the Superfund Amendments and Reauthorization Act of 1986 (SARA), § 104(i)(3), (42 U.S.C. 9604(i)(3)), directs the ATSDR Administrator to prepare Toxicological Profiles of Priority hazardous substances and, as necessary, to revise and publish each updated toxicological profile.

Comments can include additional information or reports on studies about the health effects of Set 27 substances. Although ATSDR considered key studies for each of these substances during the profile development process, the Federal Register notice solicits any relevant, additional studies, particularly unpublished data. ATSDR will evaluate the quality and relevance of such data or studies for possible inclusion into the profile. ATSDR remains committed to providing a public comment period for this document as a means to best serve public health and our clients.

DATES: Written comments on this draft Toxicological Profile must be received on or before January 14, 2016.

ADDRESSES: You may submit comments, identified by docket number ATSDR–2015–0002, by any of the following methods:


Instructions: All submissions received must include the agency name and docket number for this notice. All relevant comments will be posted without change. Because all public comments regarding ATSDR Toxicological Profiles are available for public inspection, no confidential business information or other confidential information should be submitted in response to this notice.

FOR FURTHER INFORMATION CONTACT: Ms. Delores Grant, Division of Toxicology and Human Health Sciences, Agency for Toxic Substances and Disease Registry, 1600 Clifton Rd. NE, MS F–57, Atlanta, GA 30329. Phone: (800) 232–4636 or 770–486–3351.

SUPPLEMENTARY INFORMATION: The Superfund Amendments and Reauthorization Act (SARA) (Pub. L. 99–499) amends the Comprehensive Environmental Response, Compensation, and Liability Act (CERCLA or Superfund) (42 U.S.C. 9601 et seq.) by establishing certain responsibilities for ATSDR and the U.S. Environmental Protection Agency (U.S. EPA) regarding hazardous substances most commonly found at facilities on the CERCLA National Priorities List (NPL). As part of these responsibilities, the ATSDR Administrator must prepare Toxicological Profiles for substances enumerated on the priority list of hazardous substances. This list identifies 275 hazardous substances which, according to ATSDR and U.S. EPA, pose the most significant potential threat to human health. The availability of the revised priority list of 275 hazardous substances was announced in the Federal Register on May 28, 2014 (79 FR 30613). In addition, ATSDR has the authority to prepare Toxicological Profiles for substances not found at sites on the National Priorities List, in an effort to “... establish and maintain inventory of literature, research, and studies on the health effects of toxic substances” under CERCLA Section 104(i)(1)(B). ATSDR also prepares Toxicological Profiles in response to requests for consultation under section 104(i)(4), and as otherwise necessary to support the site-specific response actions conducted by ATSDR.

Each profile will include an examination, a summary, and an interpretation of available toxicological information and epidemiological evaluations. This information and these data identify the levels of significant human exposure for the substance and for the associated health effects. The profiles must also include a determination of whether adequate information on the health effects of each substance is available (or in the process of development) in order to identify levels of significant human exposure. If adequate information is not available, ATSDR, in cooperation with the National Toxicology Program (NTP), is required to ensure the initiation of a program of research to provide such information.

SET 27 TOXICOLOGICAL PROFILES

<table>
<thead>
<tr>
<th>No.</th>
<th>Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Polychlorinated Biphenyl Ethers (PCB)</td>
</tr>
<tr>
<td>2</td>
<td>N,N-Diethyl-meta-toluamide (DEET)</td>
</tr>
<tr>
<td>3</td>
<td>Toluene Diisocyanates (mixture), Methylene diisocyanate (MDI)</td>
</tr>
<tr>
<td>4</td>
<td>Nitrates/Nitrites (NEW)</td>
</tr>
<tr>
<td>5</td>
<td>Toluene (UPDATED)</td>
</tr>
</tbody>
</table>


Donna B. Knutson, Acting Director, Office of Policy, Planning and Evaluation, National Center for Environmental Health and Agency for Toxic Substances and Disease Registry.

