C. Public Comments

Public comments are particularly invited on: Whether this collection of information is necessary for the proper performance of functions of the FAR, and whether it will have practical utility; whether our estimate of the public burden of this collection of information is accurate, and based on valid assumptions and methodology: ways to enhance the quality, utility, and clarity of the information to be collected; and ways in which we can minimize the burden of the collection of information on those who are to respond, through the use of appropriate technological collection techniques or other forms of information technology.

Obtaining Copies of Proposals: Requesters may obtain a copy of the information collection documents from the General Services Administration, Regulatory Secretariat Division (MVCB), 1800 F Street NW., Washington, DC 20405, telephone 202–501–4755. Please cite OMB Control No. 9000–0138, Contract Financing, in all correspondence.

Dated: October 5, 2016.

Lorin S. Curit,
Director, Federal Acquisition Policy Division, Office of Governmentwide Acquisition Policy, Office of Governmentwide Policy.

[FR Doc. 2016–24538 Filed 10–11–16; 8:45 am] BILLCODE 6820–0–P

DEPARTMENT OF DEFENSE

GENERAL SERVICES ADMINISTRATION

NATIONAL AERONAUTICS AND SPACE ADMINISTRATION

[OMB Control No. 9000–0096; Docket No. 2016–0053; Sequence No. 31]

Submission for OMB Review; Patents

AGENCY: Department of Defense (DOD), General Services Administration (GSA), and National Aeronautics and Space Administration (NASA).

ACTION: Notice of request for comments regarding an extension to an existing OMB clearance.

SUMMARY: Under the provisions of the Paperwork Reduction Act, the Regulatory Secretariat Division will be submitting to the Office of Management and Budget (OMB) a request to review and approve an extension of a previously approved information collection requirement concerning patents. A notice was published in the Federal Register at 81 FR 43607 on July 5, 2016. No comments were received.

DATES: Submit comments on or before November 14, 2016.

ADDRESSES: Submit comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to: Office of Information and Regulatory Affairs of OMB, Attention: Desk Officer for GSA, Room 10236, NEOB, Washington, DC 20503. Additionally submit a copy to GSA by any of the following methods:

• Regulations.gov: http://www.regulations.gov. Submit comments via the Federal eRulemaking portal by searching for “9000–0096; Patents”. Select the link “Submit a Comment” that corresponds with “Information Collection 9000–0096, Patents”. Follow the instructions provided at the “Submit a Comment” screen. Please include your name, company name (if any), and “Information Collection 9000–0096, Patents” on your attached document.

• Mail: General Services Administration, Regulatory Secretariat Division (MVCB), IC 9000–0096, 1800 F Street NW., Washington, DC 20405. Instructions: Please submit comments only and cite Information Collection 9000–0096, Patents, in all correspondence related to this collection. Comments received generally will be posted without change to http://www.regulations.gov, including any personal and/or business confidential information provided. To confirm receipt of your comment(s), please check www.regulations.gov, approximately two to three days after submission to verify posting (except allow 30 days for posting of comments submitted by mail).

FOR FURTHER INFORMATION CONTACT: Mr. Charles Gray, Procurement Analyst, Federal Acquisition Policy Division, at 703–795–6328 or email charles.gray@gsa.gov.

SUPPLEMENTARY INFORMATION:

A. Purpose

The patent coverage in Federal Acquisition Regulation (FAR) subpart 27.2 requires the contractor to report each notice of a claim of patent or copyright infringement that came to the contractor’s attention in connection with performing a Government contract (FAR 27.202–1 and 52.227–2).

The contractor is also required to report all royalties anticipated or paid in excess of $250 for the use of patented inventions by furnishing the name and address of licensor, date of license agreement, patent number, brief description of item or component, percentage or dollar rate of royalty per unit, unit price of contract item, and number of units (FAR 27.202–5, 52.227–6, and 52.227–9).

B. Annual Reporting Burden

Number of Respondents: 107.

