This rule will not 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, as specified in Executive Order 13132, because it merely proposes to approve or disapprove State rules implementing a Federal standard, and does not alter the relationship or the distribution of power and responsibilities established in the Clean Air Act. Thus, the requirements of section 6 of the Executive Order do not apply to this rule.

F. Executive Order 13175, Coordination With Indian Tribal Governments

Executive Order 13175, entitled "Consultation and Coordination with Indian Tribal Governments" (65 FR 67249, November 9, 2000), requires EPA to develop an accountable process to ensure "meaningful and timely input by Tribal officials in the development of regulatory policies that have Tribal implications." This proposed rule does not have Tribal implications, as specified in Executive Order 13175. It will not have substantial direct effects on Tribal governments, on the relationship between the Federal government and Indian Tribes, or on the distribution of power and responsibilities between the Federal government and Indian Tribes. Thus, Executive Order 13175 does not apply to this rule.

EPA specifically solicits additional comment on this proposed rule from Tribal officials.

G. Executive Order 13045, Protection of Children From Environmental Health Risks and Safety Risks

EPA interprets Executive Order 13045 (62 FR 19885, April 23, 1997) as applying only to those regulatory actions that concern health or safety risks, such that the analysis required under section 5–501 of the Executive Order has the potential to influence the regulation. This rule is not subject to Executive Order 13045, because it approves state rules implementing a Federal standard.

H. Executive Order 13211, Actions That Significantly Affect Energy Supply, Distribution, or Use

This rule is not subject to Executive Order 13211, "Actions Concerning Regulations That Significantly Affect Energy Supply, Distribution, or Use" (66 FR 28355, May 22, 2001) because it is not a significant regulatory action under Executive Order 12866. I. National Technology Transfer and Advancement Act

Section 12 of the National Technology Transfer and Advancement Act (NTTAA) of 1995 requires Federal agencies to evaluate existing technical standards when developing a new regulation. To comply with NTTAA, EPA must consider and use "voluntary consensus standards" (VCS) if available and applicable when developing programs and policies unless doing so would be inconsistent with applicable law or otherwise impractical.

The EPA believes that VCS are inapplicable to this action. Today's action does not require the public to perform activities conducive to the use of VCS.

J. Executive Order 12898: Federal Actions To Address Environmental Justice in Minority Populations and Low-Income Population

Executive Order (EO) 12898 (59 FR 7629 (Feb. 16, 1994)) establishes Federal executive policy on environmental justice. Its main provision directs Federal agencies, to the greatest extent practicable and permitted by law, to make environmental justice part of their mission by identifying and addressing, as appropriate, disproportionately high and adverse human health or environmental effects of their programs, policies, and activities on minority populations and low-income populations in the United States.

EPA lacks the discretionary authority to address environmental justice in this rulemaking.

## List of Subjects in 40 CFR Part 52

Environmental protection, Air pollution control, Intergovernmental relations, Nitrogen dioxide, Ozone, Particulate matter, Reporting and recordkeeping requirements.

**Authority:** 42 U.S.C. 7401 *et seq.* Dated: June 21, 2011.

## Jared Blumenfeld,

Regional Administrator, Region IX.  $[{\rm FR\ Doc.\ 2011-17262\ Filed\ 7-7-11;\ 8:45\ am}]$ 

BILLING CODE 6560-50-P

## **DEPARTMENT OF TRANSPORTATION**

## Federal Motor Carrier Safety Administration

49 CFR Parts 382 and 391

[Docket No. FMCSA-2011-0073]

RIN 2126-AB35

## Harmonizing Schedule I Drug Requirements

**AGENCY:** Federal Motor Carrier Safety Administration, DOT.

**ACTION:** Notice of proposed rulemaking.

**SUMMARY:** The Federal Motor Carrier Safety Administration (FMCSA) proposes to amend the physical qualifications for drivers and the instructions for the medical examination report to clarify that drivers may not use Schedule I drugs and be qualified to drive commercial motor vehicles under any circumstances. The proposal also harmonizes FMCSA's provisions regarding pre-employment and returnto-duty test refusals with corresponding Department of Transportation (DOT)wide provisions. Finally, the proposal corrects inaccurate uses of the term "actual knowledge."

**DATES:** Comments and related material must be submitted on or before September 6, 2011.

**ADDRESSES:** You may submit comments identified by docket number FMCSA–2011–0073 using any one of the following methods:

- Federal eRulemaking Portal: http://www.regulations.gov.
  - Fax: 202-493-2251.
- *Mail:* Docket Management Facility (M–30), U.S. Department of Transportation, West Building Ground Floor, Room W12–140, 1200 New Jersey Avenue, SE., Washington, DC 20590–0001.
- Hand delivery: Same as mail address above, between 9 a.m. and 5 p.m., e.t., Monday through Friday, except Federal holidays. The telephone number is 202–366–9329.

To avoid duplication, please use only one of these four methods. See the "Public Participation and Request for Comments" portion of the

**SUPPLEMENTARY INFORMATION** section below for instructions on submitting comments.

FOR FURTHER INFORMATION CONTACT: If you have questions on this proposed rule, call or e-mail Angela Ward, Nurse Consultant, Medical Programs Office, Federal Motor Carrier Safety Administration, telephone: 202–366–

3109; e-mail: angela.ward@dot.gov. If you have questions on viewing or submitting material to the docket, call Renee V. Wright, Program Manager, Docket Operations, telephone 202–366–9826.

## SUPPLEMENTARY INFORMATION:

## **Table of Contents for Preamble**

- I. Public Participation and Request for Comments
  - A. Submitting Comments
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## I. Public Participation and Request for Comments

FMCSA encourages you to participate in this rulemaking by submitting comments and related materials.

## A. Submitting Comments

If you submit a comment, please include the docket number for this rulemaking (FMCSA-2011-0073), indicate the specific section of this document to which each comment applies, and provide a reason for each suggestion or recommendation. You may submit your comments and material online or by fax, mail, or hand delivery, but please use only one of these means. FMCSA recommends that you include your name and a mailing address, an e-mail address, or a phone number in the body of your document so that the Agency can contact you if it has questions regarding your submission.

To submit your comment online, go to http://www.regulations.gov, click on the "Submit a Comment" box, which will then become highlighted in blue. In the "Document Type" drop down menu, select "Rules," insert "FMCSA-2011-0073" in the "Keyword" box, and click "Search." When the new screen appears, click on "Submit a Comment" in the "Actions" column. If you submit your comments by mail or hand delivery, submit them in an unbound format, no larger than 8½ by 11 inches, suitable for copying and electronic filing. If you submit comments by mail and would like to know that they reached the facility, please enclose a stamped, self-addressed postcard or envelope.

FMCSA will consider all comments and material received during the comment period and may change this proposed rule based on your comments. B. Viewing Comments and Documents

To view comments, as well as documents mentioned in this preamble as being available in the docket, go to http://www.regulations.gov and click on the "Read Comments" box in the upper right hand side of the screen. Then, in the "Keyword" box, insert "FMCSA-2011-0073" and click "Search." Next, click "Open Docket Folder" in the "Actions" column. Finally, in the "Title" column, click on the document you would like to review. If you do not have access to the Internet, you may view the docket online by visiting the Docket Management Facility in Room W12-140 on the ground floor of the DOT West Building, 1200 New Jersey Avenue, SE., Washington, DC 20590, between 9 a.m. and 5 p.m., e.t., Monday through Friday, except Federal holidays.

## C. Privacy Act

All comments received will be posted without change to http:// www.regulations.gov and will include any personal information you have provided. Anyone is able to search the electronic form for all comments received into any of our dockets by the name of the individual submitting the comment (or signing the comment, if submitted on behalf of an association, business, labor union, etc.). You may review DOT's complete Privacy Act Statement in the Federal Register published on January 17, 2008 (73 FR 3316), or you may visit http://edocket. access.gpo.gov/2008/pdf/E8-785.pdf.

## II. Abbreviations

CAA CFR	Clean Air Act. Code of Federal Regulations.
CMV	Commercial Motor Vehicle.
DEA	Drug Enforcement Administra- tion.
FMCSA	Federal Motor Carrier Safety Administration.
FR	Federal Register.
NEPA	National Environmental Policy Act.
OTETA	Omnibus Transportation Employee Testing Act of 1991.
U.S.C	United States Code.

## III. Background

## A. History

The Omnibus Transportation
Employee Testing Act of 1991 (OTETA),
49 U.S.C. 31306, mandated that DOT
establish a controlled substances (drug)
and alcohol testing program applicable
to regulated entities and individuals
performing safety sensitive functions.
Entitled "Procedures for Transportation
Workplace Drug and Alcohol Testing
Programs," 49 CFR part 40 contains the
DOT regulations that detail how testing

must be administered and prescribes procedures to protect the integrity of the process. The FMCSA's related drug and alcohol testing regulations are in 49 CFR part 382, "Controlled Substances and Alcohol Use and Testing."

DEA implemented the Comprehensive Drug Abuse Prevention and Control Act of 1970, often referred to as the Controlled Substances Act and the Controlled Substances Import and Export Act (21 U.S.C. 801-971), as amended. DEA published regulations implementing these statutes in 21 CFR Parts 1300 to 1399. These regulations are designed to ensure an adequate supply of controlled substances for legitimate medical, scientific, research, and industrial purposes, and to deter the diversion of controlled substances to illegal purposes. Controlled substances are drugs and other substances that have a potential for abuse and psychological and physical dependence. DEA lists controlled substances in 21 CFR part 1308. The substances are divided into five schedules. The substances listed in the schedule that are relevant to this rulemaking, Schedule I, have a high potential for abuse and have no currently accepted medical use in the United States (DEA Interim Final Rule on Electronic Prescriptions for Controlled Substances, 75 FR 16237, March 31, 2010). These substances may only be used for research, chemical analysis, or manufacture of other drugs.