[FR Doc. 2015–26321 Filed 10–15–15; 8:45 am]

BILLING CODE 4153–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifier: CMS–10003, CMS–10467, CMS–1450(UB–04), CMS–1500(08–05)]

Agency Information Collection Activities: Proposed Collection; Comment Request

AGENCY: Centers for Medicare & Medicaid Services, HHS.

ACTION: Notice.

SUMMARY: The Centers for Medicare & Medicaid Services (CMS) is announcing an opportunity for the public to comment on CMS’ intention to collect information from the public. Under the Paperwork Reduction Act of 1995 (the PRA), federal agencies are required to publish notice in the Federal Register concerning each proposed collection of information (including each proposed extension or reinstatement of an existing collection of information) and to allow 60 days for public comment on the proposed action. Interested persons are invited to send comments regarding our burden estimates or any other aspect of this collection of information, including any of the following subjects: (1) The necessity and utility of the proposed information collection for the proper performance of the agency’s functions; (2) the accuracy of the estimated burden; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) the use of automated collection techniques or other forms of information technology to minimize the information collection burden.

DATES: Comments must be received by December 15, 2015.

ADDRESSES: When commenting, please reference the document identifier or OMB control number. To be assured consideration, comments and recommendations must be submitted in any one of the following ways:

1. Electronically. You may send your comments electronically to http://www.regulations.gov. Follow the
instructions for “Comment or Submission” or “More Search Options” to find the information collection document(s) that are accepting comments.

2. By regular mail. You may mail written comments to the following address:

CMS, Office of Strategic Operations and Regulatory Affairs,
Division of Regulations Development,
Attention: Document Identifier/OMB Control Number ________
Room C4–26–05,
7500 Security Boulevard,
Baltimore, Maryland 21244–1850.

To obtain copies of a supporting statement and any related forms for the proposed collection(s) summarized in this notice, you may make your request using one of following:


2. Email your request, including your address, phone number, OMB number, and CMS document identifier, to Paperwork@cms.hhs.gov.

3. Call the Reports Clearance Office at (410) 786–1326.

FOR FURTHER INFORMATION CONTACT:
Reports Clearance Office at (410) 786–1326.

SUPPLEMENTARY INFORMATION:

Contents

This notice sets out a summary of the use and burden associated with the following information collections. More detailed information can be found in each collection’s supporting statement and associated materials (see ADDRESSES).

CMS–10003 Notice of Denial of Medical Coverage (or Payment)
CMS–10467 Evaluation of the Graduate Nurse Education Demonstration Program
CMS–1450(UB–04) Medicare Uniform Institutional Provider Bill and Supporting Regulations CMS–1500(08–05) Health Insurance Common Claims Form and Supporting Regulations

Under the PRA (44 U.S.C. 3501–3520), federal agencies must obtain approval from the Office of Management and Budget (OMB) for each collection of information they conduct or sponsor. The term “collection of information” is defined in 44 U.S.C. 3502(3) and 5 CFR 1320.3(c) and includes agency requests or requirements that members of the public submit reports, keep records, or provide information to a third party. Section 3506(c)(2)(A) of the PRA requires federal agencies to publish a 60-day notice in the Federal Register concerning each proposed collection of information, including each proposed extension or reinstatement of an existing collection of information, before submitting the collection to OMB for approval. To comply with this requirement, CMS is publishing this notice.

Information Collection

1. Type of Information Collection Request: Revision of a currently approved collection; Title of Information Collection: Notice of Denial of Medical Coverage (or Payment); Use: Medicare health plans, including Medicare Advantage plans, cost plans, and Health Care Prepayment Plans, are required to issue the CMS–10003 form when a request for either a medical service or payment is denied in whole or in part. The notice explains why the plan denied the service or payment and informs Medicare enrollees of their appeal rights. The notice is also used, as appropriate, to explain Medicaid appeal rights to full dual eligible individuals enrolled in a Medicare health plan that is also managing the individual’s Medicaid benefits. To that end, the revised notice contains bracketed text the plan will insert if the denial notice is being delivered to an enrollee who is a full dual eligible. The text in square brackets “[ ]” reflects the Federal protections for Medicaid managed care enrollees. Since a State may offer additional protections, there is also free-text space for inclusion of any State-specific protections that exceed the Federal protections. Form Number: CMS–10003 (OMB control number: 0938–0829).