Responses per Respondent: 1.

Total Annual Responses: 107.

Hours per Response: 1.17.

Estimated Total Burden Hours: 126.

Frequency: On Occasion.

Affected Public: Businesses or other-for-profit entities and not-for-profit institutions.

C. Public Comments

Public comments are particularly invited on: Whether this collection of information is necessary for the proper performance of functions of the FAR, and whether it will have practical utility; whether our estimate of the public burden of this collection of information is accurate, and based on valid assumptions and methodology: ways to enhance the quality, utility, and clarity of the information to be collected; and ways in which we can minimize the burden of the collection of information on those who are to respond, through the use of appropriate technological collection techniques or other forms of information technology.

Obtaining Copies of Proposals: Requesters may obtain a copy of the information collection documents from the General Services Administration, Regulatory Secretariat Division (MVCB), 1800 F Street NW., Washington, DC 20405, telephone 202–501–4755. Please cite OMB Control No. 9000–0096, Patents, in all correspondence.

Dated: October 5, 2016.

Lorin S. Curit,
Director, Federal Acquisition Policy Division, Office of Governmentwide Acquisition Policy, Office of Governmentwide Policy.

[FR Doc. 2016–24537 Filed 10–11–16; 8:45 am] BILLCODE 6820–14–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Statement of Organization, Functions, and Delegations of Authority

Part C (Centers for Disease Control and Prevention) of the Statement of Organization, Functions, and Delegations of Authority of the Department of Health and Human Services (45 FR 67772–76, dated October 14, 1980, and corrected at 45 FR 69296, October 20, 1980, as amended most recently at 81 FR 66284–66285,
dated September 27, 2016) is amended to reflect the reorganization of the Division of Scientific Resources, National Center for Emerging and Zoonotic Infectious Diseases, Office of Infectious Diseases, Centers for Disease Control and Prevention.

Section C–B, Organization and Functions, is hereby amended as follows:

Delete and replace the title and the mission and function statements for the Division of Scientific Resources (CVLH) and insert the following: Division of Scientific Resources (CVLH). The Division of Scientific Resources (DSR) provides products, services, and specialized expertise to CDC staff and activities in support of research and service activities: In carrying out its mission, DSR: (1) Provides animals, laboratory supplies, animal and human blood products, glassware, mammalian tissue cultures, microbiological media, special reagents, and other materials in support of research and service activities to laboratories and investigators at CDC; (2) develops and implements applied research programs to expand and enhance the use of animal models necessary to support research and diagnostic programs and to improve breeding and husbandry procedures; (3) conducts applied research in cell biology and in the expansion of tissue culture technology as a research and diagnostic tool for infectious disease activities; (4) provides services for laboratory investigators in protein and DNA synthesis and sequencing, genomic sequencing, microarrays, proteomics, and molecular modeling; (5) maintains a bank of serum and other biological specimens of epidemiological and special significance to CDC’s research and diagnostic activities; (6) obtains and distributes experimental and orphaned vaccines, drugs, antisera, antitoxins, and immune globulins; (7) manages and distributes the inventory, maintains the computerized system database, and provides general technical service support for the dispensing, lyophilizing, capping, and labeling of CDC reference reagents; (8) receives, triages, processes, and distributes specimens to CDC laboratories for reference diagnostic testing, research studies, and epidemics and reports diagnostic test results to submitting organizations; (9) manages all CDC exports and ensures compliance with regulations and serves as CDC liaison with the Department of Commerce for export related issues; (10) produces specialized reagents and kits for the detection of select agents to members of the Laboratory Response Network; (11) provides services and expertise in development of quality systems to support compliance with the Food and Drug Administration regulations on production, distribution, and use of laboratory diagnostic reagents; (12) provides liaison activities, resources, and expertise for inquiries related to animals and zoonotic diseases; and (13) provides a centralized activity for tracking requests for and distributing select agents to investigators outside of CDC in compliance with federal regulations.