Section 382.213 prohibits commercial motor vehicle (CMV) drivers from using any controlled substances when on duty or reporting for duty except when prescribed by a licensed medical practitioner who has advised the driver that the prescribed substance will not adversely affect the driver's ability to operate a CMV. Section 382.213 has remained largely unchanged since its adoption in 1994, outside of a technical amendment changing the term "physician" to "licensed medical practitioner" for the purpose of the prescription exception (61 FR 9556, March 8, 1996).

In addition to those in part 382, FMCSA has several other regulations governing drivers' use of drugs. Section 391.41(b)(12) was first promulgated in 1970, and stated that persons who "use an amphetamine, narcotic, or any habitforming drug, are not medically qualified to operate a commercial motor vehicle" (35 FR 6463, April 22, 1970). Section 391.43(f) incorporates the substance of § 391.41(b)(12) in the instructions to the medical examiner. Section 391.41(b)(12) was revised several times, most notably in 1984, when the DEA's Schedule I drugs were added to the list of drugs prohibited by

§ 391.41(b)(12) (49 FR 44215, November 5, 1984). Sections 382.213 and 391.41(b)(12) were designed to complement § 392.4, which prohibits the use of drugs by CMV drivers. Section 392.4 contains an exception for use of non-Schedule I drugs "administered to a driver by or under the instructions of a licensed medical practitioner, as defined in § 382.107 of this subchapter, who has advised the driver that the substance will not affect the driver's ability to safely operate a motor vehicle" (49 CFR 392.4).

## B. Legal Authority

FMCSA has general authority to promulgate safety standards, including those governing drivers' use of drugs while operating a CMV. The Motor Carrier Safety Act of 1984 (Pub. L. 98– 554, Title II, 98 Stat. 2832, October 30, 1984) (the 1984 Act) provides authority to regulate drivers, motor carriers, and vehicle equipment. It requires the Secretary to ensure that—(1) CMVs are maintained, equipped, loaded, and operated safely; (2) the responsibilities imposed on operators of CMVs do not impair their ability to operate the vehicles safely; (3) the physical condition of CMV operators is adequate to enable them to operate the vehicles safely; and (4) the operation of CMVs does not have a deleterious effect on the physical condition of the operators (49) U.S.C. 31136(a)). Section 211 of the 1984 Act also grants the Secretary broad power in carrying out motor carrier safety statutes and regulations to "prescribe recordkeeping and reporting requirements" and to "perform other acts the Secretary considers appropriate" (49 U.S.C. 31133(a)(8) and

The FMCSA Administrator has been delegated authority under 49 CFR 1.73(g) to carry out the functions vested in the Secretary of Transportation by 49 U.S.C. chapter 311, subchapters I and III, relating to CMV programs and safety regulation.

As stated above, OTETA (Pub. L. 102– 143, Title V, 105 Stat. 917, at 952, Oct. 28, 1991, codified at 49 U.S.C. 31306), mandated the alcohol and controlled substances (drug) testing program for DOT. OTETA required the Secretary of Transportation to promulgate regulations for alcohol and controlled substances testing for persons in safetysensitive positions in four modes of transportation—motor carrier, airline, railroad, and mass transit. Those regulations, including subsequent amendments, are codified at 49 CFR part 40, "Procedures for Transportation Workplace Drug and Alcohol Testing Programs." Part 40 prescribes drug and

alcohol testing requirements for all DOT-regulated parties, including employers of drivers with commercial driver's licenses subject to FMCSA testing requirements. FMCSA's related drug and alcohol testing regulations are in 49 CFR part 382, "Controlled Substances and Alcohol Use and Testing."

## C. Discussion of the Proposed Rule

This rulemaking is necessary to reconcile and resolve a perceived inconsistency among: §§ 382.213, 391.41(b)(12), 391.43(f), and 392.4 of the Federal Motor Carrier Safety Regulations (FMCSRs); DOT-wide drug regulations in part 40; and DEA regulations. Although § 392.4 clearly prohibits drivers from using Schedule I drugs, it has come to FMCSA's attention that some people might interpret §§ 382.213, 391.41(b)(12) and 391.43(f) to permit their use if recommended by a licensed medical practitioner. The FMCSA has always considered §§ 382.213, 391.41(b)(12), 391.43(f), and 392.4 to prohibit any and all use of Schedule I drugs by CMV drivers. In fact, Federal law prohibits Schedule I drugs from being prescribed in the United States (75 FR 16237, March 31, 2010). Schedule I drugs have a high potential for abuse and no medically accepted therapeutic use (id.). Currently, Federal law only allows for their use in research, chemical analysis, or manufacture of other drugs (id.).

In certain circumstances, a medical review officer can verify a drug test negative when he or she has information that a driver is using a drug under a physician's prescription. However, under DOT-wide rules, no medical review officer may verify a drug test negative for a Schedule I drug, even if he or she has information that a driver is using the Schedule I drug in accordance with a physician's recommendation (49 CFR 40.151(e)). Interpreting FMCSA's regulations to permit drivers to use Schedule I drugs would put the FMCSRs in direct conflict with DOT's comprehensive drug testing program under 49 CFR part 40, which does not permit drivers to use Schedule I drugs. The FMCSA does not believe this is a reasonable interpretation of the regulations. Regardless, to avoid any confusion, this rulemaking would harmonize §§ 382.213, 391.41(b)(12), 391.43(f), and 392.4 with DOT-wide regulations and DEA regulations, and make it clear that drivers may not use Schedule I drugs under any circumstances.

In addition, 49 CFR 382.211 prohibits drivers from refusing to submit to certain types of drug or alcohol tests and establishes such refusals as violations of FMCSA's drug and alcohol regulations. Currently, under DOT-wide regulations, drivers who refuse to submit to preemployment and return-to-duty tests must complete the return-to-duty process prescribed in part 40, subpart O. However, § 382.211 is inconsistent with the DOT-wide drug and alcohol rules in that it does not include refusals to submit to pre-employment and return-to-duty tests as violations. The FMCSA proposes to correct this inconsistency by adding these two types of refusals to the prohibitions at § 382.211.

Finally, FMCSA proposes changes to 49 CFR 382.201 and 382.215 to clarify the Agency's rules prohibiting an employer from using a driver about whom the employer has actual knowledge of drug or alcohol use, as defined at § 382.107. Sections 382.201 and 382.215 currently state that an employer may not allow an employee to perform safety-sensitive functions if the employer has actual knowledge that the employee has tested positive for drugs or has an alcohol concentration of .04 or greater. However, the term "actual knowledge" is defined in § 382.107 to mean the observation of alcohol or controlled substances use, and is not intended to refer to testing results. As a result, the use of the term "actual knowledge" in these sections is not appropriate. FMCSA proposes to replace the term "actual knowledge" with "knowledge" in these sections. This should clarify that these prohibitions refer to the knowledge of test results, not employer observation of prohibited conduct.

## IV. Section-by-Section Analysis

Sections 382.201 and 382.215

An employer has "actual knowledge" that an employee has used drugs or alcohol in violation of FMCSA rules when he or she directly observes or otherwise learns that a driver is using controlled substances or consuming alcohol while on duty (49 CFR 382.107). Actual knowledge, as defined at § 382.107, is distinct from an employer knowing that his or her employee-driver tested positive or refused a DOT drug or alcohol test. Because §§ 382.201 and 382.215 set forth prohibitions related to an employer's knowledge related to testing, not observation, the use of the term "actual knowledge" is not appropriate. The FMCSA proposes to replace the term "actual knowledge" with "knowledge" in these sections. This would clarify that these prohibitions refer to the knowledge of test results, not employer observation of prohibited conduct.

Section 382.211

Current § 382.211 prohibits drivers from refusing to submit to a post-accident, random, or reasonable suspicion drug or alcohol test. The Agency proposes to amend § 382.211 to also prohibit refusals for pre-employment testing and return-to-duty testing. This would make this regulation consistent with 49 CFR 40.191(a)(3).

## Section 382.213

Section 382.213 currently prohibits CMV drivers from using any drugs when on duty or reporting for duty except when prescribed by a licensed medical practitioner who has advised the driver that the prescribed substance will not adversely affect the driver's ability to operate a CMV. The Agency proposes to amend the language regarding the drugs that CMV drivers are prohibited from using in order to differentiate between Schedule I drugs and non-Schedule I drugs. The proposed changes would make it clear that Schedule I drugs may not be used by a CMV driver under any circumstances. The FMCSA's regulations would continue to permit the use of non-Schedule I drugs under limited circumstances, when prescribed by a licensed medical practitioner.

## Sections 391.41 and 391.43

Section 391.41(b)(12)(i) currently states that a driver may not use: Controlled substances on the DEA Schedule I, amphetamines, narcotics, or other habit-forming drugs. Section 391.41(b)(12)(ii) contains an exception for a substance or drug prescribed by a licensed medical practitioner who is familiar with the driver's history and work duties and has advised the driver that the prescribed substance or drug will not adversely affect his or her ability to safely operate a CMV. The FMCSA has never considered this exception to permit use of Schedule I drugs by CMV drivers under any circumstance because Federal law prohibits Schedule I drugs from being prescribed in the United States (75 FR 16237, March 31, 2010). Section 391.43(f) incorporates the substance of § 391.41(b)(12) into pages 4 and 8 of the Instructions to the Medical Examiner. The FMCSA makes no others changes to this document.

