

ESTIMATED ANNUALIZED BURDEN HOURS—Continued

Types of respondent	Number of respondents	Number of responses per respondent	Average burden per response (in hours)
Gonorrhea Case	2,880	1	8/60

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day–12–0493]

Proposed Data Collections Submitted for Public Comment and Recommendations

In compliance with the requirement of Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995 for opportunity for public comment on proposed data collection projects, the Centers for Disease Control and Prevention (CDC) will publish periodic summaries of proposed projects. To request more information on the proposed projects or to obtain a copy of the data collection plans and instruments, call 404–639–7570 and send comments to Kimberly Lane, CDC Reports Clearance Officer, 1600 Clifton Road, MS–D74, Atlanta, GA 30333 or send an email to omb@cdc.gov.

Comments are invited on: (a) Whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have

practical utility; (b) the accuracy of the agency's estimate of the burden of the proposed collection of information; (c) ways to enhance the quality, utility, and clarity of the information to be collected; and (d) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques or other forms of information technology. Written comments should be received within 60 days of this notice.

Proposed Project

2013 and 2015 National Youth Risk Behavior Surveys (YRBS)(OMB No. 0920–0493)—Reinstatement with change—National Center for HIV/AIDS, Viral Hepatitis, STD, and TB Prevention (NCHHSTP), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

The purpose of this request is to obtain OMB approval to reinstate with change, the data collection for the National Youth Risk Behavior Survey (YRBS), a school-based survey that has been conducted biennially since 1991. OMB approval for the 2009 YRBS and 2011 YRBS expired November 30, 2011 (OMB no. 0920–0493). CDC seeks a three-year approval to conduct the YRBS in Spring 2013 and Spring 2015. Minor changes incorporated into this reinstatement request include: an updated title for the information collection to accurately reflect the years in which the survey will be conducted, minor changes to the data collection

instrument, and a minor increase in the burden estimate.

The YRBS assesses priority health risk behaviors related to the major preventable causes of mortality, morbidity, and social problems among both youth and young adults in the United States. Data on health risk behaviors of adolescents are the focus of approximately 65 national health objectives in Healthy People 2020, an initiative of the U.S. Department of Health and Human Services (HHS). The YRBS provides data to measure 20 of the health objectives and 1 of the Leading Health Indicators established by Healthy People 2020. In addition, the YRBS can identify racial and ethnic disparities in health risk behaviors. No other national source of data measures as many of the Healthy People 2020 objectives addressing adolescent health risk behaviors as the YRBS. The data also will have significant implications for policy and program development for school health programs nationwide.

In Spring 2013 and Spring 2015, the YRBS will be conducted among nationally representative samples of students attending public and private schools in grades 9–12. Information supporting the YRBS also will be collected from state, district, and school-level administrators and teachers. The table below reports the number of respondents annualized over the 3-year project period.

There are no costs to respondents except their time. The total estimated annualized burden hours are 7,822.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondent	Form name	Number of respondents	Number of responses per respondent	Average burden per response (in hours)
State Administrators	State-level Recruitment Script for the Youth Risk Behavior Survey.	17	1	30/60
District Administrators	District-level Recruitment Script for the Youth Risk Behavior Survey.	80	1	30/60
School Administrators	School-level Recruitment Script for the Youth Risk Behavior Survey.	133	1	30/60
Teachers	Data Collection Checklist for the Youth Risk Behavior Survey.	435	1	15/60
Students	Youth Risk Behavior Survey	10,129	1	45/60

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day-12-12PE]

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Comments are invited on: (a) Whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have practical utility; (b) the accuracy of the agency's estimate of the burden of the proposed collection of information; (c) ways to enhance the quality, utility, and clarity of the information to be collected; and (d) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques or other forms of information technology. Written comments should be received within 60 days of this notice.

Proposed Project

Interventions to Reduce Shoulder MSDs in Overhead Assembly—New -National Institute for Occupational Safety and Health (NIOSH), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

The mission of the National Institute for Occupational Safety and Health (NIOSH) is to promote safety and health at work for all people through research and prevention. Under Public Law 91-596, sections 20 and 22 (Section 20-22,

Occupational Safety and Health Act of 1970), NIOSH has the responsibility to conduct research to advance the health and safety of workers. In this capacity, NIOSH proposes to conduct a study to assess the effectiveness and cost-benefit of occupational safety and health (OSH) interventions to prevent musculoskeletal disorders (MSDs) among workers in the Manufacturing (MNF) sector.