Office of the Director (CVLH1). (1) Manages, directs, and coordinates the activities of DSR; (2) provides leadership and guidance on policy, budget, program planning and development, program management, and operations; (3) provides DSR-wide administrative and program services and coordinates or ensures coordination with the appropriate CIOs, OID, and CDC staff offices on administrative and program matters; (4) provides liaison with other governmental agencies, international organizations, and other outside groups; (5) coordinates, in collaboration with the appropriate CIOs, OID, and CDC components, laboratory activities relating to support of outbreak investigations or laboratory-based research including but not limited to specimen management, biological reagents, and laboratory supplies; (6) maintains a formulary of investigational and licensed drugs and biologicals that are distributed to approved physicians for the prevention, control, and/or treatment of rare, tropical, or exceptional diseases; (7) advises the Director, NCEZID, on policy matters concerning DSR activities; and (8) coordinates technical services for laboratory activities of CDC programs including procurement of glassware and laboratory supplies.

Comparative Medicine Branch (CVLH2). (1) Acquires and distributes laboratory animals for research; (2) provides appropriate housing, husbandry, and psychological enrichment for all research animals; (3) provides veterinary services, including clinical and surgical support, for the laboratory animals; (4) develops standard operating procedures for animal care and use in accordance with the policies established by the American Association for Accreditation of Laboratory Animal Care, the Animal Welfare Act, The Guide for the Care & Use of Laboratory Animals, and the CDC International Animal Care & Use Committee; (5) conducts applied research to improve the care and use of animals in research and collaborates on research projects that use laboratory animals; (6) provides consultation and laboratory animal technology training to investigators, technical staff and animal care personnel; (7) provides oversight, support and investigator training for the graphical animal information technology protocol development and animal tracking database; (8) coordinates technical services for laboratory activities of CDC programs including processing and distribution of glassware and related items, laboratory waste decontamination and disposal, laundry services, and materiel management; and (9) provides autoclave label production services.

Biotechnology Core Facility Branch (CVLH3). (1) provides state-of-the-art next-generation genomic sequencing and metagenomics analysis of infectious and biothreat agents; (2) provides optical mapping to produce high resolution whole-genome maps for strain typing, molecular epidemiology, comparative genomics, and quality control for whole genome sequence assembly; (3) provides computational analysis of genomics sequencing data, bioinformatics, and biological computing; (4) provides qualitative and quantitative proteomic analyses (identification of expressed proteins by mass spectrometry); analysis of functionally-relevant post-translational modifications of proteins; (5) provides mass spectrometry-based positive identification of bacteria and fungi (BiotyperTM, Bruker Daltonics); (6) provides synthetic oligonucleotide chemistry in support of development of rapid diagnostic tests and characterization of pathogens and their hosts; (7) provides synthetic peptide chemistry in support of studies of immune response and antigen-antibody interactions; (8) provides biotechnology seminars and methods evaluation; (9) provides laboratory equipment design and repair services to all CDC; and (10) collaborates on research related to STD transmitted infections as chronic infectious diseases.

Reagent and Diagnostic Services Branch (CVLH4). (1) maintains laboratory water treatment systems to ensure quality of CDC reagent grade laboratory water; (2) produces, develops, evaluates and distributes custom microbiological and cell culture media, buffers and chemical reagent, mammalian and insect cell cultures, hybridomas, monoclonal and polyclonal antibodies, and in vitro diagnostic products for diagnostic research purposes, proficiency testing, pandemic preparedness, outbreak response and surveillance activities; (3) collaborates with subject matter experts.
in regulatory compliant development, production, packaging, storing and distribution of BSL2/BSL3 reagents, select agents, novel immuno-chemical reagents and reference diagnostic reagents; (4) provides dispensing, lyophilizing, label production, and device assembly services; (5) improves the process of bench-top development and in-house pilot scale production providing immediate availability for distribution, preventing backorders and streamlining commercialization; (6) maintains CDC’s Biological Reference Reagent Inventory, mammalian cell line repository and a serviceable inventory at the DSR Continuity of Operations storage facility; (7) provides centralized specimen management services for diagnostic, reference, and outbreak investigations; maintains a bank of biological specimens of epidemiological significance to CDC’s research and diagnostic activities; manages and tracks systems of specimen collections; (8) receives, triages, processes, stores and distributes specimens to CDC laboratories for reference diagnostic testing, research studies, and reports diagnostic and surveillance test results to submitting organizations; (9) serves as central facility for acquisition and distribution of fresh human blood, blood products, and serum in bulk; (10) packages and ships infectious substances and other materials, ensuring compliance with regulations for shipping clinical specimens, infectious substances, and other materials; (11) manages all CDC exports and ensures compliance with regulations and serves as CDC liaison with Department of Commerce for export related issues; and (12) provides consultation in all of the above technical services.