Section 391.41(b)(12) and the Instructions for Medical Examiners at § 391.43(f) currently do not differentiate between Schedule I and non-Schedule I drugs for the purpose of the prescription exception. The prescription exception currently states that a CMV driver may use a substance or drug that is prescribed by a licensed medical

practitioner who is familiar with the driver's medical history and has advised the driver that the prescribed substance or drug will not adversely affect the driver's ability to safely operate a CMV. The Agency proposes to amend these sections to clarify that this exception only applies to non-Schedule I prescribed substances, amphetamines, narcotics, or other habit-forming drugs.

## V. Regulatory Analyses

Regulatory Planning and Review

This action does not meet the criteria for a "significant regulatory action," either as specified in Executive Order 12866 as supplemented by Executive Order 13563 (76 FR 3821, January 18, 2011) or within the meaning of the DOT regulatory policies and procedures (44 FR 1103, February 26, 1979). The estimated economic costs of the proposed rule do not exceed the \$100 million annual threshold nor does the Agency expect the proposed rule to have substantial Congressional or public interest. Therefore, this proposed rule has not been formally reviewed by the Office of Management and Budget. No expenditures would be required of the affected population because the proposed rule would only clarify existing rules, amend inconsistencies in FMCSA's current regulations, and harmonize them with the DOT-wide regulations and DEA regulations.

## Regulatory Flexibility Act

The Regulatory Flexibility Act of 1980 (5 U.S.C. 601-612) requires Federal agencies to consider the effects of the regulatory action on small business and other small entities and to minimize any significant economic impact. The term "small entities" comprises small businesses and not-for-profit organizations that are independently owned and operated and are not dominant in their fields, as well as governmental jurisdictions with populations of less than 50,000. Accordingly, DOT policy requires an analysis of the impact of all regulations on small entities and mandates that agencies strive to lessen any adverse effects on these businesses.

Under the Regulatory Flexibility Act, as amended by the Small Business Regulatory Enforcement Fairness Act of 1996, the proposed rule is not expected to have a significant economic impact on a substantial number of small entities because the proposed rule would only clarify existing rules, amend inconsistencies in FMCSA's current regulations, and harmonize them with the DOT-wide regulations and DEA regulations. Accordingly, I certify that a

regulatory flexibility analysis is not necessary.

Assistance for Small Entities

Under section 213(a) of the Small **Business Regulatory Enforcement** Fairness Act of 1996 (Pub. L. 104-121), FMCSA wants to assist small entities in understanding this proposed rule so that they can better evaluate its effects on them and participate in the rulemaking initiative. If the proposed rule would affect your small business, organization, or governmental jurisdiction and you have questions concerning its provisions or options for compliance, please consult the FMCSA point of contact, Angela Ward, listed in the FOR **FURTHER INFORMATION CONTACT** section of this proposed rule. FMCSA will not retaliate against small entities that question or complain about this proposed rule or any policy or action of the Agency.

Small businesses may send comments on the actions of Federal employees who enforce or otherwise determine compliance with Federal regulations to the Small Business and Agriculture Regulatory Enforcement Ombudsman and the Regional Small Business Regulatory Fairness Boards. The Ombudsman evaluates these actions annually and rates each agency's responsiveness to small business. If you wish to comment on actions by employees of FMCSA, call 1–888–REG–FAIR (1–888–734–3247).

Unfunded Mandates Reform Act of 1995

The Unfunded Mandates Reform Act of 1995 (2 U.S.C. 1531-1538) requires Federal agencies to assess the effects of their discretionary regulatory actions. In particular, the Act addresses actions that may result in the expenditure by a State, local, or Tribal government, in the aggregate, or by the private sector of \$140.8 million (which is the value of \$100 million in 2010 after adjusting for inflation) or more in any 1 year. This proposed rule would not result in such expenditure; FMCSA expects the effects of this proposed rule to be minimal because the proposed rule would only clarify existing rules, amend inconsistencies in FMCSA's current regulations, and harmonize them with the DOT-wide regulations and DEA regulations.

## Paperwork Reduction Act

This proposed rule would call for no new collection of information under the Paperwork Reduction Act of 1995 (44 U.S.C. 3501–3520). Privacy Impact Assessment

FMCSA conducted a Privacy Threshold Analysis for the Notice of Proposed Rulemaking (NPRM) and determined that this proposed rule is not a privacy-sensitive rulemaking because if promulgated as a final rule it would not require any collection, maintenance, or dissemination of Personally Identifiable Information from or about members of the public.

## Executive Order 13132 (Federalism)

A rule has implications for federalism under Executive Order 13132, Federalism, if it has a substantial direct effect on State or local governments and would either preempt State law or impose a substantial direct cost of compliance on States or localities. FMCSA has analyzed this proposed rule under that Order and has determined that it does not have implications for federalism.

## Executive Order 12630 (Taking of Private Property)

This proposed rule would not effect a taking of private property or otherwise have taking implications under Executive Order 12630, Governmental Actions and Interference with Constitutionally Protected Property Rights.

## Executive Order 12988 (Civil Justice Reform)

This proposed rule meets applicable standards in sections 3(a) and 3(b)(2) of Executive Order 12988, Civil Justice Reform, to minimize litigation, eliminate ambiguity, and reduce burden.

## Executive Order 13045 (Protection of Children)

FMCSA has analyzed this proposed rule under Executive Order 13045, Protection of Children from Environmental Health Risks and Safety Risks. This proposed rule is not an economically significant rule and would not create an environmental risk to health or risk to safety that might disproportionately affect children.

## Executive Order 13211 (Energy Effects)

FMCSA has analyzed this proposed rule under Executive Order 13211, Actions Concerning Regulations That Significantly Affect Energy Supply, Distribution, or Use. The Agency has determined that it is not a "significant energy action" under that order because it is not a "significant regulatory action" under Executive Order 12866 and is not likely to have a significant adverse effect on the supply, distribution, or use of energy.

## Technical Standards

The National Technology Transfer and Advancement Act (NTTAA) (15 U.S.C. 272 note) directs agencies to use voluntary consensus standards in their regulatory activities unless the agency provides Congress, through the Office of Management and Budget, with an explanation of why using these standards would be inconsistent with applicable law or otherwise impractical. Voluntary consensus standards are technical standards (e.g., specifications of materials, performance, design, or operation; test methods; sampling procedures; and related management systems practices) that are developed or adopted by voluntary consensus standards bodies. This proposed rule does not use technical standards. Therefore, FMCSA did not consider the use of voluntary consensus standards.

## National Environmental Policy Act

FMCSA analyzed this NPRM for the purpose of the National Environmental Policy Act of 1969 (NEPA) (42 U.S.C. 4321 et seq.) and determined under our environmental procedures Order 5610.1, published February 24, 2004 (69 FR 9680), that this proposed action does not have any effect on the quality of the environment. Therefore, this NPRM is categorically excluded from further analysis and documentation in an environmental assessment or environmental impact statement under FMCSA Order 5610.1, paragraph 6(r) of Appendix 2. The Categorical Exclusion under paragraph 6(y)(6) relates to "regulations implementing employer controlled substances and alcohol use and testing procedures \* \* \*," which is the focus of this rulemaking. A Categorical Exclusion determination is available for inspection or copying in the regulations.gov Web site listed under ADDRESSES.

In addition to the NEPA requirements to examine impacts on air quality, the Clean Air Act (CAA) as amended (42 U.S.C. 7401 et seq.) also requires FMCSA to analyze the potential impact of its actions on air quality and to ensure that FMCSA actions conform to State and local air quality implementation plans. The additional contributions to air emissions are expected to fall within the CAA de minimis standards and are not expected to be subject to the Environmental Protection Agency's General Conformity Rule (40 CFR parts 51 and 93).

FMCSA seeks comment on these determinations.

## **List of Subjects**

49 CFR Part 382

Administrative practice and procedure, Alcohol abuse, Drug abuse, Drug testing, Highway safety, Motor carriers, Penalties, Safety, Transportation.

## 49 CFR Part 391

Alcohol abuse, Drug abuse, Drug testing, Highway safety, Motor carriers, Reporting and recordkeeping requirements, Safety, Transportation.

For the reasons stated in the preamble, FMCSA proposes to amend 49 CFR, parts 382 and 391 as follows:

## **PART 382—CONTROLLED** SUBSTANCES AND ALCOHOL USE **AND TESTING**

1. The authority citation for part 382 continues to read as follows:

Authority: 49 U.S.C. 31133, 31136, 31301 et seq., 31502; and 49 CFR 1.73.

## § 382.201 [Amended]

- 2. Amend § 382.201 by removing the word "actual" between the words "having" and "knowledge."
  3. Revise § 382.211 to read as follows:

## § 382.211 Refusal to submit to a required alcohol or controlled substances test.

No driver shall refuse to submit to a pre-employment controlled substance test required under § 382.301, a postaccident alcohol or controlled substance test required under § 382.303, a random alcohol or controlled substances test required under § 382.305, a reasonable suspicion alcohol or controlled substance test required under § 382.307, a return-to-duty alcohol or controlled substances test required under § 382.309, or a follow-up alcohol or controlled substance test required under § 382.311. No employer shall permit a driver who refuses to submit to such tests to perform or continue to perform safety-sensitive functions.

4. Revise § 382.213 to read as follows:

## § 382.213 Controlled substance use.

- (a) No driver shall report for duty or remain on duty requiring the performance of safety sensitive functions when the driver uses any controlled substance identified in 21 CFR 1308.11.
- (b) No driver shall report for duty or remain on duty requiring the performance of safety-sensitive functions when the driver uses any non-Schedule I drug except when the use is pursuant to the instructions of a licensed medical practitioner, as defined in § 382.107, who is familiar with the driver's medical history and

has advised the driver that the substance will not adversely affect the driver's ability to safely operate a commercial motor vehicle.

