

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Submission for OMB Review; Comment Request

Title: Work Participation and TANF/WIA Coordination Project.

OMB No.: New collection.

Description: The Administration for Children and Families (ACF) is

proposing an information collection activity as part of the Work Participation and TANF/WIA Coordination Project. The proposed information collection consists of semi-structured interviews with key state/and or local Temporary Assistance for Needy Families (TANF) and Work Investment Act (WIA) respondents on questions of engagement in additional work activities and expenditures of other benefits and services as well as questions concerning TANF/WIA Coordination. Through this

information collection, ACF seeks to elucidate the data presented in reports submitted by states to the ACF Office of Family Assistance (OFA) as required by the Claims Resolution Act of 2010. This collection is separate from the state reports to OFA required by the Act. In addition, it will provide documentation of positive TANF/WIA coordination activities.

Respondents: State and/or local administrators responsible for the TANF and WIA Programs.

ANNUAL BURDEN ESTIMATES

Instrument	Annual number of respondents	Number of responses per respondent	Average burden hours per response	Total annual burden hours
Discussion Guide for Use with State TANF officials	40	2	8	640

Estimated Total Annual Burden Hours: 640.

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:* OPRE Reports Clearance Officer. All requests should be identified by the title of the information collection. E-mail address: OPREinfocollection@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 Fax: 202-395-6974 *Attn:* Desk Officer for the Administration for Children and Families.

Dated: June 7, 2011.

Steven M. Hanmer,

OPRE Reports Clearance Officer.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2011-D-0378]

Draft Guidance for Industry and Food and Drug Administration Staff; Establishing the Performance Characteristics of In Vitro Diagnostic Devices for the Detection of Methicillin-Resistant *Staphylococcus Aureus* for Culture-Based Devices; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the availability of the draft guidance entitled "Establishing the Performance Characteristics of In Vitro Diagnostic Devices for the Detection of Methicillin-Resistant *Staphylococcus Aureus* for Culture-Based Devices." This draft guidance document provides industry and Agency staff with recommendations for studies for establishing the performance characteristics of in vitro diagnostic devices for the detection of methicillin-resistant *S. aureus* (MRSA), including those for the detection or detection and differentiation of MRSA versus *S. aureus* (SA) in either human specimens or bacterial growth detected by continuous monitoring blood culture systems. This draft guidance is not final nor is it in effect at this time.

DATES: Although you can comment on any guidance at any time (see 21 CFR 10.115(g)(5)), to ensure that the Agency considers your comment on this draft guidance before it begins work on the final version of the guidance, submit

either electronic or written comments on the draft guidance by September 13, 2011.

ADDRESSES: Submit written requests for single copies of the draft guidance document entitled "Establishing the Performance Characteristics of In Vitro Diagnostic Devices for the Detection of Methicillin-Resistant *Staphylococcus Aureus* (MRSA) for Culture-Based Devices" to the Division of Small Manufacturers, International, and Consumer Assistance, Center for Devices and Radiological Health, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 66, rm. 4613, Silver Spring, MD 20993-0002. Send one self-addressed adhesive label to assist that office in processing your request, or fax your request to 301-847-8149. See the **SUPPLEMENTARY INFORMATION** section for information on electronic access to the guidance.

Submit electronic comments on the draft guidance to <http://www.regulations.gov>. Submit written comments to the Division of Dockets Management (HFA-305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. Identify comments with the docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT: Alexandra Wong, Center for Devices and Radiological Health, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 66, rm. 5502, Silver Spring, MD 20993-0002, 301-796-6210.

SUPPLEMENTARY INFORMATION:

I. Background

FDA is issuing this draft guidance to provide industry and Agency staff with recommendations for studies for establishing the performance