personal information; obtain the prior consent of the child's parent in order to engage in such collection, use, and disclosure, with limited exceptions; provide reasonable means for the parent to obtain access to the information and to direct its deletion; and, establish procedures that protect the confidentiality, security, and integrity of personal information collected from children.

Burden Statement¹

- 1. Estimated annual hours burden: 17,500 hours
 - (a) New entrant web operators' disclosure burden: 16,800 hours
 - (b) Safe harbor applicant reporting requirements: 100 hours, rounded (for an estimated one additional safe harbor applicant)
 - (c) Annual audit and report for safe harbor programs: 800 hours
 - (d) Safe harbor program recordkeeping requirements: 0 or minimal
- 2. Estimated annual labor costs: \$5,342,500
 - (a) New entrant web operators' disclosure burden: \$5,297,600
 - (b) Safe harbor applicant reporting requirements: \$18,500
 - (c) Annual audit and report for safe harbor programs: \$26,400
 - (d) Safe harbor program recordkeeping requirements: \$0 or marginal
- 3. Estimated annual non-labor costs: \$0

Request for Comments

You can file a comment online or on paper. For the Commission to consider your comment, we must receive it on or before January 8, 2016. Write "COPPA Rule: Paperwork Comment, FTC File No. 155408" on your comment. Your comment—including your name and your state—will be placed on the public record of this proceeding, including to the extent practicable, on the public Commission Web site, at http://www.ftc. gov/os/publiccomments.shtm. As a matter of discretion, the Commission tries to remove individuals' home contact information from comments before placing them on the Commission Web site.

Because your comment will be made public, you are solely responsible for making sure that your comment does not include any sensitive personal information, like anyone's Social Security number, date of birth, driver's

license number or other state identification number or foreign country equivalent, passport number, financial account number, or credit or debit card number. You are also solely responsible for making sure that your comment doesn't include any sensitive health information, like medical records or other individually identifiable health information. In addition, don't include any "[t]rade secret or any commercial or financial information . . . which is privileged or confidential'' as provided in Section 6(f) of the FTC Act 15 U.S.C. 46(f), and FTC Rule 4.10(a)(2), 16 CFR 4.10(a)(2). In particular, don't include competitively sensitive information such as costs, sales statistics, inventories, formulas, patterns devices, manufacturing processes, or customer names.

If you want the Commission to give your comment confidential treatment, you must file it in paper form, with a request for confidential treatment, and you have to follow the procedure explained in FTC Rule 4.9(c)).² Your comment will be kept confidential only if the FTC General Counsel grants your request in accordance with the law and the public interest.

Postal mail addressed to the Commission is subject to delay due to heightened security screening. As a result, we encourage you to submit your comments online. To make sure that the Commission considers your online comment, you must file it at *https:// ftcpublic.commentworks.com/ftc/ coppapra2*, by following the instructions on the web-based form. When this Notice appears at *http:// www.regulations.gov/#!home*, you also may file a comment through that Web site.

If you file your comment on paper, write "COPPA Rule: Paperwork Comment, FTC File No. 155408" on your comment and on the envelope, and mail it to the following address: Federal Trade Commission, Office of the Secretary, 600 Pennsylvania Avenue NW., Suite CC-5610 (Annex J), Washington, DC 20580, or deliver your comment to the following address: Federal Trade Commission, Office of the Secretary, Constitution Center, 400 7th Street SW., 5th Floor, Suite 5610 (Annex J), Washington, DC 20024. If possible, submit your paper comment to the Commission by courier or overnight service.

Comments on the information collection requirements subject to review under the PRA should additionally be submitted to OMB. If sent by U.S. mail, they should be addressed to Office of Information and Regulatory Affairs, Office of Management and Budget, Attention: Desk Officer for the Federal Trade Commission, New Executive Office Building, Docket Library, Room 10102, 725 17th Street NW., Washington, DC 20503. Comments sent to OMB by U.S. postal mail, however, are subject to delays due to heightened security precautions. Thus, comments instead should be sent by facsimile to (202) 395-5806.

The FTC Act and other laws that the Commission administers permit the collection of public comments to consider and use in this proceeding as appropriate. The Commission will consider all timely and responsive public comments that it receives on or before January 8, 2016. For information on the Commission's privacy policy, including routine uses permitted by the Privacy Act, see http://www.ftc.gov/ftc/ privacy.htm.

David C. Shonka,

Principal Deputy General Counsel. [FR Doc. 2015–30935 Filed 12–8–15; 8:45 am] BILLING CODE 6750–01–P

DEPARTMENT OF DEFENSE

GENERAL SERVICES ADMINISTRATION

NATIONAL AERONAUTICS AND SPACE ADMINISTRATION

[OMB Control No. 9000-0070; Docket 2015-0055; Sequence 26]

Information Collection; Payments

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

ACTION: Notice of request for comments regarding the extension of a previously 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 payments.