(c) No employer having actual knowledge that a driver has used a controlled substance shall permit the driver to perform or continue to perform a safety-sensitive function.

(d) Ån employer may require a driver to inform the employer of any therapeutic drug use.

## § 382.215 [Amended]

5. Amend § 382.215 by removing the word "actual" between the words "having" and "knowledge."

## PART 391—QUALIFICATIONS OF DRIVERS AND LONGER COMBINATION VEHICLE (LCV) DRIVER INSTRUCTORS

6. The authority citation for part 391 continues to read as follows:

**Authority:** 49 U.S.C. 322, 504, 508, 31133, 31136, and 31502; sec. 4007(b) of Pub. L. 102–240, 105 Stat. 2152; sec. 114 of Pub. L. 103–311, 108 Stat. 1673, 1677; sec. 215 of Pub. L. 106–159, 113 Stat. 1767; and 49 CFR 1.73

7. Amend § 391.41 by revising paragraphs (b)(12)(i) and (ii) to read as follows:

## § 391.41 Physical qualifications for drivers.

\* \* \* \* \* \* (b) \* \* \*

(12)(i) Does not use any controlled substance identified in 21 CFR 1308.11 Schedule I, an amphetamine, a narcotic, or other habit-forming drug.

(ii) Does not use any non-Schedule I controlled substance except when the use is pursuant to the instructions of a licensed medical practitioner, as defined in § 382.107, who is familiar with the driver's medical history and

has advised the driver that the substance will not adversely affect the driver's ability to safely operate a commercial motor vehicle.

\* \* \* \* \*

8. Amend § 391.43(f) by removing the Medical Examination Report for Commercial Driver Fitness Determination, form 649–F (6045), and adding in its place the following form, to read as follows:

## § 391.43 Medical examination; certificate of physical examination.

\* \* \* \* \* \*

(f) \* \* \*

BILLING CODE 4910-EX-P

# Medical Examination Report FOR COMMERCIAL DRIVER FITNESS DETERMINATION

649-F (6045)

Fainting, dizziness Sleep disorders, pauses in breathing while asleep, daytime sleepiness, loud Stroke or paralysis Missing or impaired hand, arm, foot, leg, Regular, frequent alcohol use Narcotic or habit forming drug use For any YES answer, indicate onset date, diagnosis, treating physician's name and address, and any current limitation. List all medications (including State of Issue Date of Exam Spinal injury or disease Chronic low back pain New Certification Recertification Follow-up ŝ Yes Driver License No. Driver completes this section, but medical examiner is encouraged to discuss with driver ☐ diet ☐ pills ☐ insulin Nervous or psychiatric disorders, e.g., severe depression Lung disease, emphysema, asthma, chronic bronchitis Sex □□⊠ Digestive problems
Diabetes or elevated blood sugar controlled by: Age Loss of, or altered consciousness Kidney disease, dialysis Birthdate M/D/Y Work Tel: ( ) Home Tel: ( Social Security No. Yes DRIVER'S INFORMATION Driver completes this section over-the-counter medications) used regularly or recently City, State, Zip Code Eye disorders or impaired vision (except corrective lenses) Ear disorders, loss of hearing or balance Heart disease or heart attack; other cardiovascular condition Heart surgery (valve replacement/bypass, angioplasty, Any illness or injury in the last 5 years? Head/Brain injuries, disorders or illnesses Seizures, epilepsy medication Driver's Name (Last, First, Middle) pacemaker)
High blood pressure
Muscular disease
Shortness of breath HEALTH HISTORY res No Address

certify that the above information is complete and true. I understand that inaccurate, false or missing information may invalidate the examination and my Date Driver's Signature Medical Examiner's Certificate.

Medical Examiner's Comments on Health History (The medical examiner must review and discuss with the driver any "yes" answers and potential hazards of medications, including over-the-counter medications, while driving. This discussion must be documented below.)

TESTIN(	TESTING (Medical Examiner complet	niner comple		es Section 3 through 7) Name: Last	7) Name: Last,	First,		Middle	ď.	
3. VISION		l: At least 20/40 and in each eye. The	acuity (Snellen he use of corre	in each eye wit ective lenses sho	Standard: At least 20/40 acuity (Snellen) in each eye with or without correction. At least 70 degrees peripheral in horizontal meridian measured in each eye. The use of corrective lenses should be noted on the Medical Examiner's Certificate.	At least 70 degr dical Examiner's	ees periph Certificate	eral in horiz	ontal mer	idian
INSTRUCTIC ratio with 20 habitually we	INSTRUCTIONS: When other than the Snellen chart is us ratio with 20 as numerator and the smallest type read at 2 habitually wears contact lenses, or intends to do so while	ne Snellen chart is unallest type read at tends to do so while	used, give test re 20 feet as denon e driving, sufficier	sults in Snellen-con ninator. If the applic ntevidence of good	INSTRUCTIONS: When other than the Snellen chart is used, give test results in Snellen-comparable values. In recording distance vision, use 20 feet as normal. Report visual acuity as a ratio with 20 as numerator and the smallest type read at 20 feet as denominator. If the applicant wears corrective lenses, these should be worn while visual acuity is being tested. If the driver habitually wears contact lenses, or intends to do so while driving, sufficient evidence of good tolerance and adaptation to their use must be obvious. Monocular drivers are not qualified.	distance vision, u these should be ν their use must be α	ise 20 feet as vorn while vis obvious. <b>Mor</b>	s normal. Rep ualacuity is be n <b>ocular drive</b> l	ort visual ac eing tested. rs are not c	cuity as a If the drive <b>qualified.</b>
Numerical	Numerical readings must be provided.	ovided.			Applicant can recognize and distinguish among traffic control	and distinguish a	mong traffic	control	اا	Yes
ACUITY	UNCORRECTED	CORRECTED	HORIZONTAL	HORIZONTAL FIELD OF VISION	signals and devices showing standard red, green, and amber colors ?	owing standard rec	ı, green, and	amber colors		<b>8</b>
Right Eye	20/	20/	Right Eye	0	Applicant meets visual acuity requirement only when wearing:	acuity requirem	ent only wh	en wearing:		
Left Eye	20/	20/	Left Eye	0	Corrective Lenses	S				
Both Eyes	20/	20/			Monocular Vision: Yes	Yes No				
Complete n	Complete next line only if vision testing is done by a	esting is done by	an opthalmolog	n opthalmologist or optometrist						
Date of Examination	1	Name of Ophthalmologist or	or Optometrist (print)	print) Tel. No.	License	License No./ State of Issue	O	Signature	an an	
<b>INSTRUCT</b>	INSTRUCTIONS: To convert auditeduencies tested and divide by 3	Crieck in realing and used for tests. convert audiometric test results from ISO towide by 3.	from ISO to ANS	SI, -14 dB from ISO	INSTRUCTIONS: To convert audiometric test results from ISO to ANSI, -14 dB from ISO for 500Hz, -10dB for 1,000 Hz, -8.5 dB for 2000 Hz. To average, add the readings for 3 from less tested and divide by 3	ırd. İz, -8.5 dB for 2000	) Hz. To ave	To average, add the readings for 3	readings fo	r 3
Numerical re	Numerical readings must be recorded.	orded.				Right Ear	ar.	Left	Left Ear	
a) Record di	a) Record distance from individual at which	l at which Right ear	ear Left Ear	L	b) If audiometer is used, record hearing loss in	ig loss in 500 Hz	1000 Hz	2000 Hz 500	500 Hz 1000	1000 Hz 2000 Hz
torced whisp	forced whispered voice can first be heard.	e neard.	/ Feet	) Feet	cibels. (acc. to ANSI Z24.5-1951)	Average	.i.	Ave	Average:	
5. BLOOD	BLOOD PRESSURE/ PULSE RATE		nerical reading	is must be record	Numerical readings must be recorded. Medical Examiner should take at least two readings to confirm BP.	should take at le	ast two rea	adings to co	onfirm BP.	
Blood	Systolic Diastolic	lic Reading	D	Category	<b>Expiration Date</b>		Rec	Recertification		
Pressure		140-159/	66-06/6	Stage 1	1 year		۲ . ۲ .	1 year if <140/90	0.	:
Driver qua	Driver qualified if <140/90.		10.0				e 4	One-time certificate for 3 months if 141-159/91-99.	cate for 3 r	nonths if
Pulse Rate:	e: ☐ Regular ☐ Irregular	ılar 160-179/	9/100-109	Stage 2	One-time certificate for 3 months.	3 months.	<u> </u>	1 year from date of exam if ≤140/90	of exam i	f≤140/90
Record Pulse Rate:	Ise Bate	>180/11	10	Stage 3	6 months from date of exam if <140/90	xam if <140/90	6 m	6 months if < 140/90	06/0	
6. LABORA	LABORATORY AND OTHER TEST FINDINGS	TEST FINDINGS	Numerica	Numerical readings must be recorded	ecorded.	URINE SPECIMEN	SP. GR.	PROTEIN	BLOOD	SUGAR
Urinalysis is rerule out any ur	Urinalysis is required. Protein, blood or sugar in the urine may be an indication for further testing to trule out any underlying medical problem.	or sugar in the urine im.	may be an indic	ation for further test	ing to					
Other Testing	Other Testing (Describe and record)									

Middle,

First,

7. PHYSICAL EXAMINATION Height: (in.) Weight: (lbs.) Name:	Las
AMINATION	Name:
AMINATION	(lbs.)
AMINATION	(in.) Weigh
7. PHYSICAL EXAMINATION	Height:
7	PHYSICAL EXAMINATION
	7.

