (offset requirements may be exempted or delayed), or
  - (iv) expected to generate de minimis costs.
- **d. Not subject to, not significant**: Is a NPRM or final rule AND is neither an Executive Order 13771 regulatory action nor an Executive Order 13771 deregulatory action.
- **e. Other**: At the time of designation, either the available information is too preliminary to determine E.O. 13771 status or other reasonable circumstances preclude a preliminary Executive Order 13771 designation.
- **f. Independent agency**: Is an action an independent agency anticipates issuing and thus is not subject to Executive Order 13771.

Major: A rule is “major” under 5 U.S.C. 801 (Pub. L. 104–121) if it has resulted or is likely to result in an annual effect on the economy of $100 million or more or meets other criteria specified in that Act.

Unfunded Mandates: Whether the rule is covered by section 202 of the Unfunded Mandates Reform Act of 1995 (Pub. L. 104–4). The Act requires that, before issuing an NPRM likely to result in a mandate that may result in expenditures by State, local, and tribal governments, in the aggregate, or by the private sector of more than $100 million in 1 year, the agency prepare a written statement on federal mandates addressing costs, benefits, and intergovernmental consultation.

Legal Authority: The sections of the United States Code (U.S.C.), Public Law (Pub. L.), Executive Order (E.O.), or common name of the law that authorizes the regulatory action.

Legal Deadline: An indication of whether the rule is subject to a statutory or judicial deadline, the date of that deadline, and whether the deadline pertains to a Notice of Proposed Rulemaking, a Final Action, or some other action.

Abstract: A brief description of the problem the action will address.

Timetable: The dates and citations (if available) for all past steps and a projected date for at least the next step for the regulatory action. A date displayed in the form 05/00/20 means the agency is predicting the month and year the action will take place but not the day it will occur. For some entries, the timetable indicates that the date of the next action is “to be determined.”

Regulatory Flexibility Analysis Required: Indicates whether EPA has prepared or anticipates preparing a regulatory flexibility analysis under section 603 or 604 of the RFA.

Generally, such an analysis is required for proposed or final rules subject to the RFA that EPA believes may have a significant economic impact on a substantial number of small entities.

**Small Entities Affected**: Indicates whether the rule is anticipated to have any effect on small businesses, small governments or small nonprofit organizations.

**Government Levels Affected**: Indicates whether the rule may have any effect on levels of government and, if so, whether the affected governments are State, local, tribal, or Federal.

**Federalism Implications**: Indicates whether the action is expected to have substantial direct effects on the States, on the relationship between the National Government and the States, or on the distribution of power and responsibilities among the various levels of government.

**Energy Impacts**: Indicates whether the action is a significant energy action under Executive Order 13211.

**Sectors Affected**: Indicates the main economic sectors regulated by the action. The regulated parties are identified by their North American Industry Classification System (NAICS) codes. These codes were created by the Census Bureau for collecting, analyzing, and publishing statistical data on the U.S. economy. There are more than 1,000 NAICS codes for sectors in agriculture, mining, manufacturing, services, and public administration.

**International Trade Impacts**: Indicates whether the action is likely to have international trade or investment effects, or otherwise be of international interest.

**Agency Contact**: The name, address, phone number, and email address, if available, of a person who is knowledgeable about the regulation.

**Additional Information**: Other information about the action including docket information.

**URLs**: For some actions, the internet addresses are included for reading copies of rulemaking documents, submitting comments on proposals, and getting more information about the rulemaking and the program of which it is a part. (Note: To submit comments on proposals, you can go to the associated electronic docket, which is housed at www.regulations.gov. Once there, follow the online instructions to access the docket in question and submit comments. A docket identification [ID] number will assist in the search for materials.)

**RIN**: The Regulation Identifier Number is used by OMB to identify and track rulemakings. The first four digits of the RIN identify the EPA office with lead responsibility for developing the action.
D. What tools are available for mining Regulatory Agenda data and for finding more about EPA rules and policies?

1. Federal Regulatory Dashboard
   The [https://www.reginfo.gov/searchable database, maintained by the Regulatory Information Service Center and OIRA](https://www.reginfo.gov/) allows users to view the Regulatory Agenda database ([https://www.reginfo.gov/public/do/eAgendaMain](https://www.reginfo.gov/public/do/eAgendaMain)), which includes search, display, and data transmission options.

2. Subject Matter EPA websites
   Some actions listed in the Agenda include a URL for an EPA-maintained website that provides additional information about the action.

