

targeted investment that would not serve the needs of State VR agencies, their clients, or the public at large.

The Department has also concluded that it would be contrary to the public interest to have a lapse in the provision of TA and CE currently provided by the TACE Centers. Allowing funding to lapse before a new TA and CE delivery system can be implemented would leave State VR agencies and their partners without necessary supports in the event that critical needs arise.

For these reasons, the Secretary proposes to waive the requirements in 34 CFR 75.250, which prohibit project periods exceeding five years, and the requirements in 34 CFR 75.261(c)(2), which limits the extension of a project period if the extension involves the obligation of additional Federal funds, and to issue continuation awards to the ten current TACE grantees for a total amount not to exceed \$9,000,000. Under this proposal, the eight current TACE grantees with project periods ending on September 30, 2013, would receive funding to operate for an additional 12 months. The two current TACE grantees with project periods ending on December 21, 2013, would receive funds for an additional nine months. Consequently, the expiration date for all 10 grants would be September 30, 2014. Waiving these regulations and issuing these continuation awards will ensure that TA and CE to State VR agencies and their partners will not be interrupted. It will also ensure that the Department has adequate opportunity to solicit feedback and develop a coordinated TA and CE strategy that will best meet the needs of State VR agencies and their partners.

With this proposed extension of project period and waiver, each TACE Center will be required to continue to carry out activities during the year of the continuation award consistent with the scope, goals, and objectives of the grantee's application as approved in the 2008 competition.

Regulatory Flexibility Act Certification

The Secretary certifies that this proposed extension of project period and waiver would not have a significant economic impact on a substantial number of small entities. The only entities that would be affected are the ten current grantees receiving Federal funds to serve as the TACE Centers and any other potential applicants.

The Secretary certifies that the proposed waivers and extensions would not have a significant economic impact on these entities because the proposed waivers and extensions impose minimal compliance costs to extend projects already in existence, and the activities

required to support the additional year of funding would not impose additional regulatory burdens or require unnecessary Federal supervision.

Paperwork Reduction Act of 1995

This notice of proposed extension of project period and waiver does not contain any information collection requirements.

Intergovernmental Review: This program is subject to the requirements of Executive Order 12372 and the regulations in 34 CFR part 79. One of the objectives of the Executive order is to foster an intergovernmental partnership and a strengthened federalism. The Executive order relies on processes developed by State and local governments for coordination and review of proposed Federal financial assistance. This document provides early notification of our specific plans and actions for this program.

Accessible Format: Individuals with disabilities can obtain this document in an accessible format (e.g., braille, large print, audiotape, or compact disc) on request to the contact person listed under **FOR FURTHER INFORMATION CONTACT**.

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Dated: March 12, 2013.

Michael Yudin,

Acting Assistant Secretary for Special Education and Rehabilitative Services.

[FR Doc. 2013-06077 Filed 3-14-13; 8:45 am]

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ARCHITECTURAL AND TRANSPORTATION BARRIERS COMPLIANCE BOARD

36 CFR Part 1195

[Docket No. ATBCB-2012-0003]

RIN 3014-AA40

Medical Diagnostic Equipment Accessibility Standards Advisory Committee

AGENCY: Architectural and Transportation Barriers Compliance Board.

ACTION: Notice of advisory committee meeting.

SUMMARY: The Medical Diagnostic Equipment Accessibility Standards Advisory Committee will hold its fifth meeting. On July 5, 2012, the Architectural and Transportation Barriers Compliance Board (Access Board) established the advisory committee to make recommendations to the Board on matters associated with comments received and responses to questions included in a previously published Notice of Proposed Rulemaking (NPRM) on Medical Diagnostic Equipment Accessibility Standards.

DATES: The Committee will meet on March 26, 2013 from 10:00 a.m. to 5:00 p.m. and on March 27, 2013 from 9:00 a.m. to 2:30 p.m.

ADDRESSES: The meeting will be held at the Access Board's Conference Room, 1331 F Street NW., Suite 800, Washington, DC 20004-1111.

FOR FURTHER INFORMATION CONTACT: Rex Pace, Office of Technical and Information Services, Architectural and Transportation Barriers Compliance Board, 1331 F Street NW., Suite 1000, Washington, DC 20004-1111. Telephone number (202) 272-0023 (Voice); (202) 272-0052 (TTY). Electronic mail address: pace@access-board.gov.

SUPPLEMENTARY INFORMATION: On July 5, 2012, the Access Board established an advisory committee to make recommendations to the Board on matters associated with comments received and responses to questions included in a previously published NPRM on Medical Diagnostic Equipment Accessibility Standards. See 77 FR 6916 (February 9, 2012). The NPRM and information related to the proposed standards are available on the Access Board's Web site at: <http://www.access-board.gov/medical-equipment.htm>.

The advisory committee will hold its fifth meeting on March 26 and 27, 2013. The agenda includes the following:

- Review of previous committee work;
- Review and discussion of subcommittee work and recommendations;
- Continued discussion on recommendations for transfer surface height and Transfer support location and configuration
- Consideration of issues proposed by committee members; and
- Discussion of administrative issues.

The preliminary meeting agenda, along with information about the committee, is available at the Access Board's Web site (<http://www.access-board.gov/medical-equipment.htm>).

