aspiration or bone marrow biopsy, but not peripheral blood smears alone. Examples of these disorders are myelodysplastic syndromes, aplastic anemia, granulocytopenia, and myelofibrosis. Acquired disorders of bone marrow failure may result from viral infections, chemical exposure, or immunologic disorders.

- 2. The hospitalizations in 107.10A do not all have to be for the same complication of bone marrow failure. They may be for three different complications of the disorder. Examples of complications that may result in hospitalization include uncontrolled bleeding, anemia, and systemic bacterial, viral, or fungal infections.
- 3. For 107.10B, the requirement of life-long RBC transfusions to maintain life in myelodysplastic syndromes or aplastic anemias has the same meaning as it does for beta thalassemia major. (See 107.00C4.)
- F. How do we evaluate bone marrow or stem cell transplantation under 107.17?

We will consider you to be disabled for 12 months from the date of bone marrow or stem cell transplantation, or we may consider you to be disabled for a longer period if you are experiencing any serious post-transplantation complications, such as graft-versus-host (GVH) disease, frequent infections after immunosuppressive therapy, or significant deterioration of organ systems. We do not restrict our determination of the onset of disability to the date of the transplantation in 107.17. We may establish an earlier onset of disability due to your transplantation if evidence in your case record supports such a finding.

G. How do we consider your symptoms, including your pain, severe fatigue, and malaise?

Your symptoms, including pain, severe fatigue, and malaise, may be important factors in our determination whether your hematological disorder meets or medically equals a listing, or in our determination whether you otherwise have marked and severe functional limitations. We cannot consider your symptoms unless you have medical signs or laboratory findings showing the existence of a medically determinable impairment(s) that could reasonably be expected to produce the symptoms. If you have such an impairment(s), we will evaluate the intensity, persistence, and functional effects of your symptoms using the rules throughout 107.00 and in our other regulations. (See sections 416.928 and 416.929 of this chapter.) Additionally, when we assess the credibility of your complaints about your symptoms and their functional effects, we will not draw any inferences from the fact that you do not receive treatment or that you are not following treatment without considering all of the relevant evidence in your case record, including any explanations you provide on why you are not receiving or following treatment.

H. How do we evaluate episodic events in hematological disorders?

Some of the listings in this body system require a specific number of events within a consecutive 12-month period. (See 107.05, 107.08, and 107.10A.) When we use such

criteria, a consecutive 12-month period means a period of 12 consecutive months, all or part of which must occur within the period we are considering in connection with your application or continuing disability review. These events must occur at least 30 days apart to ensure that we are evaluating separate events.

I. How do we evaluate hematological disorders that do not meet one of these listings?

- 1. These listings are only common examples of hematological disorders that we consider severe enough to result in marked and severe functional limitations. If your disorder does not meet the criteria of any of these listings, we must consider whether you have a disorder that satisfies the criteria of a listing in another body system. For example, we will evaluate hemophilic joint deformity under 101.00; polycythemia vera under 103.00, 104.00, or 111.00; chronic iron overload resulting from repeated RBC transfusion (transfusion hemosiderosis) under 103.00, 104.00, or 105.00; and the effects of intracranial bleeding or stroke under 111.00 or 112.00.
- 2. If you have a severe medically determinable impairment(s) that does not meet a listing, we will determine whether your impairment(s) medically equals a listing. (See section 416.926 of this chapter.) Hematological disorders may be associated with disorders in other body systems, and we consider the combined effects of multiple impairments when we determine whether they medically equal a listing. If your impairment(s) does not medically equal a listing, we will also consider whether it functionally equals the listings. (See section 416.926a of this chapter.) We use the rules in § 416.994a of this chapter when we decide whether you continue to be disabled.

107.01 Category of Impairments, Hematological Disorders

107.05 *Hemolytic anemias,* including sickle cell disease, thalassemia, and their variants (see 107.00C), with:

A. Documented painful (vaso-occlusive) crises requiring parenteral (intravenous or intramuscular) narcotic medication, occurring at least six times within a 12-month period with at least 30 days between crises.

OR

B. Complications of hemolytic anemia requiring at least three hospitalizations within a 12-month period and occurring at least 30 days apart. Each hospitalization must last at least 48 hours, which can include hours in a hospital emergency department or comprehensive sickle cell disease center immediately before the hospitalization (see 107.00C2).

OR

C. Hemoglobin measurements of 7.0 grams per deciliter (g/dL) or less, occurring at least three times within a 12-month period with at least 30 days between measurements.