Even if a condition does not disqualify a driver, the medical examiner may consider deferring the driver temporarily. Also, the driver should be advised to take the necessary steps to correct the condition as soon as possible particularly if the condition, if neglected, could result in more serious illness that might affect driving. The presence of a certain condition may not necessarily disqualify a driver, particularly if the condition is controlled adequately, is not likely to worsen a is readily amenable to treatment.

Check YES if there are any abnormalities. Check NO if the body system is normal. Discuss any YES answers in detail in the space below, and indicate whether it would affect the driver's ability to operate a commercial motor vehicle safely. Enter applicable item number before each comment. If organic disease is present, note that it has been compensated for. See <u>Instructions to the Medical Examiner</u> for guidance.

<b>BODY SYSTEM</b>	CHECK FOR:	YES* NO	BODY SYSTEM	CHECK FOR: YES*	ON *S
1. General Appearance	Marked overweight, tremor, signs of alcoholism, problem drinking, or drug abuse.		7. Abdomen and Viscera	Enlarged liver, enlarged spleen, masses, bruits, hemia, sionificant abdominal wall muscle	
2. Eyes	Pupillary equality, reaction to light, accommodation, ocular motility, ocular muscle imbalance, extraocular movement, nystagmus, exophthalmos. Ask about retinopathy, cataracts,		8. Vascular System	weakness. Abnormal pulse and amplitude, cartoid or	
	aphakia, glaucoma, macular degeneration and refer to a specialist if appropriate.		9. Genito-urinary System	Hernias.	
3. Ears	Scarring of tympanic membrane, occlusion of external canal, perforated eardrums.		10. Extremities- Limb impaired. Driver may	Loss or impairment of leg, foot, toe, arm, hand, finger, Perceptible limp, deformities, atrophy, modernor anotherical chicking adoms	
4. Mouth and I hroat	Irremediable deformities likely to interfere with breathing or swallowing.		be subject to SPE certificate if otherwise qualified.	wear less, paralysis, chooling, ederna, hypotonia. Insufficient grasp and prehension in upper limb to maintain steering wheel grip. Insufficient mobility and strength in lower limb.	
5. Heart	Murmurs, extra sounds, enlarged heart, pacemaker, implantable defibrillator.		11. Spine, other	to operate pedals properly.  Previous surgery, deformities, limitation of	
6. Lungs and chest, not including breast examination	Abnormal chest wall expansion, abnormal respiratory rate, abnormal breath sounds including wheezes or alveolar rales, impaired respiratory function, cyanosis. Abnormal findings on physical exam may require further testing such as pulmonary tests and/ or xray of chest.		musculoskeletal 12. Neurological	Indidori, teridentess. Impaired equilibrium, coordination or speech pattern; asymmetric deep tendon reflexes, sensory or positional abnormalities, abnormal patellar and Babinki's reflexes, ataxia.	
*COMMENTS:					
Note certification star	Note certification status here. See Instructions to the Medical Examiner for guidance.	g.	☐ Wearing corrective lense	lense	
☐ Meets standards in 49 CF	Meets standards in 49 CFR 391.41; qualifies for 2 year certificate Does not meet standards		Accompanied by aexemption at time of certification	waiver/ exemption. Driver must present	present
☐ Meets standar  Driver qualified	Meets standards, but periodic monitoring required due to Driver qualified only for: ☐3 months ☐6 months ☐1 year ☐ Other		Skill Performance I  Driving within an of Qualified by opera	Skill Performance Evaluation (SPE) Certificate Driving within an exempt intracity zone (See 49 CFR 391.62) Qualified by operation of 49 CFR 391.64	

If meets standards, complete a Medical Examiner's Certificate as stated in 49 CFR 39143(h). (Driver must carry certificate when operating a commercial vehicle.)

Medical Examiner's signature Medical Examiner's name

Temporarily disqualified due to (condition or medication):

Return to medical examiner's office for follow up on

Telephone Number

# 49 CFR 391.41 Physical Qualifications for Drivers

## THE DRIVER'S ROLE

Responsibilities, work schedules, physical and emotional demands, and lifestyles among commercial drivers vary by the type of driving that they do. Some of the main types of drivers include the following: turn around or short relay (drivers return to their home base each evening); long relay (drivers drive 9-11 hours and then have at least a 10-hour offduty period), straight through haul (cross country drivers); and team drivers (drivers share the driving by alternating their 5-hour driving periods and 5-hour rest periods.)

cargo in order to compensate for the lost time; and environmental conditions such as excessive vibration, noise, and extremes in temperature. Transporting passengers or hazardous The following factors may be involved in a driver's performance of duties: abrupt schedule changes and rotating work schedules, which may result in irregular sleep patterns and a driver beginning a trip in a fatigued condition; long hours; extended time away from family and friends, which may result in lack of social support; tight pickup and delivery urregularity in work, rest, and eating patterns, adverse road, weather and traffic conditions, which may cause delays and lead to hurriedly loading or unloading materials may add to the demands on the commercial driver.

lifting heavy tarpaulins to cover open top trailers. The above tasks demand agility, the ability to bend and stoop, the ability to maintain a crouching position to inspect the underside There may be duties in addition to the driving task for which a driver is responsible and needs to be fit. Some of these responsibilities are: coupling and uncoupling trailer(s) from the tractor, loading and unloading trailer(s) (sometimes a driver may lift a heavy load or unload as much as 50,000 lbs. of freight after sitting for a long period of time without any stretching period); inspecting the operating condition of tractor and/or trailer(s) before, during and after delivery of cargo; lifting, installing, and removing heavy tire chains; and, of the vehicle, frequent entering and exiting of the cab, and the ability to climb ladders on the tractor and/or trailer(s).

In addition, a driver must have the perceptual skills to monitor a sometimes complex driving situation, the judgment skills to make quick decisions, when necessary, and the manipulative skills to control an oversize steering wheel, shift gears using a manual transmission, and maneuver a vehicle in crowded areas.

## §391.41 PHYSICAL QUALIFICATIONS FOR

- photographic copy, of a medical examiner's certificate that (a) A person shall not drive a commercial motor vehicle provided in §391.67, has on his person the original, or a unless he is physically qualified to do so and, except as he is physically qualified to drive a commercial motor
- (b) A person is physically qualified to drive a motor vehicle if that person:
  - Certificate (formerly Limb Waiver Program) pursuant to (1) Has no loss of a foot, a leg, a hand, or an arm, or has been granted a Skill Performance Evaluation (SPE)
- arm, foot, or leg which interferes with the ability to perform normal tasks associated with operating a commercial motor (3) Has no established medical history or clinical diagnosis thrombosis, or any other cardiovascular disease of a variety of diabetes mellitus currently requiring insulin for control; associated with operating a commercial motor vehicle; or known to be accompanied by syncope, dyspnea, collapse, vehicle; or any other significant limb defect or limitation which interferes with the ability to perform normal tasks interferes with prehension or power grasping; or (ii) An has been granted a SPE Certificate pursuant to §391.49. (2) Has no impairment of: (i) A hand or finger which (4) Has no current clinical diagnosis of myocardial infarction, angina pectoris, coronary insufficiency,
- (5) Has no established medical history or clinical diagnosis

- ability to control and drive a commercial motor vehicle of a respiratory dysfunction likely to interfere with his
- (6) Has no current clinical diagnosis of high blood pressure likely to interfere with his ability to operate a commercial motor vehicle safely.
- (7) Has no established medical history or clinical diagnosis neuromuscular, or vascular disease which interferes with his ability to control and operate a commercial motor of rheumatic, arthritic, orthopedic, muscular,
- (8) Has no established medical history or clinical diagnosis of epilepsy or any other condition which is likely to cause loss of consciousness or any loss of ability to control a commercial motor vehicle;
- or psychiatric disorder likely to interfere with his ability to (10) Has distant visual acuity of at least 20/40 (Snellen) in (9) Has no mental, nervous, organic, or functional disease drive a commercial motor vehicle safely;
- meridian in each eye, and the ability to recognize the colors corrective lenses, distant binocular acuity of at least 20/40 of traffic signals and devices showing standard red, green (Snellen) in both eyes with or without corrective lenses, separately corrected to 20/40 (Snellen) or better with field of vision of at least 70degrees in the horizontal each eye without corrective lenses or visual acuity
- ear not less than 5 feet with or without the use of a hearing (11) First perceives a forced whispered voice in the better aid, or, if tested by use of an audiometric device, does not

- have an average hearing loss in the better ear greater than 40 decibels at 500 Hz, 1,000 Hz and 2,000 Hz with or without a hearing device when the audiometric device is calibrated to the American National Standard (formerly ASA Standard) Z24.5-1951;
- 21 CFR 1308.11 Schedule I, an amphetamine, a narcotic, or is familiar with the driver's medical history and has advised licensed medical practitioner, as defined in § 382.107, who (12)(i) Does not use any controlled substance identified in (ii) Does not use any non-Schedule I controlled substance except when the use is pursuant to the instructions of a other habit-forming drug.
- (13) Has no current clinical diagnosis of alcoholism.

the driver that the substance will not adversely affect the

driver's ability to safely operate a commercial motor

# **INSTRUCTIONS TO THE MEDICAL EXAMINER**

## General Information

commerce according to the requirements in 49 CFR 391.41-49. Therefore, the medical examiner must be knowledgeable of these requirements and making the qualification determination. The medical examiner should be qualification to operate a commercial motor vehicle (CMV) in interstate guidelines developed by the FMCSA to assist the medical examiner in familiar with the driver's responsibilities and work environment and is The purpose of this examination is to determine a driver's physical referred to the section on the form, The Driver's Role.