Frequency: Occasionally; Affected Public: Private sector (Business and other for-profit and Not-for-profit institutions); Number of Respondents: 730; Total Annual Responses: 33,574,293; Total Annual Hours: 5,593,477. (For policy questions regarding this collection contact Staci Paige at 410–786–2045. For all other issues call 410–786–1326.)

2. Type of Information Collection Request: Revision of a currently approved information collection; Title of Information Collection: Evaluation of the Graduate Nurse Education Demonstration Program; Use: The Graduate Nurse Education (GNE) Demonstration is mandated under Section 5509 of the Affordable Care Act (ACA) under title XVIII of the Social Security Act (42 U.S.C. 1395 et seq.). According to Section 5509 of the ACA, the five selected demonstration sites receive Medicare’s reasonable costs for the provision of qualified clinical training to advance practice registered nurses.” Section 5509 of the ACA also states that an evaluation of the graduate nurse education demonstration must be completed no later than October 17, 2017. This evaluation includes analysis of the following: (1) Growth in the number of advanced practice registered nurses (APRNs) with respect to a specific base year as a result of the demonstration; (2) growth for each of the following specialties: clinical nurse specialist, nurse practitioner, certified nurse anesthetist, certified nurse-midwife; and (3) costs to the Medicare program as result of the demonstration.

All information collected through the Evaluation of the GNE project will be used to meet the requirements specified under the ACA Section 5509. We will also use the information to determine the overall effectiveness of the GNE project. The process evaluation seeks to understand how the demonstration is implemented overall, how that implementation has changed over time, which aspects of the demonstration have been successful or unsuccessful, and what plans the sites have for the remainder of the implementation and after the demonstration formally ends. The process evaluation will answer both quantitative and qualitative questions. Form Number: CMS–10467 (OMB control number: 0938–1212); Frequency: Annually; Affected Public: State, Local, or Tribal Governments; Private sector (Business and other for-profit and Not-for-profit institutions); Number of Respondents: 104; Total Annual Responses: 104; Total Annual Hours: 802. (For policy questions regarding this collection contact Pauline Karikari-Martin at 410–786–1040.)

3. Type of Information Collection Request: Extension of a currently approved collection; Title of Information Collection: Medicare Uniform Institutional Provider Bill and Supporting Regulations in 42 CFR 424.5; Use: Section 42 CFR 424.5(a)(5) requires providers of services to submit a claim for payment prior to any Medicare reimbursement. Charges billed are coded by revenue codes. The bill specifies diagnoses according to the International Classification of Diseases, Ninth Edition (ICD–9–CM) code. Inpatient procedures are identified by ICD–9–CM codes, and outpatient procedures are described using the CMS Common Procedure Coding System (HCPCS). These are standard systems of identification for all major health insurance claims payers. Submission of information on the CMS–1450 permits Medicare intermediate claims and outpatient claims to receive consistent data for proper payment. Form Numbers: CMS–1450 (UB–04)
4. Type of Information Collection Request: Extension of a currently approved collection; Title of Information Collection: Health Insurance Common Claims Form and Supporting Regulations at 42 CFR part 424, Subpart C; Use: The Form CMS–1500 answers the needs of many health insurers. It is the basic form prescribed by CMS for the Medicare program for claims from physicians and suppliers. The Medicaid State Agencies, CHAMPUS/TriCare, Blue Cross/Blue Shield Plans, the Federal Employees Health Benefit Plan, and several private health plans also use it; it is the de facto standard "professional" claim form.