3. Deregulatory Actions and Regulatory Reform
   EPA maintains a list of its deregulatory actions under development, as well as those that are completed, at [https://www.epa.gov/laws-regulations/epa-deregulatory-actions](https://www.epa.gov/laws-regulations/epa-deregulatory-actions). Additional information about EPA’s regulatory reform activity is available to the public at [https://www.epa.gov/laws-regulations/regulatory-reform](https://www.epa.gov/laws-regulations/regulatory-reform).

4. Public Dockets
   When EPA publishes either an Advance Notice of Proposed Rulemaking (ANPRM) or a Notice of Proposed Rulemaking (NPRM) in the Federal Register, the Agency typically establishes a docket to accumulate materials developed throughout the development process for that rulemaking. The docket serves as the repository for the collection of documents or information related to that particular Agency action or activity. EPA most commonly uses dockets for rulemaking actions, but dockets may also be used for RFA section 610 reviews of rules with significant economic impacts on a substantial number of small entities and for various non-rulemaking activities, such as Federal Register documents seeking public comments on draft guidance, policy statements, information collection requests under the PRA, and other non-rule activities. Docket information should be in that action’s agenda entry. All of EPA’s public dockets can be located at [www.regulations.gov](http://www.regulations.gov).

III. Review of Regulations Under 610 of the Regulatory Flexibility Act

A. Reviews of Rules With Significant Impacts on a Substantial Number of Small Entities

Section 610 of the RFA requires that an agency review, within 10 years of promulgation, each rule that has or will have a significant economic impact on a substantial number of small entities. At this time, EPA is initiating one section 610 review.

<table>
<thead>
<tr>
<th>Review title</th>
<th>RIN</th>
<th>Docket ID No.</th>
<th>Status</th>
</tr>
</thead>
</table>

EPA has established an official public docket for this 610 review. Comments received on this 610 review can be submitted at [https://www.regulations.gov/](https://www.regulations.gov/) with docket identification number EPA–HQ–OAR–2019–0168.

B. What other special attention does EPA give to the impacts of rules on small businesses, small governments, and small nonprofit organizations?

For each of EPA’s rulemakings, consideration is given to whether there will be any adverse impact on any small entity. EPA attempts to fit the regulatory requirements, to the extent feasible, to the scale of the businesses, organizations, and governmental jurisdictions subject to the regulation.

IV. Thank You for Collaborating With Us

Finally, we would like to thank those of you who choose to join with us in making progress on the complex issues involved in protecting human health and the environment. Collaborative efforts such as EPA’s open rulemaking process are a valuable tool for addressing the problems we face, and the regulatory agenda is an important part of that process.

Dated: March 11, 2019.
Brittany Bolen, Associate Administrator, Office of Policy.

### 10—PRERULE STAGE

<table>
<thead>
<tr>
<th>Sequence No.</th>
<th>Title</th>
<th>Regulation Identifier No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>173</td>
<td>Section 610 Review of Renewable Fuels Standard Program (Section 610 Review)</td>
<td>2060–AU44</td>
</tr>
</tbody>
</table>

### 35—FINAL RULE STAGE

<table>
<thead>
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<th>Title</th>
<th>Regulation Identifier No.</th>
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<tbody>
<tr>
<td>174</td>
<td>Review of Dust-Lead Hazard Standards and the Definition of Lead-Based Paint</td>
<td>2070–AJ82</td>
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</table>
### ENVIRONMENTAL PROTECTION AGENCY (EPA)

**10**

**Prerule Stage**

**173. • Section 610 Review of Renewable Fuels Standard Program** *(Section 610 Review)*

E.O. 13771 Designation: Not subject to, not significant.

Legal Authority: 5 U.S.C. 610

Abstract: This notice indicates that EPA will review this action pursuant to section 610 of the Regulatory Flexibility Act (5 U.S.C. 610). As part of this review, EPA would consider and solicit comments on the following factors: (1) The continued need for the rule; (2) the nature of complaints or comments received concerning the rule; (3) the complexity of the rule; (4) the extent to which the rule overlaps, duplicates, or conflicts with other Federal, State, or local government rules; and (5) the degree to which the technology, economic conditions or other factors.

**Timetable:**

<table>
<thead>
<tr>
<th>Action</th>
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<tbody>
<tr>
<td>Final Rule .....</td>
<td>05/00/19</td>
<td>75 FR 14669</td>
</tr>
<tr>
<td>Begin Review ...</td>
<td>05/00/19</td>
<td>75 FR 14669</td>
</tr>
</tbody>
</table>

Regulatory Flexibility Analysis Required: No.