common prescriptions and over-the-counter medications relative to the side to read warning labels on all medications. History of certain conditions may effects and hazards of these medications while driving. Educate the driver be cause for rejection, particularly if required by regulation, or may indicate conducting the physical examination, the medical examiner should discuss the need for additional laboratory tests or more stringent examination perhaps by a medical specialist. These decisions are usually made by the medical examiner in light of the driver's job responsibilities, work schedule In addition to reviewing the Health History section with the driver and and potential for the conditions to render the driver unsafe.

condition, if neglected, could develop into a serious illness that could affect Medical conditions should be recorded even if they are not cause for appropriate remedial care. This advice is especially needed when a denial, and they should be discussed with the driver to encourage

regulations, the certificate is valid for two years, unless the driver has medical examiner signs the medical certificate which the driver must carry If the medical examiner determines that the driver is fit to drive and is also able to perform non-driving responsibilities as may be required, the a medical condition that does not prohibit driving but does require information (a vision exemption, qualifying drivers under 49 CFR 391.64 should be done carefully and at least as complete as is indicated by the should be issued for a shorter length of time. The physical examination more frequent monitoring. In such situations, the medical certificate with his/her license. The certificate must be dated. Under current attached form. Contact the FMCSA at (202) 366-1790 for further

## Interpretation of Medical Standards

(FMCSA) has published recommendations called Advisory Criteria to help qualifications for commercial driving. These recommendations have been condensed to provide information to medical examiners that (1) is directly relevant to the physical examination and (2) is not already included in the medical examination form. The specific regulation is printed in italics and medical examiners in determining whether a driver meets the physical commercial drivers, the Federal Motor Carrier Safety Administration Since the issuance of the regulations for physical qualifications of it's reference by section is highlighted.

## Federal Motor Carrier Safety Regulations -Advisory Criteria-

## Loss of Limb: §391.41(b)(1)

A person is physically qualified to drive a commercial motor vehicle if that person

Has no loss of a foot, leg, hand or an arm, or has been granted a Skill Performance Evaluation (SPE) Certificate pursuant to Section 391.49.

## Limb Impairment:

## §391.41(b)(2)

A person is physically qualified to drive a commercial motor vehicle if that person:

with prehension or power grasping; or (ii) An arm, foot, or leg associated with operating a commercial motor vehicle; or (iii) Any other significant limb defect or limitation which interferes Has no impairment of: (i) A hand or finger which interferes operating a commercial motor vehicle, or (iv) Has been granted a Skill Performance Evaluation (SPE) Certificate pursuant to Section 391.49. which interferes with the ability to perform normal tasks with the ability to perform normal tasks associated with

commercial motor vehicle is subject to the Skill Performance A person who suffers loss of a foot, leg, hand or arm or whose limb impairment in any way interferes with the safe performance of normal tasks associated with operating a Evaluation Certification Program pursuant to section

still present, and thus restrictions may be included on individual SPE certificates when a State Director for the FMCSA determines operate a commercial motor vehicle. Since there are no medical equipment modifications have been developed to compensate impairment to qualify under the Federal Motor Carrier Safety aids equivalent to the original body or limb, certain risks are (formerly the Limb Waiver Program) was designed to allow 391.49, assuming the person is otherwise qualified. With the advancement of technology, medical aids and they are necessary to be consistent with safety and public persons with the loss of a foot or limb or with functional Regulations (FMCSRs) by use of prosthetic devices or for certain disabilities. The SPE Certification Program equipment modifications which enable them to safely interest

accompanied by a SPE certificate. The driver and the employing (391.41(b)(3) through (13)), the medical examiner must check operates a motor vehicle in interstate or foreign commerce without a curent SPE certificate for his/her physical disability. motor carrier are subject to appropriate penalty if the driver on the medical certificate that the driver is qualified only if If the driver is found otherwise medically qualified

## §391.41(b)(3)

A person is physically qualified to drive a commercial motor Has no established medical history or clinical diagnosis of vehicle if that person:

and space. Individuals who require insulin for control have diabetes mellitus currently requiring insulin for control.

Diabetes mellitus is a disease which, on occasion, can much or too little insulin, or food intake not consistent with (drowsiness, semiconsciousness, diabetic coma or insulin result in a loss of consciousness or disorientation in time conditions which can get out of control by the use of too symptoms of hyperglycemic or hypoglycemic reactions the insulin dosage. Incapacitation may occur from

complicated process requiring insulin, syringe, needle, alcohol sponge and a sterile technique. Factors related to the FMCSA has consistently held that a diabetic who uses long-haul commercial motor vehicle operations, such as stress, and concomitant illness, compound the dangers, insulin for control does not meet the minimum physical fatigue, lack of sleep, poor diet, emotional conditions, The administration of insulin is, within itself, a requirements of the FMCSRs.

prescribed for diabetic individuals to help stimulate natural controlled by the use of oral medication and diet, then an may call (202) 366-1790 for an application for a diabetes individual may be qualified under the present rule. CMV drivers who do not meet the Federal diabetes standard Hypoglycemic drugs, taken orally, are sometimes (See Conference Report on Diabetic Disorders and body production of insulin. If the condition can be exemption.

Commercial Drivers and Insulin-Using Commercial Motor Vehicle Drivers at:

http://www.fmcsa.dot.gov/rulesregs/medreports.htm)

## Cardiovascular Condition §391.41(b)(4)

A person is physically qualified to drive a commercial angina pectoris, coronary insufficiency, thrombosis or any Has no current clinical diagnosis of myocardial infarction, other cardiovascular disease of a variety known to be motor vehicle if that person:

accompanied by syncope, dyspnea, collapse or congestive

specifically designed to encompass: "a clinical diagnosis cardiovascular condition which has not fully stabilized The term "has no current clinical diagnosis of" is regardless of the time limit The term "known to be of" (1) a current cardiovascular condition, or (2) a cardiac failure.

accompanied by" is designed to include a clinical diagnosis cardiac failure; and/or (2) which is likely to cause syncope a cardiovascular disease (1) which is accompanied by symptoms of syncope, dyspnea, collapse or congestive dyspnea, collapse or congestive cardiac failure.

However, the subjective decision of whether the nature and severity of an individual's condition will likely cause electrocardiogram (ECG), no residual complications and no symptoms of cardiovascular insufficiency is on an individua basis and qualification rests with the medical examiner and certified that he or she have a normal resting and stress infarction, thrombosis, etc.), it is suggested before a driver syncope, dyspnea, collapse, or congestive cardiac failure. driver who has a current cardiovascular disease which is It is the intent of the FMCSRs to render unqualified, a physical limitations, and is taking no medication likely to occurrence of cardiovascular insufficiency (myocardial accompanied by and/or likely to cause symptoms of the motor carrier. In those cases where there is an interfere with safe driving.

underlying medical condition(s) which require treatment and of the driver and should not, by its use, medically disqualify medical treatment which can improve the health and safety recommendations regarding the physical qualification of the commercial driver. The emphasis should be on the unqualifying. Implantable cardioverter defibrillators are the general health of the driver. The FMCSA should be disqualifying due to risk of syncope. Coumadin is a Coronary artery bypass surgery and pacemaker implantation are remedial procedures and thus, not contacted at (202) 366-1790 for additional drivers on coumadin

(See Cardiovasular Advisory Panel Guidelines for the Medical examination of Commercial Motor Vehicle Drivers at: http://www.fmcsa.dot.gov/rulesregs/medreports.htm)

## Respiratory Dysfunction

## §391.41(b)(5)

Has no established medical history or clinical diagnosis of a A person is physically qualified to drive a commercial motor respiratory dysfunction likely to interfere with ability to vehicle if that person:

safety. Even the slightest impairment in respiration function under emergency conditions (when greater oxygen supply is necessary for performance) may be detrimental to safe Since a driver must be alert at all times, any change in his or her mental state is in direct conflict with highway control and drive a commercial motor vehicle safely. driving

optimum dose is achieved, provided lower extremity venous examiner detects a respiratory dysfunction, that in any way is likely to interfere with the driver's ability to safely control and drive a commercial motor vehicle, the driver must be referred to a specialist for further evaluation and therapy. Anticoagulation therapy for deep vein thrombosis and/or There are many conditions that interfere with oxygen examinations remain normal and the treating physician emphysema, chronic asthma, carcinoma, tuberculosis, pulmonary thromboembolism is not unqualifying once exchange and may result in incapacitation, including chronic bronchitis and sleep apnea. If the medical gives a favorable recommendation.

See Conference on Pulmonary/Respiratory Disorders http://www.fmcsa.dot.gov/rulesregs/medreports.htm and Commercial Drivers at:

## §391.41(b)(6)

A person is physically qualified to drive a commercial motor likely to interfere with ability to operate a commercial motor Has no current clinical diagnosis of high blood pressure vehicle if that person:

Hypertension alone is unlikely to cause sudden collapse; Cardiovascular Advisory Guidelines for the Examination of damage, particularly cerebral vascular disease, is present. CMV Drivers, which used the Sixth Report of the Joint National Committee on Detection, Evaluation, and however, the likelihood increases when target organ This regulatory criteria is based on FMCSA's Treatment of High Blood Pressure (1997).

thereafter and should be at or less than 140/90. If less than 140-159 mmHg and/or a diastolic BP of 90-99 mmHg. The Stage 1 hypertension corresponds to a systolic BP of hypertension-related acute incapacitation and may be 160/100, certification may be extended one time for 3 Certification examinations should be done annually medically certified to drive for a one-year period. driver with a BP in this range is at low risk for months.

driver demonstrates a BP value of 140/90 or less, he or she diastolic is considered Stage 2 hypertension, and the driver pressure to less than or equal to 140/90. A blood pressure in this range is an absolute indication for anti-hypertensive Provided treatment is well tolerated and the may be certified for one year from date of the initial exam. A blood pressure of 160-179 systolic and/or 100-109 institution of treatment. The driver is given a one time certification of three months to reduce his or her blood is not necessarily unqualified during evaluation and The driver is certified annually thereafter drug therapy.

