

Respondent's research is currently being supervised by KU; Respondent shall ensure that a plan for supervision of his PHS-related duties is submitted to ORI for approval either within two weeks of this Agreement becoming final or prior to receiving or applying for PHS funds if such support is not current at the time this Agreement is completed; the supervision plan must be designed to ensure the scientific integrity of his research contribution; because of the ongoing review of Respondent's research by KU, ORI will only require a summary report on the first and second anniversary of the Agreement detailing how KU has ensured that Respondent's research and language in PHS grant applications and reports of PHS-supported research have been verified to be his own and accurately reported; Respondent agrees to maintain responsibility for compliance with the agreed upon supervision plan;

(2) that this annual summary, provided by any institution employing him, shall provide assurance that each application for PHS funds, or report, manuscript, or abstract involving PHS-supported research in which Respondent was involved, was based on actual experiments or was otherwise legitimately derived, that the data, procedures, and methodology were accurately reported in the application, report, manuscript, or abstract, and that the text in such submissions was his own or properly cited the source of copied language and ideas; and

(3) to exclude himself from serving in any advisory capacity to PHS including, but not limited to, service on any PHS advisory committee, board, and/or peer review committee, or as a consultant.

FOR FURTHER INFORMATION CONTACT: Director, Division of Investigative Oversight, Office of Research Integrity, 1101 Wootton Parkway, Suite 750, Rockville, MD 20852, (240) 453-8800.

John Dahlberg,

Director, Division of Investigative Oversight, Office of Research Integrity.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifier: CMS-10368]

Agency Information Collection Activities: Submission for OMB Review; Comment Request

AGENCY: Centers for Medicare & Medicaid Services, HHS.

In compliance with the requirement of section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Centers for Medicare & Medicaid Services (CMS), Department of Health and Human Services, is publishing the following summary of proposed collections for public comment. Interested persons are invited to send comments regarding this burden estimate 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 function; (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.

1. *Type of Information Collection Request:* New collection (request for a new OMB control number); *Title of Information Collection:* Dental Action Plan Template for Medicaid and CHIP Programs; *Use:* CMS is responsible for administering the Federal Medicaid program and the Children's Health Insurance Program (CHIP). As part of the Federal Medicaid program, CMS oversees the Early and Periodic Screening, Diagnostic, and Treatment (EPSDT) benefit to assure that all requirements are met. The provision of dental services to EPSDT-eligible individuals is required under section 1905(r)(3) of the Social Security Act. In addition, section 1902(a)(43)(D)(iii) requires that CMS collect information on dental services furnished to eligible individuals. Section 501(e) of CHIPRA imposed new data reporting requirements for the CHIP program by requiring certain dental data to be reported in 2011 on the CHIP annual report. Dental data for CHIP is unavailable as the requirement to report this data is new for CHIP programs. CMS intends to use the information provided in the template to help inform us of the States' activities undertaken to achieve the national oral health goals for

Medicaid and CHIP. CMS will use the information to routinely follow-up with States on the achievement of their goals and activities and will share that information with other States. The template has been revised since the publication of the 60-day notice by clarifying instructions and by making minor changes. The supporting documents have not been changed; *Form No.:* CMS-10368 (OCN 0938-NEW); *Frequency:* Once; *Affected Public:* State, Local, or Tribal Governments; *Number of Respondents:* 69; *Total Annual Responses:* 69; *Total Annual Hours:* 4,485. (For policy questions regarding this collection contact Cindy Ruff at (410) 786-5916. For all other issues call (410) 786-1326.)

To obtain copies of the supporting statement and any related forms for the proposed paperwork collections referenced above, access CMS Web site address at <http://www.cms.hhs.gov/PaperworkReductionActof1995>, or email your request, including your address, phone number, OMB number, and CMS document identifier, to Paperwork@cms.hhs.gov, or call the Reports Clearance Office on (410) 786-1326.

To be assured consideration, comments and recommendations for the proposed information collections must be received by the OMB desk officer at the address below, no later than 5 p.m. on *January 23, 2012*. OMB, Office of Information and Regulatory Affairs, Attention: CMS Desk Officer, Fax Number: (202) 395-6974, email: OIRA_submission@omb.eop.gov.

Dated: December 16, 2011.

Martique Jones,

Director, Regulations Development Group, Division-B, Office of Strategic Operations and Regulatory Affairs.

[FR Doc. 2011-33098 Filed 12-22-11; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifier: CMS-1880 and CMS-1882]

Agency Information Collection Activities: Proposed Collection; Comment Request

AGENCY: Centers for Medicare & Medicaid Services, HHS.

In compliance with the requirement of section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Centers for Medicare & Medicaid

Services (CMS) is publishing the following summary of proposed collections for public comment. Interested persons are invited to send comments regarding this burden estimate 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.