Agency Contact: Julia Burch, Environmental Protection Agency, Office of Air and Radiation, 1200 Pennsylvania Ave. NW, Washington, DC 20460, Phone: 202 564–0961, Email: burch.julia@epa.gov.

Jessica Mroz, Environmental Protection Agency, Office of Air and Radiation, 1200 Pennsylvania Ave. NW, Washington, DC 20460, Phone: 202 564–1094, Email: mroz.jessica@epa.gov.

RIN: 2060–AU44

### ENVIRONMENTAL PROTECTION AGENCY (EPA)

**35**

**Final Rule Stage**

**174. Review of Dust-Lead Hazard Standards and the Definition of Lead-Based Paint**


Abstract: Addressing childhood lead exposure is a priority for EPA. As part of EPA’s efforts to reduce childhood lead exposure, EPA evaluated the current dust-lead hazard standards (DLHS) and the definition of lead-based paint (LBP). Based on this evaluation, EPA proposed to change the dust-lead hazard standards from 40 µg/ft² and 250 µg/ft² to 10 µg/ft² and 100 µg/ft² on floors and window sills, respectively. These standards apply to most pre-1978 housing and child-occupied facilities, such as day care centers and kindergarten facilities. In addition, EPA proposed to make no change to the definition of lead-based paint because the Agency currently lacks sufficient information to support such a change. The proposed rule was issued in compliance with the December 27, 2017, decision of the Ninth Circuit, and the subsequent March 26, 2018, order that directed the EPA “to issue a proposed rule within ninety (90) days from the filed date of this order”. EPA is reviewing the comments received and developing a final rule.

**Timetable:**

<table>
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<th>Action</th>
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<td>07/02/18</td>
<td>83 FR 30889</td>
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<td>NPRM Comment</td>
<td>08/16/18</td>
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<tr>
<td>Period End.</td>
<td>05/00/19</td>
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</tr>
<tr>
<td>Final Rule ........</td>
<td>05/00/19</td>
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</table>

Regulatory Flexibility Analysis Required: Yes.

Agency Contact: John Yowell, Environmental Protection Agency, Office of Chemical Safety and Pollution Prevention, Mail Code 7404T, Washington, DC 20460, Phone: 202 564–1213, Email: yowell.john@epa.gov.

Marc Edmonds, Environmental Protection Agency, Office of Chemical Safety and Pollution Prevention, 1200 Pennsylvania Avenue NW, Mail Code 7404T, Washington, DC 20460, Phone: 202 566–0758, Email: edmonds.marc@epa.gov.

RIN: 2070–AJ82

### ENVIRONMENTAL PROTECTION AGENCY (EPA)

**35**

**Long-Term Actions**

**175. N-Methylpyrrolidone; Regulation of Certain Uses Under TSCA Section 6(a)**


Abstract: Section 6(a) of the Toxic Substances Control Act provides authority for EPA to ban or restrict the manufacture (including import), processing, distribution in commerce, and use of chemical substances, as well as any manner or method of disposal. Section 26(l)(4) of TSCA authorizes EPA to issue rules under TSCA section 6 for chemicals listed in the 2014 update to the TSCA Work Plan for Chemical Assessments for which EPA published completed risk assessments prior to June 22, 2016, consistent with the scope of the completed risk assessment. N-methylpyrrolidone (NMP) is used in paint and coating removal in commercial processes and consumer products. In the March 2015 TSCA Work Plan Chemical Risk Assessment for NMP, EPA characterized risks from use of this chemical in paint and coating removal. On January 19, 2017, EPA preliminarily determined that the use of NMP in paint and coating removal poses an unreasonable risk of injury to health.

**RIN: 2070–AK07**
EPA also co-proposed two options for NMP in paint and coating removal. The first co-proposal would prohibit the manufacture, processing, and distribution in commerce of NMP for all consumer and most commercial paint and coating removal and the use of NMP for most commercial paint and coating removal. The second co-proposal would require commercial users of NMP for paint and coating removal to establish a worker protection program and not use paint and coating removal products that contain greater than 35% NMP by weight, with certain exceptions; and require processors of products containing NMP for paint and coating removal to reformulate products such that they do not exceed 35% NMP by weight, to identify gloves that provide effective protection for the formulation, and to provide warnings and instructions on any paint and coating removal products containing NMP. In the final rule for methylene chloride in consumer paint and coating removal (RIN 2070–AK07), EPA explained that the Agency was not finalizing the proposed regulation for NMP as part of that action. NMP use in paint and coating removal will be incorporated into the risk evaluation currently being conducted under TSCA section 6(b).