OR

D. Beta thalassemia major requiring lifelong RBC transfusions at least once every 6 weeks to maintain life (see 107.00C4).

107.08 Disorders of thrombosis and hemostasis, including hemophilia and

thrombocytopenia (see 107.00D), with complications requiring at least three hospitalizations within a 12-month period and occurring at least 30 days apart. Each hospitalization must last at least 48 hours, which can include hours in a hospital emergency department or comprehensive hemophilia treatment center immediately before the hospitalization (see 107.00D2).

107.10 Disorders of bone marrow failure, including myelodysplastic syndromes, aplastic anemia, granulocytopenia, and myelofibrosis (see 107.00E), with:

A. Complications of bone marrow failure requiring at least three hospitalizations within a 12-month period and occurring at least 30 days apart. Each hospitalization must last at least 48 hours, which can include hours in a hospital emergency department immediately before the hospitalization (see 107.00E2).

OR

B. Myelodysplastic syndromes or aplastic anemias requiring life-long RBC transfusions at least once every 6 weeks to maintain life (see 107.00E3).

107.17 Hematological disorders treated by bone marrow or stem cell transplantation (see 107.00F). Consider under a disability for at least 12 consecutive months from the date of transplantation. After that, evaluate any residual impairment(s) under the criteria for the affected body system.

[FR Doc. 2015–08849 Filed 4–16–15; 8:45 am]

DEPARTMENT OF THE TREASURY

Internal Revenue Service

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26 CFR Part 1

[TD 9674]

RIN 1545-BM07

Guidelines for the Streamlined Process of Applying for Recognition of Section 501(c)(3) Status; Correction

AGENCY: Internal Revenue Service (IRS), Treasury.

ACTION: Final rule; correction.

SUMMARY: This document contains a correction to final and temporary regulations (TD 9674) that were published in the **Federal Register** on Wednesday, July 2, 2014 (79 FR 37630). The final and temporary regulations provide guidance to eligible organizations seeking recognition of taxexempt status under section 501(c)(3) of the Internal Revenue Code.

DATES: This correction is effective April 17, 2015 and applicable July 2, 2014.

FOR FURTHER INFORMATION CONTACT: James R. Martin and Robin Ehrenberg, at (202) 317–5800 (not a toll free number).

SUPPLEMENTARY INFORMATION:

Background

The final and temporary regulations (TD 9674) that are the subject of this correction are under section 501(c)(3) of the Internal Revenue Code.

Need for Correction

As published, the final and temporary regulation (TD 9674) contains an error and is in need of clarification.

Correction of Publication

In FR Doc. 2014–15623 appearing on page 37630 in the **Federal Register** of Wednesday, July 2, 2014, the following correction is made:

§1.508-1T [Corrected]

On page 37632, the amendatory instruction reading "Par. 7. Section 1.508–1T is revised to read as follows: " is corrected to read "Par. 7. Section 1.508–1T is added to read as follows:".

Martin V. Franks,

Chief, Publications and Regulations Branch, Legal Processing Division, Associate Chief Counsel (Procedure and Administration). [FR Doc. 2015–08856 Filed 4–16–15; 8:45 am]

BILLING CODE 4830-01-P

ENVIRONMENTAL PROTECTION AGENCY

40 CFR Part 52

[EPA-R04-OAR-2014-0867; FRL-9926-41-Region-4]

Approval and Promulgation of Implementation Plans; Alabama: Non-Interference Demonstration for Federal Low-Reid Vapor Pressure Requirement for the Birmingham Area

AGENCY: Environmental Protection

Agency (EPA).

ACTION: Final rule.

SUMMARY: The Environmental Protection Agency (EPA) is approving the State of Alabama's November 14, 2014, State Implementation Plan (SIP) revision, submitted through the Alabama Department of Environmental Management (ADEM), in support of the State's request that EPA change the Federal Reid Vapor Pressure (RVP) requirements for Jefferson and Shelby Counties (hereinafter referred to as the "Birmingham Area" or "Area"). Alabama's November 14, 2014, SIP revision evaluates whether changing the Federal RVP requirements in this Area would interfere with the Area's ability to meet the requirements of the Clean Air Act (CAA or Act). Specifically, Alabama's SIP revision concludes that relaxing the Federal RVP requirement

from 7.8 pounds per square inch (psi) to 9.0 psi for gasoline sold between June 1 and September 15 of each year in the Area would not interfere with attainment or maintenance of the national ambient air quality standards (NAAQS) or with any other CAA requirement. EPA has determined that Alabama's November 14, 2014, SIP revision is consistent with the CAA.