Musculoskeletal disorders (MSDs) represent a major proportion of injury/illness incidence and cost in the U.S. Manufacturing (MNF) sector. In 2008, 29% of non-fatal injuries and illnesses involving days away from work (DAW) in the MNF sector involved MSDs and the MNF sector had some of the highest rates of MSD DAW cases. The rate for the motor vehicle manufacturing sub-sector (NAICS 3361) was among the highest of MNF sub sectors, with MSD DAW rates that were higher than the general manufacturing MSD DAW rates from 2003-2007. In automotive manufacturing, overhead conveyance of the vehicle chassis requires assembly line employees to use tools in working postures with the arms elevated. These postures are believed to be associated with symptoms of upper limb discomfort, fatigue, and impingement syndromes (Fischer *et al.*, 2007). Overhead working posture, independent of the force or load exerted with the hands, may play a role in the development in these conditions. However, recent studies suggest a more significant role of localized shoulder muscle fatigue in contributing to these disorders. Fatigue of the shoulder muscles may result in changes in normal shoulder kinematics (motion) that affect risk for shoulder impingement disorders (Ebaugh *et al.*, 2006; Chopp *et al.*, 2010).

The U.S. Manufacturing sector has faced a number of challenges including an overall decline in jobs, an aging workforce, and changes in organizational management systems. Studies have indicated that the average age of industrial workers is increasing and that older workers may differ from younger workers in work capacity, injury risk, severity of injuries, and speed of recovery (Kenny *et al.*, 2008; Gall *et al.*, 2004; Restrepo *et al.*, 2006). As the average age of the industrial population increases and newer systems of work organization (such as lean manufacturing) are changing the nature of labor-intensive work, prevention of MSDs will be more critical to protecting older workers and maintaining productivity.

This study will evaluate the efficacy of two intervention strategies for

reducing musculoskeletal symptoms and pain in the shoulder attributable to overhead assembly work in automotive manufacturing. These interventions are, (1) an articulating spring-tensioned tool support device that unloads from the worker the weight of the tool that would otherwise be manually supported, and, (2) a targeted exercise program intended to increase individual employees' strength and endurance in the shoulder and upper arm stabilizing muscle group. As a primary prevention strategy, the tool support engineering control approach is preferred; however, a cost-efficient opportunity exists to concurrently evaluate the efficacy of a preventive exercise program intervention. Both of these intervention approaches have been used in the Manufacturing sector, and preliminary evidence suggests that both approaches may have merit. However, high quality evidence demonstrating their effectiveness, by way of controlled trials, is lacking.

This project will be conducted as a partnership between NIOSH and Toyota Motors Engineering & Manufacturing North America, Inc. (TEMA), with the intervention evaluation study taking place at the Toyota Motor Manufacturing Kentucky, Inc. (TMMK) manufacturing facility in Georgetown, Kentucky. The prospective intervention evaluation study will be conducted using a group-randomized controlled trial multi-time series design. Four groups of 25-30 employees will be established to test the two intervention treatment conditions (tool support, exercise program), a combined intervention treatment condition, and a control condition. The four groups will be comprised of employees working on two vehicle assembly lines in different parts of the facility, on two work shifts (first and second shift). Individual randomization to treatment condition is not feasible, so a group-randomization (by work unit) will be used to assign the four groups to treatment and control conditions. Observations will be made over the 10-month study period and questionnaires will include the Shoulder Rating Questionnaire (SRQ), Disabilities of the Arm, Shoulder and Hand (DASH) questionnaire, a Standardized Nordic Questionnaire for body part discomfort, and a Work Organization Questionnaire. In addition to the questionnaires a shoulder-specific functional capacity evaluation test battery will be administered at 90 and 210 days, immediately pre- and post-intervention, to confirm the efficacy of the targeted exercise program in improving shoulder capacity.