DATES: Submit comments on or before February 8, 2016.

¹ This discussion and the associated burden estimates concern strictly recurring compliance obligations under the COPPA Rule. Details underlying the estimates within this Burden Statement can be found in the September 25, 2015 **Federal Register** Notice.

² In particular, the written request for confidential treatment that accompanies the comment must include the factual and legal basis for the request, and must identify the specific portions of the comment to be withheld from the public record. *See* FTC Rule 4.9(c), CFR 4.9(c), 16 CFR 4.9(c).

ADDRESSES: Submit comments identified by Information Collection 9000–0070, Payments, by any of the following methods:

• Regulations.gov: http:// www.regulations.gov. Submit comments via the Federal eRulemaking portal by searching the OMB control number. Select the link "Submit a Comment" that corresponds with "Information Collection 9000–0070, Payments". Follow the instructions provided at the "Submit a Comment" screen. Please include your name, company name (if any), and "Information Collection 9000– 0070, Payments" on your attached document.

• *Mail:* General Services Administration, Regulatory Secretariat Division (MVCB), 1800 F Street NW., Washington, DC 20405. ATTN: Ms. Flowers/IC 9000-0070, Payments.

Instructions: Please submit comments only and cite Information Collection 9000-0070, Payments, 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:

Kathy Hopkins, Procurement Analyst, Office of Acquisition Policy, GSA at 202-969-7226 or email at *kathlyn.hopkins@gsa.gov*.

SUPPLEMENTARY INFORMATION:

A. Purpose

Firms performing under Federal contracts must provide adequate documentation to support requests for payment under these contracts. The documentation may range from a simple invoice to detailed cost data. The information is usually submitted once, at the end of the contract period or upon delivery of the supplies, but could be submitted more often depending on the payment schedule established under the contract (see FAR 52.232-1 through 52.232-4, and FAR 52.232-6 through 52.232–11). The information is used to determine the proper amount of payments to Federal contractors.

B. Annual Reporting Burden

Respondents: 80,000. Responses per Respondent: 120. Total Responses: 9,600,000. Hours per Response: .25. Total Burden Hours: 2,400,000.

C. Public Comments

Public comments are particularly invited on: Whether this collection of information is necessary for the proper performance of functions of the Federal Acquisition Regulations (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.9000–0070, Payments, in all correspondence.

Edward Loeb,

Acting Director, Federal Acquisition Policy Division, Office of Governmentwide Acquisition Policy, Office of Acquisition Policy, Office of Governmentwide Policy. [FR Doc. 2015–30969 Filed 12–8–15; 8:45 am] BILLING CODE 6820–EP–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 80 FR 73766–73769, dated November 25, 2015) is amended to reflect the reorganization of the Division of Global HIV/AIDS, Center for Global Health, Centers for Disease Control and Prevention.

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

Delete in its entirety the title and the mission and function statements for the *Division of Global HIV/AIDS (CWG)* and insert the following:

Division of Global HIV and TB (CWG). The Division of Global HIV and TB (DGHT) provides technical assistance to host governments, working through its strong partnerships with Ministries of Health and local and international partners to implement integrated HIV/ AIDS clinical and preventive services and systems; develop and strengthen laboratory services; and provide epidemiologic science, informatics, and research support to develop sustainable public health systems in resourceconstrained countries. DGHT: (1) Provides leadership, management, and services to DGHT country offices; (2) implements integrated evidence-based prevention, care, and treatment programs and services; (3) evaluates program costs, cost effectiveness and impact to assist with prioritization, inform program planning, and appropriate rates of program expansion, and strengthens capacity for sustainable, high quality research and service implementation to indigenous partners and Ministries of Health; (4) builds sustainable public health capacity in laboratory services and systems; (5) ensures epidemiologic and scientific excellence in HIV/AIDS programs; (6) contributes to the broader scientific body of knowledge in global public health by systematically evaluating the scope and quality of global HIV/AIDS and TB programs; (7) implements operations and effectiveness research to inform the design of current and future programs as well as optimize allocation of human and financial resources; (8) strengthens in-country capacity to design and implement HIV/AIDS surveillance systems and surveys; (9) builds host government public health management capacity and trains incountry public heath workforce with the goal of long-term program sustainability; (10) supports host government capacity to monitor and evaluate the process, outcome, and impact of HIV prevention, care, and treatment programs; and (11) helps countries respond to public health emergencies, assisting in response planning and implementation with Ministries of Health and other international partners.

Office of the Director (CWG1). (1) Provides strategic leadership, guidance, management and oversight to all DGHT programs and ensures coordination and communication across its branches and with other CDC programs including CDC/Washington; U.S. Government (USG) agencies, including the Department of Health and Human Services (HHS), the United States Agency for International Development (USAID), and Department of State (DoS);