DATES: This rule will be effective April 17, 2015.

ADDRESSES: EPA has established a docket for this action under Docket Identification No. EPA-R04-OAR-2014-0867. All documents in the docket are listed on the www.regulations.gov Web site. Although listed in the index, some information is not publicly available, i.e., Confidential Business Information or other information whose disclosure is restricted by statute. Certain other material, such as copyrighted material, is not placed on the Internet and will be publicly available only in hard copy form. Publicly available docket materials are available either electronically through www.regulations.gov or in hard copy at the Air Regulatory Management Section (formerly the Regulatory Development Section), Air Planning and Implementation Branch (formerly the Air Planning Branch), Air, Pesticides and Toxics Management Division, U.S. Environmental Protection Agency, Region 4, 61 Forsyth Street SW., Atlanta, Georgia 30303-8960. EPA requests that if at all possible, you contact the person listed in the FOR **FURTHER INFORMATION CONTACT** section to schedule your inspection. The Regional Office's official hours of business are Monday through Friday, 8:30 a.m. to 4:30 p.m., excluding Federal holidays.

FOR FURTHER INFORMATION CONTACT: Richard Wong of the Air Regulatory Management Section, in the Air Planning and Implementation Branch, Air, Pesticides and Toxics Management Division, U.S. Environmental Protection Agency, Region 4, 61 Forsyth Street SW., Atlanta, Georgia 30303–8960. Mr. Wong may be reached by phone at (404) 562–8726 or via electronic mail at wong.richard@epa.gov.

SUPPLEMENTARY INFORMATION:

I. What is the background for this final action?

The Birmingham Area was originally designated as a 1-hour ozone nonattainment area by EPA on March 3, 1978 (43 FR 8962). A 7.8 psi Federal RVP requirement was first applied to the Area during the high ozone season given its status as a marginal nonattainment area for the 1-hour ozone

standard. Subsequently, in order to comply with the 1-hour ozone NAAQS, Alabama opted to implement a state RVP requirement of 7.0 psi for gasoline sold in the Birmingham Area during the high ozone season. EPA incorporated the state RVP requirement of 7.0 psi for gasoline sold in the Birmingham Area into the Alabama SIP on November 7, 2001. See 66 FR 56218. The Area attained the 1-hour ozone NAAQS and was redesignated to attainment for the 1-hour ozone on March 12, 2004, based on 2001-2003 ambient air quality monitoring data. See 69 FR 11798. Alabama's 1-hour ozone redesignation request did not include a request to remove the 7.0 psi state RVP requirement for the Birmingham Area from the SIP nor a request to relax the 7.8 psi Federal RVP standard.

On April 30, 2004, EPA designated and classified areas for the 8-hour ozone NAAQS that was promulgated on July 18, 1997, as unclassifiable/attainment or nonattainment for the new 8-hour ozone NAAQS. See 69 FR 23857. The Birmingham Area was designated as nonattainment for the 1997 8-hour ozone NAAQS with a design value of 0.087 parts per million (ppm). The Area was redesignated to attainment for the 1997 8-hour ozone NAAQS in a final rulemaking on May 12, 2006. See 71 FR 27631. Alabama's 1997 8-hour ozone redesignation request did not include a request for the removal of the 7.8 psi Federal RVP standard, nor did it include a request to change the 7.0 psi state RVP requirement for the Birmingham Area. However, to support its request for redesignation to attainment for the 1997 8-hour ozone NAAQS, Alabama took a conservative approach and estimated emissions using a 9.0 psi RVP in its modeling supporting the State's maintenance demonstration.

On March 2, 2012, Alabama submitted a SIP revision requesting that EPA remove the State's 7.0 psi RVP requirement for the Area from the SIP. EPA approved Alabama's March 2, 2012, SIP revision on April 20, 2012. See 77 FR 23619. In EPA's final rulemaking to remove the State RVP requirement, EPA noted that the action did not remove the 7.8 psi Federal RVP requirement for the Birmingham Area. Effective July 20, 2012, EPA designated the Birmingham Area as unclassifiable/ attainment for the 2008 8-hour ozone NAAQS. See 77 FR 30088 (April 30, 2012). Although the Birmingham Area is designated as attainment, the federal 7.8 psi RVP requirement remains in place.

Alabama is now requesting that EPA remove the federal 7.8 psi RVP requirement for the Birmingham Area, and it submitted a SIP revision on