One important and effective means to promote and support the initiation and maintenance of breastfeeding is through the health care system. While the few studies on breastfeeding practices at intra-partum care facilities in individual states and facilities show significant variation in practices, it is not currently possible to assess and monitor breastfeeding-related practices and policies in hospitals and free-standing childbirth centers across the United States with data currently available.

CDC plans to conduct an assessment of breastfeeding-related maternity care practices in intra-partum care facilities in the United States and Territories to provide information to individual facilities, state health departments, and CDC on the extent to which facilities are providing effective breastfeeding-related maternity care. The assessment will provide detailed information on general facility characteristics related to maternity care such as facility policies related to breastfeeding-related maternity care practices, practices

related to the training of health care staff on breastfeeding instruction, management and support, rooming-in, infant supplementation, and discharge from facility. CDC will provide facilityspecific information based on the assessment to the individual facilities and state-specific information to state health departments. The information from the survey can be used by facilities to evaluate and modify breastfeedingrelated maternity care practices, and by states and CDC to inform and target programs and policies to improve breastfeeding-related maternity care practices at intra-partum care facilities.

Approximately 4,375 facilities providing maternity care in the United States and Territories will be mailed a survey every other year in this study. The survey will be administered for the first time in 2007 and for the second time in 2009. Survey content will be similar in each of the administrations to examine changes in practices and policies over time. It is expected that approximately 3,700 facilities will

complete the thirty-minute questionnaire in each administration. The facilities will be identified from the American Hospital Association's Annual Survey of Hospitals (AHA) and the National Association of Childbearing Centers (NACC). A fiveminute screening telephone call will be made prior to survey administrations to all facilities identified as providing maternity care in AHA and NACC to ensure they are currently providing maternity care, to identify possible satellite clinics providing maternity care, and to identify survey respondent in each of the facilities. The respondents will have the option of either responding by mail or through a Webbased system. The survey will provide detailed information about breastfeeding-related maternity care practices and policies at hospitals and free-standing birth centers. There are no costs to respondents other than their time. The approximate annualized burden hours are 1,484 hours.

ESTIMATED ANNUALIZED BURDEN HOURS

Questionnaire/respondents	Number of respondents	Number of responses/ respondent	Average burden per response (in hours)
Screening call to facilities that have at least one registered maternity bed (2006)	1240	1 1 1 1	5/60 30/60 5/60 30/60

Dated: January 31, 2007.

Joan F. Karr,

Acting Reports Clearance Officer, Centers for Disease Control and Prevention.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30 Day-07-06BI]

Agency Forms Undergoing Paperwork Reduction Act Review

The Centers for Disease Control and Prevention (CDC) publishes a list of information collection requests under review by the Office of Management and Budget (OMB) in compliance with the Paperwork Reduction Act (44 U.S.C. Chapter 35). To request a copy of these requests, call the CDC Reports Clearance Officer at (404) 639–4766 or send an email to omb@cdc.gov. Send written comments to CDC Desk Officer, Office of

Management and Budget, Washington, DC or by fax to (202) 395–6974. Written comments should be received within 30 days of this notice.

Proposed Project

Determining Stakeholder Awareness and the Use and Impact of Products Developed by the Evaluation of Genomic Applications in Practice and Prevention (EGAPP) Model Project—New—National Center for Chronic Disease Prevention and Health Promotion/National Office of Public Health Genomics (NOPHG), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

More than 1,000 genetic tests are currently available in clinical practice. Most are used for diagnosis of rare genetic diseases, but a growing number have population-based applications, and the potential for broad public health impact.

Å number of issues have been raised about the current status of genetic testing implementation, including the need to develop evidence to establish

validity and utility of genetic tests before tests are commercialized. Advisory panels, professional organizations, and clinical experts have produced recommendations on the development and clinical implementation of safe and effective genetic tests. In response to the need for a coordinated approach for effectively integrating genomic tests into clinical practice and health policy, CDC's National Office of Public Health Genomics (NOPHG) initiated the (Evaluation of Genomic Applications in Practice and Prevention) EGAPP model project in 2004 to establish a systematic, evidence-based process for assessing genetic tests in transition from research to practice. To support this goal, an independent, non-federal, multidisciplinary EGAPP Working Group was established to identify, prioritize, and select genetic tests to be reviewed; establish review methods and processes; monitor progress of the reviews; and develop conclusions and recommendations based on the evidence.