Committee meetings are open to the public and interested persons can attend the meetings and communicate their views. Members of the public will have opportunities to address the committee on issues of interest to them during public comment periods scheduled on each day of the meeting.

The meetings will be accessible to persons with disabilities. An assistive listening system, computer assisted real-time transcription (CART), and sign language interpreters will be provided. Persons attending the meetings are requested to refrain from using perfume, cologne, and other fragrances for the comfort of other participants (see www.access-board.gov/about/policies/fragrance.htm for more information). Also, persons wishing to provide handouts or other written information to the committee are requested to provide electronic formats to Rex Pace via email prior to the meetings so that alternate formats can be distributed to committee members.

David M. Capozzi,
Executive Director.

[FR Doc. 2013-05936 Filed 3-14-13; 8:45 am]

BILLING CODE 8150-01-P

ENVIRONMENTAL PROTECTION AGENCY

40 CFR Part 52

[EPA-R05-OAR-2012-0650; FRL-9789-8]

Approval and Promulgation of Air Quality Implementation Plans; Indiana; Consent Decree Requirements

AGENCY: Environmental Protection Agency (EPA).

ACTION: Proposed rule.

SUMMARY: EPA is proposing to approve a portion of Indiana's construction

permit rule for sources subject to the state operating permit program regulations at 40 CFR part 70. These provisions authorize the state to incorporate terms from Federal consent decrees or Federal district court orders into these construction permits. EPA is also approving public notice requirements for these permit actions. These rules will help streamline the process for making Federal consent decree and Federal district court order requirements permanent and Federally enforceable.

DATES: Comments must be received on or before April 15, 2013.

ADDRESSES: Submit your comments, identified by Docket ID No. EPA-R05-OAR-2012-0650, by one of the following methods:

1. *www.regulations.gov*: Follow the on-line instructions for submitting comments.
2. *Email*: damico.genevieve@epa.gov.
3. *Fax*: (312) 385-5501.
4. *Mail*: Genevieve Damico, Chief, Air Permits Section, Air Programs Branch (AR-18J), U.S. Environmental Protection Agency, 77 West Jackson Boulevard, Chicago, Illinois 60604.
5. *Hand Delivery*: Genevieve Damico, Chief, Air Permits Section, Air Programs Branch (AR-18J), U.S. Environmental Protection Agency, 77 West Jackson Boulevard, Chicago, Illinois 60604. Such deliveries are only accepted during the Regional Office normal hours of operation, and special arrangements should be made for deliveries of boxed information. The Regional Office official hours of business are Monday through Friday, 8:30 a.m. to 4:30 p.m., excluding Federal holidays.

Please see the direct final rule which is located in the Rules section of this **Federal Register** for detailed instructions on how to submit comments.

FOR FURTHER INFORMATION CONTACT: Sam Portanova, Environmental Engineer, Air Permits Section, Air Programs Branch (AR-18J), Environmental Protection Agency, Region 5, 77 West Jackson Boulevard, Chicago, Illinois 60604, (312) 886-3189, portanova.sam@epa.gov.

SUPPLEMENTARY INFORMATION: In the Final Rules section of this **Federal Register**, EPA is approving the State's SIP submittal as a direct final rule without prior proposal because the Agency views this as a noncontroversial submittal and anticipates no adverse comments. A detailed rationale for the approval is set forth in the direct final rule. If no adverse comments are received in response to this rule, no

further activity is contemplated. If EPA receives adverse comments, the direct final rule will be withdrawn and all public comments received will be addressed in a subsequent final rule based on this proposed rule. EPA will not institute a second comment period. Any parties interested in commenting on this action should do so at this time. Please note that if EPA receives adverse comment on an amendment, paragraph, or section of this rule and if that provision may be severed from the remainder of the rule, EPA may adopt as final those provisions of the rule that are not the subject of an adverse comment. For additional information, see the direct final rule which is located in the Rules section of this **Federal Register**.

Dated: March 4, 2013.

Susan Hedman,

Regional Administrator, Region 5.

[FR Doc. 2013-05953 Filed 3-14-13; 8:45 am]

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ENVIRONMENTAL PROTECTION AGENCY

40 CFR Part 52

[EPA-R03-OAR-2013-0113; FRL-9790-9]

Approval and Promulgation of Air Quality Implementation Plans; West Virginia; Prevention of Significant Deterioration

AGENCY: Environmental Protection Agency (EPA).

ACTION: Proposed rule.

SUMMARY: EPA is proposing to disapprove a narrow portion of a State Implementation Plan (SIP) revision submitted by the State of West Virginia on August 31, 2011. EPA is proposing this action because a narrow portion of the submittal does not satisfy the Federal requirement for the inclusion of condensable emissions of particulate matter (condensables) within the definition of "regulated new source review (NSR) pollutant." Additionally, because West Virginia's August 31, 2011 SIP revision does not adequately account for condensable emissions within the definition of "regulated NSR pollutant," EPA is also proposing to disapprove specific Prevention of Significant Deterioration (PSD) portions of related infrastructure submissions required by the Clean Air Act (CAA) to implement, maintain, and enforce the 1997 fine particulate matter (PM_{2.5}) and ozone National Ambient Air Quality Standards (NAAQS), the 2006 PM_{2.5} NAAQS, and the 2008 lead and ozone