1. *Type of Information Collection Request*: Extension without change of a currently approved collection; *Title of Information Collection*: Certification as a Supplier of Portable X-Ray and Portable X-Ray Survey Report Form and Supporting Regulations at 42 CFR Part 486.100–486.110; *Use*: CMS–1880 is utilized as an application to be completed by suppliers of portable X-ray services requesting participation in the Medicare program. This form initiates the process of obtaining a decision as to whether the conditions of coverage are met as a portable X-ray supplier. It also promotes data reduction or introduction to, and retrieval from, the Certification and Survey Provider Enhanced Reporting (CASPER) by the CMS Regional Offices (ROs).

CMS–1882 is used by the State survey agency to provide data collected during an on-site survey of a supplier of portable X-ray services to determine compliance with the applicable conditions of participation and to report this information to the Federal Government. The form is primarily a coding worksheet designed to facilitate data reduction and retrieval into the ASPEN system at the CMS ROs. The form includes basic information on compliance (i.e., met, not met, explanatory statements) and does not require any descriptive information regarding the survey activity itself. CMS has the responsibility and authority for certification decisions which are based on supplier compliance with the applicable conditions of participation. The information needed to make these decisions is available to CMS only through the use of information abstracted from the survey report form; *Form Numbers*: CMS–1880 (Request for Certification as a Supplier of Portable X-ray Services), CMS–1882 (Medicare/Medicaid Portable X-ray Survey Report), and OCN 0938–0027; *Frequency*: Occasionally; *Affected Public*: State,

Local, or Tribal Governments; *Number of Respondents*: 579; *Total Annual Responses*: 86; *Total Annual Hours*: 151. (For policy questions regarding this collection contact Georgia Johnson at (410) 786–6859. For all other issues call (410) 786–1326.)

To obtain copies of the supporting statement and any related forms for the proposed paperwork collections referenced above, access CMS' Web site address at <http://www.cms.hhs.gov/PaperworkReductionActof1995>, or email your request, including your address, phone number, OMB number, and CMS document identifier, to Paperwork@cms.hhs.gov, or call the Reports Clearance Office on (410) 786–1326.

In commenting on the proposed information collections please reference the document identifier or OMB control number. To be assured consideration, comments and recommendations must be submitted in one of the following ways by February 21, 2012:

1. *Electronically*. You may submit 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) 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.

Dated: December 16, 2011.

Martique Jones,

Director, Regulations Development Group, Division B, Office of Strategic Operations and Regulatory Affairs.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Indian Health Service

Notice of a Project Waiver of Section 1605 (Buy American Requirement) of the American Recovery and Reinvestment Act of 2009 (ARRA)

AGENCY: Indian Health Service (IHS), HHS.

ACTION: Notice.

SUMMARY: The IHS is hereby granting waivers of the Buy American requirements of ARRA Section 1605 under the authority of Section

1605(b)(1) [applying the Buy American provision would be inconsistent with the public interest] to the Alaska Native Tribal Health Consortium (ANTHC) for the specific Alaska projects listed in this notice for the purchase of a foreign manufactured equipment to be installed on those sanitation facilities construction projects. This is a project specific waiver and only applies to the use of the specified product for the ARRA projects listed in this notice. Any other ARRA recipient that wishes to use the same product must apply for a separate waiver based on project specific circumstances. Based upon information submitted by the ANTHC professional engineering staff, it has been determined that applying the Buy American provision would be inconsistent with the public interest. The IHS is making this determination based on the review and recommendation of the IHS Alaska Area Office. This action permits the purchase of a foreign manufactured item for the projects specified in this notice.

Waivers are granted for the ARRA funded ANTHC projects in these Alaska communities: City of Angoon for water treatment plant media filters, City of Buckland for Flygt submersible wastewater pumps and appurtenances, City of Chignik for water treatment plant measuring equipment, City of Chuathbaluk for Grundfos pumps for its water and sewer systems, City of Deering for Grundfos pumps for its infiltration gallery and filtration plant, and City of Hooper Bay for Toyotomi Fuel Lift pumps for its existing Toyotomi fuel oil fired hotwater heaters.

DATES: *Effective Date*: Upon publication.

SUPPLEMENTARY INFORMATION: In accordance with ARRA Section 1605(c) and Section 176.80 of the rules of the Office of Management and Budget (OMB) (2 CFR 176.80), the IHS hereby provides notice that it is granting a limited waiver of the requirements of section 1605(a) of Public Law 111–5, Buy American requirements, based on the public interest authority of section 1605(b)(1), to allow the use of non-domestic iron, steel, and manufactured goods in eligible sanitation facilities construction projects.

ARRA 1605(a) prohibits the use of Recovery Act funds for the construction, alteration, maintenance, or repair of a public building or public work unless all of the iron, steel, and manufactured goods used in the project are produced in the United States, or unless a waiver is granted by the head of the Federal department or agency. ARRA 1605(b) provides that the Buy American requirement shall not apply in any case