Timetable:

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<tr>
<th>Action</th>
<th>Date</th>
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<tbody>
<tr>
<td>NPRM</td>
<td>01/17/17</td>
<td>82 FR 7464</td>
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<tr>
<td>Final Rule</td>
<td>To Be Determined</td>
<td></td>
</tr>
</tbody>
</table>

Regulatory Flexibility Analysis

Required: Yes.

Agency Contact: Niva Kramek, Environmental Protection Agency, Office of Chemical Safety and Pollution Prevention, 1200 Pennsylvania Avenue NW, Mail Code 7405M, Washington, DC 20460, Phone: 202 564–4830, Email: kramek.niva@epa.gov.

Joel Wolf, Environmental Protection Agency, Office of Chemical Safety and Pollution Prevention, 1200 Pennsylvania Avenue NW, Mail Code 7405M, Washington, DC 20460, Phone: 202 564–0432, Email: wolf.joel@epa.gov.

RIN: 2070–AK46

ENVIRONMENTAL PROTECTION AGENCY (EPA)

35

Completed Actions

176. Methylene Chloride; Regulation of Paint and Coating Removal for Consumer Use Under TSCA Section 6(a)


Abstract: Section 6(a) of the Toxic Substances Control Act provides authority for EPA to ban or restrict the manufacture (including import), processing, distribution in commerce, and use of chemical substances, as well as any manner or method of disposal. Section 26(l)(4) of TSCA authorizes EPA to publish proposed and final rules under TSCA section 6(a) that are consistent with the scope of completed TSCA Work Plan chemical risk assessments completed before June 22, 2016 and that are consistent with other applicable requirements of TSCA section 6. Methylene chloride is used in paint and coating removal in commercial processes and consumer products. In the August 2014 TSCA Work Plan Chemical Risk Assessment for methylene chloride, EPA characterized risks from use of these chemicals in paint and coating removal. On January 19, 2017, EPA preliminarily determined that the use of methylene chloride in paint and coating removal poses an unreasonable risk of injury to health. EPA also proposed prohibitions and restrictions on the manufacture, processing, and distribution in commerce of methylene chloride for all consumer and most types of commercial paint and coating removal and on the use of methylene chloride in commercial paint and coating removal in specified sectors. In the final rule published on March 27, 2019, EPA determined that the use of methylene chloride in consumer paint and coating removal presents an unreasonable risk of injury to health due to acute human lethality but, exercising its discretion under section 26(l)(4), EPA did not finalize such a determination concerning the use of methylene chloride in commercial paint and coating removal and did not finalize that portion of the proposed rule. To address the unreasonable risk to consumers of acute human lethality, the final rule prohibits the manufacture (including import), processing, and distribution in commerce of methylene chloride for consumer paint and coating removal, including distribution to and by retailers; requires manufacturers (including importers), processors, and distributors, except for retailers, of methylene chloride for any use to provide downstream notification of these prohibitions; and requires the retention of certain records. While EPA proposed to identify the use of methylene chloride in commercial furniture refinishing as presenting an unreasonable risk, EPA intends to further evaluate this and other commercial paint and coating removal uses and develop an appropriate regulatory risk management approach under the process for risk evaluations for existing chemicals under TSCA. Although N-methylpyrrolidone (NMP) was included in the January 2017 proposed rule, EPA intends to address NMP use in paint and coating removal in the risk evaluation for NMP and to consider any resulting risk reduction requirements in a separate regulatory action (RIN 2070–AK46).

Timetable:

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<tr>
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<tr>
<td>NPRM</td>
<td>01/19/17</td>
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<td>08/30/17</td>
<td>82 FR 41256</td>
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<tr>
<td>Final Rule</td>
<td>03/27/19</td>
<td>84 FR 11420</td>
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<tr>
<td>Final Rule Effec-</td>
<td>05/28/19</td>
<td>84 FR 11420</td>
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tive.

Regulatory Flexibility Analysis

Required: Yes.

Agency Contact: Joel Wolf, Environmental Protection Agency, Office of Chemical Safety and Pollution Prevention, 1200 Pennsylvania Avenue NW, Mail Code 7405M, Washington, DC 20460, Phone: 202 564–0432, Email: wolf.joel@epa.gov.

RIN: 2070–AK07

[FR Doc. 2019–11858 Filed 6–21–19; 8:45 am]

BILLING CODE 6560–50–P