Sherri Berger,
Chief Operating Officer, Centers for Disease Control and Prevention.

[FR Doc. 2016–24640 Filed 10–11–16; 8:45 am]

BILLING CODE 4160–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Agency Recordkeeping/Reporting Requirements Under Emergency Review by the Office of Management and Budget (OMB)

Title: Provision of Child Support Services in IV-D cases under the Hague Child Support Convention; Federally Approved Forms.

OMB No.: New Collection.

Description: On January 1, 2017, the 2007 Hague Convention on the International Recovery of Child Support and Other Forms of Family Maintenance will enter into force for the United States. This Convention contains groundbreaking provisions that, for the first time on a worldwide scale, will establish uniform, simple, fast, and inexpensive procedures for the processing of international child support cases. Once the Convention is in effect, U.S. states will process child support cases with other countries that have ratified the Convention under the requirements of the Convention and article 7 of the Uniform Interstate Family Support Act (UIFSA 2008). In order to comply with the Convention, the U.S. must implement the Convention’s case processing forms.

State and Federal law require states to use federally-approved case processing forms. Section 311(b) of UIFSA 2008, which has been enacted by all 50 states, the District of Columbia, Guam, Puerto Rico and the Virgin Islands, requires States to use forms mandated by Federal law. 45 CFR 303.7 also requires child support programs to use federally-approved forms in intergovernmental IV-D cases unless a country has provided alternative forms as a part of its chapter in a Caseworker’s Guide to Processing Cases with Foreign Reciprocating Countries.

Respondents: State agencies administering a child support program under title IV-D of the Social Security Act.

<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>
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<tr>
<td>Annex I: Transmittal form under Article 12(2)</td>
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<td>Annex II: Acknowledgment form under Article 12(3)</td>
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<td>Annex A: Abstract of Decision</td>
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<td>Annex A: Statement of Enforceability of Decision</td>
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<td>19</td>
<td>.5</td>
<td>135</td>
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<tr>
<td>Annex A: Statement of Proper Notice</td>
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<td>659</td>
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<td>Annex A: Status of Application Report</td>
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<td>.33</td>
<td>659</td>
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<td>Annex B: Application for Enforcement of a Decision Made or Recognized in the Requested State, including restricted information on the applicant</td>
<td>54</td>
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<td>Annex B: Status of Application Report, Article 12</td>
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<td>Annex D: Application for Modification of a Decision, including Restricted Information on the Applicant</td>
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<td>Annex E: Financial Circumstances Form</td>
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Estimated Total Annual Burden Hours: 13,478.

Additional Information: ACF is requesting that OMB grant a 180 day approval for this information collection under procedures for emergency processing by October 20, 2016. A copy of this information collection, with applicable supporting documentation, may be obtained by calling the Administration for Children and Families, Reports Clearance Officer, Robert Sargis at (202) 690–7275.

Comments and questions about the information collection described above should be directed to the Office of Information and Regulatory Affairs, Attn: OMB Desk Officer for ACF, Office of Management and Budget, Paperwork Reduction Project, 725 17th Street NW.,