110 (diastolic) is considered Stage 3, high risk for an acute BP-related event. The driver may **not** be qualified, even temporarily, until reduced to 140/90 or less and treatment is well tolerated. The driver may be certified for 6 months and biannually (every 6 months) thereafter if at recheck BP is A blood pressure at or greater than 180 (systolic) and 140/90 or less.

examiner does not know the severity of hypertension prior Annual recertification is recommended if the medical to treatment.

An elevated blood pressure finding should be confirmed by at least two subsequent measurements on different

hazards of these medications while driving. Side effects of have side effects, the importance of which must be judged other risk factors. Most antihypertensive medications also pharmacologic modalities as well as counseling to reduce on an individual basis. Individuals must be alerted to the somnolence or syncope are particulary undesirable in Freatment includes nonpharmacologic and

Evaluation is warranted if patient is persistently hypertensive Secondary hypertension is based on the above stages.

Some causes of secondary hypertension may be amenable to surgical on maximal or near-maximal doses of 2-3 pharmacologic agents. See Cardiovascular Advisory Panel Guidelines for the Medical Examination of Commercial Motor Vehicle Drivers at: http://www.fmcsa.dot.gov/rulesregs/medreports.htm) ntervention or specific pharmacologic disease

## Neuromuscular or Vascular Disease §391.41(b)(7) Rheumatic, Arthritic, Orthopedic, Muscular,

A person is physically qualified to drive a commercial motor vehicle if

arthritic, orthopedic, muscular, neuromuscular or vascular disease which Has no established medical history or clinical diagnosis of rheumatic, interferes with the ability to control and operate a commercial motor vehicle safely

sensations (paresthesia), decreased muscular tone (hypotonia), visual have more insidious onsets and display symptoms of muscle wasting eventually interfere with the ability to safely operate a motor vehicle. Certain diseases are known to have acute episodes of transient disturbances and pain which may be suddenly incapacitating. With pronounced and remain for longer periods of time. Other diseases many instances these diseases are degenerative in nature or may muscle weakness, poor muscular coordination (ataxia), abnormal incapacitate a person but may restrict his/her movements and each recurring episode, these symptoms may become more (atrophy), swelling and paresthesia which may not suddenly result in deterioration of the involved area.

present initially but may manifest itself over time); and (4) the likelihooc required, a certificate for a shorter period of time may be issued. (See then he/she has an established history of that disease. The physician or loss of strength); (2) the degree of limitation present (such as range nature and severity of the individual's condition (such as sensory loss when examining an individual, should consider the following: (1) the of sudden incapacitation. If severe functional impairment exists, the driver does not qualify. In cases where more frequent monitoring is Conference on Neurological Disorders and Commercial Drivers at: Once the individual has been diagnosed as having a rheumatic, arthritic, orthopedic, muscular, neuromuscular or vascular disease, of motion); (3) the likelihood of progressive limitation (not always http://www.fmcsa.dot.gov/rulesregs/medreports.htm)

## Epilepsy §391.41(b)(8)

A person is physically qualified to drive a commercial motor vehicle if that person.

Has no established medical history or clinical diagnosis of epilepsy or any other condition which is likely to cause loss of consciousness or any loss of ability to control a motor vehicle.

consciousness or any loss of ability to control a motor vehicle.

Epilepsy is a chronic functional disease characterized by seizures or episodes that occur without warning, resulting in loss of voluntary control which may lead to loss of consciousness and/or seizures. Therefore, the following drivers cannot be qualified: (1) a driver who has a medical history of epilepsy; (2) a driver who has a current clinical diagnosis of epilepsy; or (3) a driver who is taking

If an individual has had a sudden episode of a nonepileptic seizure or loss of consciousness of unknown cause which did not require antiseizure medication, the decision as to whether that person's condition will likely cause loss of consciousness or loss of ability to control a motor vehicle is made on an individual basis by the medical examiner in consultation with the treating physician. Before certification is considered, it is suggested that a 6 month waiting period elapse from the time of the episode. Following the waiting period, it is suggested that the individual have a complete neurological examination. If the results of the examination are negative and antiseizure medication is not required, then the driver may be qualified.

In those individual cases where a driver has a seizure or an episode of loss of consciousness that resulted from a known medical condition (e.g., drug reaction, high temperature, acute infectious disease, dehydration or acute metabolic disturbance), certification should be deferred until the driver has fully recovered from that condition and has no existing residual complications, and not taking antiseizure medication.

Drivers with a history of epilepsy/seizures off antiseizure medication and seizure-free for 10 years may be qualified to drive a CMV in interstate commerce. Interstate drivers with a history of a single unprovoked seizure may be qualified to drive a CMV in Interstate commerce if seizure-free and off antiseizure medication for a 5-year period or more.

(See Conference on Neurological Disorders and Commercial

http://www.fmcsa.dot.gov/rulesregs/medreports.htm)

## Mental Disorders §391.41(b)(9)

A person is physically qualified to drive a commercial motor vehicle if that person:

Has no mental, nervous, organic or functional disease or psychiatric disorder likely to interfere with ability to drive a motor vehicle safely.

Emotional or adjustment problems contribute directly to an individual's level of memory, reasoning, attention, and judgment These problems often underlie physical disorders. A variety of functional disorders can cause drowsiness, dizziness, confusion, weakness or paralysis that may lead to incoordination, inattention, loss of functional control and susceptibility to accidents while driving. Physical fatigue, hadadache, impaired coordination, recurring physical fatigue, and chronic "nagging" pain may be present to such a degree that certification for commercial driving is inadvisable. Somatic and psychosomatic complaints should be thoroughly examined when determining an individual's overall fitness to drive. Disorders of a periodically incapacitating nature, even in the early stages of development, may warrant disqualification.

Many bus and truck drivers have documented that "nervous trouble" related to neurotic, personality, or emotional or adjustment problems is responsible for a significant fraction of their preventable accidents. The degree to which an individual is able to appreciate, evaluate and adequately respond to environmental strain and emotional stress is critical when assessing an individual's mental alertness and flexibility to cope with the stresses of commercial motor vehicle driving.

When examining the driver, if should be kept in mind that individuals who live under chronic emotional upsets may have deeply ingrained maladaptive or erraits behavior patterns. Excessively antagonistic, instinctive, impulsive, openly aggressive, paranoid or severely depressed behavior greatly interfere with the driver's ability to drive safely. Those emotional instability (schizophrenia, affective psychoses, paranoia, anxiety or depressive neuroses) may warrant or emotional instability (schizophrenia, affective psychoses, paranoia, anxiety or depressive neuroses) may warrant or exigualification. Careful consideration should be given to the side effects and interactions of medications in the overall qualification determination. See Psychiatric Conference Report for specific recommendations on the use of medications and potential hazards for driving.

Vision

## §391.41(b)(10)

A person is physically qualified to drive a commercial motor vehicle if that person:

Has distant visual acuity of at least 20/40 (Snellen) in each eye with or without corrective lenses or visual acuity separately corrected to 20/40 (Snellen) or better with corrective lenses, distant binocular acuity of at least 20/40 (Snellen) in both eyes with or without corrective lenses, field of vision of at least 70 degrees in the horizontal meridian in each eye, and the ability to recognize the colors of traffic signals and devices showing standard red, green, and amber.

The term "ability to recognize the colors of" is interpreted to mean if a person can recognize and distinguish among traffic control signals and devices showing standard, evel, green and amber, he or she meets the minimum standard, even though he or she may have some type of color perception deficiency. If certain color perception tests are administered, (such as Ishinara, Pseudoisochromatic, Yarn) and doubtful findings are discovered, a controlled test using signal red, green and amber may be employed to determine the driver's ability to recognize these colors.

to indicate that the driver has good tolerance and is well adapted to their use. Use of a contact lens in one eye for nealistance visual acuity and another lens in the other eye for nealistance visual acuity and another lens in the other eye for nealistance visual acuity and another lens in the other eye for nealistance is not acceptable, nor telescopic lenses acceptable for the driving of commercial motor vehicles. If an individual meets the criteria by the use of glasses or contact lenses, the following statement shall appear on the Medical Examiner's Certificate: "Qualified only if wearing

Contact lenses are permissible if there is sufficient evidence

corrective lenses."

CMV drivers who do not meet the Federal vision standard may call (202) 366-1790 for an application for a vision exemption.

(See Visual Disorders and Commercial Drivers at: http://www.fmcsa.dot.gov/rulesregs/medreports.htm)

Hearing

## §391.41(b)(11)

A person is physically qualified to drive a commercial motor vehicle if that person:

First perceives a forced whispered voice in the better ear at not less than 5 feet with or without the use of a hearing aid, or, if tested by use of an audiometric device, does not have an everage hearing loss in the better ear greater than 40 decibels at 500 Hz, 1,000 Hz, and 2,000 Hz with or without a hearing aid when the audiometric devices is calibrated to American National

http://www.fmcsa.dot.gov/rulesregs/medreports.htm

Drivers at:

Standard (formerly ADA Standard) Z24.5-1951.
Since the prescribed standard under the FMCSRs is the American Standards Association (ANSI), it may be necessary to convert the audiometric results from the ISO standard to the ANSI standard. Instructions are included on the Medical Examination report form.

fan individual meets the criteria by using a hearing aid, the driver must wear that hearing aid and have it in operation at all times while driving. Also, the driver must be in possession of a spare power source for the hearing aid.