Medicare carriers use the data collected on the CMS–1500 and the CMS–1490S to determine the proper amount of reimbursement for Part B medical and other health services (as listed in section 1861(s) of the Social Security Act) provided by physicians and suppliers to beneficiaries. The CMS–1500 is submitted by physicians/suppliers for all Part B Medicare. Serving as a common claim form, the CMS–1500 can be used by other third-party payers (commercial and nonprofit health insurers) and other Federal programs (e.g., CHAMPUS/TriCare, Railroad Retirement Board (RRB), and Medicaid). However, as the CMS–1500 displays data items required for other third-party payers in addition to Medicare, the form is considered too complex for use by beneficiaries when they file their own claims. Therefore, the CMS–1490S (Patient's Request for Medicare Payment) was explicitly developed for easy use by beneficiaries who file their own claims. The form can be obtained from any Social Security office or Medicare carrier, Form Number: CMS–1500 (08/05), CMS–1490–S (OMB control number: 0938–0999) Frequency: On occasion; Affected Public: State, Local, or Tribal Governments, Private sector (Business or other-for-profit and Not-for-profit institutions); Number of Respondents: 53,111; Total Annual Responses: 181,909,654; Total Annual Hours: 1,567,455. (For policy questions regarding this collection contact Matt Klischer at 410–786–7488.)

Dated: October 13, 2015.

William N. Parham, III,
Director, Paperwork Reduction Staff, Office of Strategic Operations and Regulatory Affairs.

[FR Doc. 2015–26390 Filed 10–15–15; 8:45 am]
BILLING CODE 4120–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Submission for OMB Review; Comment Request

Title: Native Language Preservation and Maintenance Grant Application Template Pilot (Funding Application Submission Tool (F.A.S.T. form))

OMB No.: 0938–0999

Description: The proposed F.A.S.T. form is intended to be used by applicants in the Administration for Native Americans' Native American Language Preservation and Maintenance grant competition in FY 2016. The F.A.S.T. form is proposed to be piloted as a consolidated and streamlined preformatted electronic application form that is user-friendly and has an interactive interface providing structure and clarity for applicants. The proposed F.A.S.T. form is not intended to replace the Funding Opportunity Announcement (FOAs) which will still function as the full text of all funding opportunities for which applications are sought and considered by the Administration for Native Americans.

The proposed F.A.S.T. form will be used in a pilot capacity in just one Administration for Native Americans’ discretionary program area: Native American Language Preservation and Maintenance. All applicants applying for funding in that program area will be required to use the F.A.S.T. form during the pilot competition proposed for FY 16 unless they request and receive approval to submit a paper application. By using the F.A.S.T. form no applicant will be required to provide any information beyond what is already required by the FOA. Additionally, free training and technical assistance will be available to all applicants on use of the F.A.S.T. form.

ANA intends to use the project proposals submitted via the F.A.S.T. form to make funding decisions for Native American Language Preservation and Maintenance grant awards made in the FY 2016 pilot year. In addition, ANA will solicit feedback from applicants and panel reviewers to obtain feedback on the results, outcomes, and their recommendations regarding the F.A.S.T. form as a user friendly method of applying for funding opportunities. If the pilot is successful in making it easier for applicants to apply, ANA will consider potentially expanding use of the F.A.S.T. form to all Administration for Native Americans’ discretionary funding areas in subsequent years.

Respondents: 40.

ANNUAL BURDEN ESTIMATES

<table>
<thead>
<tr>
<th>Instrument</th>
<th>Number of Respondents</th>
<th>Number of Responses per Respondent</th>
<th>Average Burden Hours per Response</th>
<th>Total Burden Hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>F.A.S.T. form</td>
<td>40</td>
<td>28</td>
<td>.50</td>
<td>14</td>
</tr>
</tbody>
</table>

Estimated Total Annual Burden Hours: 560.

Additional Information

Copies of the proposed collection may be obtained by writing to the Administration for Children and Families, Office of Planning, Research and Evaluation, 370 L’Enfant Promenade SW., Washington, DC 20447, Attn: ACF Reports Clearance Officer. All requests should be identified by the title of the information collection. Email address: infocollection@acf.hhs.gov.

OMB Comment

OMB is required to make a decision concerning the collection of information between 30 and 60 days after publication of this document in the Federal Register. Therefore, a comment is best assured of having its full effect if OMB receives it within 30 days of publication. Written comments and recommendations for the proposed information collection should be sent directly to the following: Office of Management and Budget, Paperwork Reduction Project, Email: OIRA_SUBMISSION@OMB.EOP.GOV,