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For the whispered voice test, the individual should be stationed at least 5 feet from the examiner with the ear being tested furned toward the examiner. The other ear is covered. Using the breath which remains after a normal expiration, the examiner whispers words or random numbers such as 66, 18,

23, etc. The examiner should not use only sibilants (s sounding materials). The opposite ear should be tested in the same manner. If the individual fails the whispered voice test, the audiometric test should be administered.

If an individual meets the criteria by the use of a hearing aid, the following statement must appear on the Medical Examiner's Certificate "Qualified only when wearing a hearing aid." (See Hearing Disorders and Commercial Motor Vehicle Drivers at:

http://www/fmcsa.dot.gov/rulesregs/medreports.htm)

## **Drug Use**

## §391.41(b)(12)

A person is physically qualified to drive a commercial motor vehicle if that person does not use any controlled substance identified in 21 CFR 1308.11, an amphetamine, a narcotic, or other habit-forming drug. A driver may use a non-Schedule I substance or drug, if the substance or drug, if the substance or drug is prescribed by a licensed medical practitioner who: (A) is familiar with the driver's medical history, and assigned duties; and (B) has advised the driver that the prescribed substance or drug will not adversely affect the driver's ability to safely operate a commercial motor vehicle.

This exception does not apply to methadone. The intent of the medical certification process is

to medically evaluate a driver to ensure that the driver has no medical condition which interferes with the safe performance of driving tasks on a public road. If a driver uses a Schedule I drug or other substance, an amphetamine, a narcotic, or any other habit-forming drug, it may be cause for the driver to be found medically unqualified. Motor carriers are encouraged to obtain a practitioner's written statement about the effects on transportation safety of the use of a particular drug.

A test for controlled substances is not required as part of this biennial certification process. The FMCSA or the driver's employer should be contacted directly for information on controlled substances and alcohol testing under Part 382 of the FMCSRs.

The term "uses" is designed to encompass instances of prohibited drug use determined by a physician through established medical means. This may or may not involve body fluid testing. If body fluid testing takes place, positive test results should be confirmed by a second test of greater specificity. The term "habit-forming" is intended to include any drug or medication generally recognized as capable of becoming habitual, and which may impair the user's ability to operate a commercial motor vehicle safely.

The driver is medically unqualified for the duration of the prohibited drug(s) use and until second examination shows the driver is free

from the prohibited drug(s) use. Recertification may involve a substance abuse evaluation, the successful completion of a drug rehabilitation program, and a negative drug test result. Additionally, given that the certification period is normally two years, the examiner has the option to certify for a period of less than 2 years if this examiner determines more frequent monitoring is required.

(See Conference on Neurological Disorders and Commercial Drivers and Conference on Psychiatric Disorders and Commercial Drivers at:

http://www.fmcsa.dot.gov/rulesregs/medreports.

htm)

## Alcoholism §391.41(b)(13)

A person is physically qualified to drive a commercial motor vehicle if that person:

Has no current clinical diagnosis of alcoholism

The form "ourset clinical diagnosis of" is

The term "current clinical diagnosis of" is specifically designed to encompass a current alcoholic illness or those instances where the individual's physical condition has not fully stabilized, regardless of the time element. If an individual shows signs of having an alcohol-use problem, he or she should be referred to a specialist. After counseling and/or treatment, he or she may be considered for certification.

BILLING CODE 4910-EX-C

Issued on: July 5, 2011.

## William Bronrott,

Deputy Administrator.

[FR Doc. 2011-17192 Filed 7-7-11; 8:45 am]

BILLING CODE 4910-EX-P

## **DEPARTMENT OF TRANSPORTATION**

## **Federal Railroad Administration**

49 CFR Chapter II [Docket No. FRA-2009-0038]

## **Risk Reduction Program**

RIN 2130-AC11

**AGENCY:** Federal Railroad Administration (FRA), Department of

Transportation (DOT).

**ACTION:** Notice of public hearings.

**SUMMARY:** FRA is announcing public hearings to provide interested persons an opportunity to discuss the development of a regulation requiring certain railroads to develop a Risk Reduction Program (RRP). The Rail Safety Improvement Act of 2008 requires the development and implementation of railroad safety risk reduction programs. Risk reduction is a comprehensive, system-oriented approach to safety that (1) determines an operation's level of risk by identifying and analyzing applicable hazards and (2) develops plans to mitigate that risk. Each RRP is statutorily required to be supported by a risk analysis and a Risk Reduction Program Plan (RRPP), which must include a Technology Implementation Plan and a Fatigue Management Plan.

**DATES:** To encourage participation, two public hearings will be held. A public hearing will be held on July 19, 2011, in Chicago, and a public hearing will be held on July 21, 2011, in Washington, DC. At both locations, the times of the public hearings will be from 9 a.m. to 4 p.m.

ADDRESSES: Public Hearings. The public hearing in Chicago will be held at the W Chicago City Center Hotel located at 172 West Adams, in the Great Room I, Plateau. The public hearing in Washington, DC, will be held at the Doubletree Hotel located at 1515 Rhode Island Avenue, NW., in the Terrace Ballroom.

Attendance: Any persons wishing to make a statement at the hearing should notify FRA's Docket Clerk, Michelle Silva, by telephone, e-mail, or in writing, at least five business days before the date of the hearing. Ms.

Silva's contact information is as follows: FRA, Office of Chief Counsel, Mail Stop 10, 1200 New Jersey Avenue, SE., Washington, DC 20590; *telephone*: 202–493–6030; *e-mail*:

michelle.silva@dot.gov. For information on facilities or services for persons with disabilities or to request special assistance at the meetings, please contact by telephone or e-mail as soon as possible, Wendy A. Noble Burns at 202–493–6304 or wendy.noble@dot.gov.

## FOR FURTHER INFORMATION CONTACT:

Miriam Kloeppel, Staff Director, Risk Reduction Program Division, Office of Safety Analysis, FRA, 1200 New Jersey Avenue, SE., Mail Stop 25, Washington, DC 20590; telephone: 202–493–6224; e-mail: miriam.kloeppel@dot.gov; or Matthew L. Navarrete, Trial Attorney, Office of Chief Counsel, FRA, 1200 New Jersey Avenue, SE., Mail Stop 10, Washington, DC 20590; telephone: 202–493–0138; e-mail:

matthew.navarrete@dot.gov.

**SUPPLEMENTARY INFORMATION:** Interested parties are invited to present oral statements and to proffer information and views at the hearings. The hearings will be informal and will be conducted by a representative designated by FRA in accordance with FRA's Rules of Practice (49 CFR 211.25). The hearings will be non-adversarial proceedings; therefore, there will be no cross examination of persons presenting statements or proffering evidence. An FRA representative will make an opening statement outlining the scope of each hearing. After all initial statements have been completed, those persons wishing to make a brief rebuttal will be given the opportunity to do so in the same order in which the initial statements were made. Additional procedures, as necessary for the conduct of the hearings, will be announced at the hearings. The purpose of these hearings is to receive oral comments in response to an Advanced Notice of Proposed Rulemaking (ANPRM) that requested public comment on a potential risk reduction rulemaking. See 75 FR 76345-76351, Dec. 8, 2010. A transcript of the discussions will be made part of the public docket in this proceeding.

Public Participation Procedures. Any person wishing to participate in one of the public hearings should notify the Docket Clerk by mail or at the address or fax number provided in the Attendance section at least five working days prior to the date of the hearing and submit three copies of the oral statement that he or she intends to make at the proceeding. The notification should identify the party the person represents,

the particular subject(s) the person plans to address, and the time requested. The notification should also provide the Docket Clerk with the participant's mailing address and other contact information. FRA reserves the right to limit participation in the hearings of persons who fail to provide such notification. FRA reserves the right to limit the duration of presentations if necessary to afford all persons with the opportunity to speak.

## **Background**

In § 103 of the Rail Safety
Improvement Act of 2008, Public Law
110–432, 122 Stat. 4854 (Oct. 16, 2008)
(codified at 49 U.S.C. 20156)
(hereinafter RSIA), Congress directed
the Secretary of Transportation to issue
a regulation by October 16, 2012,
requiring certain railroads to develop an
RRP. While the statute vests certain
responsibilities with the Secretary of the
U.S. DOT (Secretary), the Secretary has
since delegated those responsibilities to
the FRA Administrator. See 49 CFR
1.49(00); 74 FR 26981 (June 5, 2009); see
also 49 U.S.C. 103(g).

Each railroad subject to the regulation would have to develop and implement an RRP approved by FRA. See 49 U.S.C. 20156(a)(1). This RRP is required to be supported by an RRPP. See 49 U.S.C. 20156(d)(2). FRA would conduct an annual review to ensure that each railroad has complied with its RRP. See 49 U.S.C. 20156(a)(3). The RSIA mandates that the following three categories of railroads be required to develop and implement an FRA-approved RRP:

- (1) Class I railroads;
- (2) Railroad carriers with inadequate safety performance, as determined by the Secretary; and
- (3) Railroad carriers that provide intercity rail passenger or commuter rail passenger transportation (passenger railroads).

See 49 U.S.C. 20156(a)(1).

Railroads not required to implement RRPs under the RSIA would be permitted to voluntarily submit plans meeting the requirements of any final RRP regulation for FRA review and approval. See 49 U.S.C. 20156(a)(4).

On December 8, 2010, FRA published an ANPRM soliciting public comment on how FRA can best develop a risk reduction regulation based upon the RSIA's requirements. See 75 FR 76345—76351. The ANPRM discussed certain major components that must be included in the final rule under the RSIA and identified various approaches that FRA could take in developing the rule. The purpose of these hearings is to