

cases of public health emergency. Estimated average time to complete this form is 1 hour. Based on data regarding the requests received since the last submission, CDC estimates that 5 requests per respondent will be received on an annual basis.

In addition to the standardized forms, this regulation also outlines situations in which an entity must notify or may make a request of the HHS Secretary in writing. An entity may apply to the HHS Secretary for an expedited review of an individual by the Attorney General (42 CFR 73.10(e)). To apply for this expedited review, an entity must submit a request in writing to the HHS Secretary establishing the need for such action. The estimated time to gather the information and submit this request is 30 minutes. CDC has not developed standardized forms to use in the above situations. Rather, the entity should provide the information as requested in the appropriate section of the regulation.

An entity may also apply to the HHS Secretary for an exclusion of an

attenuated strain of a select agent or toxin that does not pose a severe threat to public health and safety (42 CFR 73.3(e)(1) and 73.4(e)(1)). The estimated time to gather the information and submit this request is 1 hour.

As part of the requirements of the Responsible Official, the Responsible Official is required to conduct regular inspections (at least annually) of the laboratory where select agents or toxins are stored. Results of these self-inspections must be documented (42 CFR 73.9(a)(5)). CDC estimates, that, on average, such documentation will take 1 hour.

As part of the training requirements of this regulation, the entity is required to record the identity of the individual trained, the date of training, and the means used to verify that the employee understood the training (42 CFR 73.15(c)). Estimated time for this documentation is 2 hours per principal investigator.

An individual or entity may request administrative review of a decision denying or revoking certification of registration or an individual may appeal

a denial of access approval (42 CFR 73.20). This request must be made in writing and within 30 calendar days after the adverse decision. This request should include a statement of the factual basis for the review. CDC estimates the time to prepare and submit such a request is 4 hours.

An entity must implement a system to ensure that certain records and databases are accurate and that the authenticity of records may be verified (42 CFR 73.17(b)). The time to implement such a system is estimated to average 4 hours.

Prior to issuance of a certificate of registration, CDC inspects entities to ensure compliance with this regulation (42 CFR 73.18). As part of the inspection process, the entity may need to respond to written requests from CDC. CDC estimates the time to prepare and submit a response for the inspection is 8 hours. To estimate the burden, we use the total number of registered entities since each entity will be inspected at least once during the course of their registration.

ESTIMATED ANNUALIZED BURDEN HOURS

CFR reference	Form	Number of respondents	Responses per respondent	Average burden per response (in hours)	Total annual burden (in hours)
73.7(d)	Registration Application	5	1	5	25
73.7(h)(1)	Amendment to Registration Application	264	5	1	1,320
73.19(a)(b)	Notification of Theft, Loss, or Release form.	60	1	1	60
73.5 & 73.6(d-e)/ 73.3 & 73.4(e)(1).	Request for Exemption/Exclusion	5	1	1	5
73.16	Request to Transfer Select Agent or Toxin.	264	4	2	2,112
73.5 & 73.6(a)(b)	Report of Identification of Select Agent or Toxin form.	264	10	1	2,640
73.10(e)	Request expedited review	10	1	1	10
73.9(a)(5)	Documentation of self-inspection	264	1	1	264
73.15(c)	Documentation of training	264	1	2	528
73.20	Administrative Review	15	1	4	60
73.17	Ensure secure recordkeeping system	264	1	4	1,056
73.18	Inspections	264	1	8	2,112
Total					9,657

Dated: March 4, 2008.
 Maryam I. Daneshvar,
 Reports Clearance Officer, Centers for Disease Control and Prevention.
 [FR Doc. E8-5256 Filed 3-14-08; 8:45 am]
 BILLING CODE 4163-18-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day-0920-0630]

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 404-639-5960 and send comments to Dr. Maryam Daneshvar, 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

Work Organization Predictors of Depression in Women—Reinstatement—The National Institute for Occupational Safety and Health (NIOSH), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

Depression is a costly and debilitating occupational health problem. Research has indicated that the costs to an organization of treatment for depression can rival those for heart disease, and major and minor depressive disorders which have been found to be associated with more disability days than other types of health diagnoses. This may be of particular relevance for working women. Various national and international studies indicate that

women in developed countries experience depression at up to twice the rate of men. Studies that have examined this gender difference have focused on social, personality, and genetic explanations while few have explored factors in the workplace that may contribute to the gender differential. Examples of workplace factors that may contribute to depression among women include: Additive workplace and home responsibilities, lack of control and authority, and low paying and low status jobs. Additionally, women are much more likely to face various types of discrimination in the workplace than men, ranging from harassment to inequalities in hiring and promotional opportunities, and these types of stressors have been strongly linked with psychological distress and other negative health outcomes. On the positive side, organizations that are judged by their employees to value diversity and employee development engender lower levels of employee stress, and those that enforce policies against discrimination have more committed employees. Such organizational practices and policies may be beneficial for employee mental health, particularly the mental health of women.

This research focuses on the following questions: (1) Which work organization factors are most predictive of depression in women, and (2) are there measurable work organization factors that confer

protection against depression in women employees?

The research uses a repeated measures, prospective design with data collection at three points (baseline and 1-year and 2-year follow-ups). A 45-minute survey is being administered by telephone to 314 women and men at 16 different organizations. The survey contains questions about traditional job stressors (e.g., changes in workload, social support, work roles), stressors not traditionally examined, but which may be linked with depressive symptoms among women (e.g., roles and responsibilities outside of the workplace, discrimination, career issues) depression symptoms, and company policies, programs and practices. In our previous collection (2002), one Human Resource (HR) representative at each company was also surveyed about company policies, programs and practices. No HR representatives will be contacted for this survey. Analyses will determine which work organization factors are linked with depressive symptoms and what effect the organizational practices/policies of interest have on depression. Findings from this prospective study will also help target future intervention efforts to reduce occupationally-related depression in women workers. This study is being renewed in order to finish data collection. There will be no cost to respondents other than their time.

ESTIMATED ANNUALIZED BURDEN TABLE

Respondents	No. of respondents	No. of responses per respondent	Average burden per response (in hours)	Total burden (in hours)
Employees	314	1	45/60	236

Dated: March 5, 2008.
Maryam I. Daneshvar,
Acting Reports and Clearance Officer, Centers for Disease Control and Prevention.
 [FR Doc. E8-5257 Filed 3-14-08; 8:45 am]
 BILLING CODE 4163-18-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day-08-0109]

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) 639-5960 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

Respiratory Protective Devices—42 CFR 84—Regulation—(0920-0109)—Reinstatement—National Institute for Occupational Safety and Health (NIOSH), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

The regulatory authority for the National Institute for Occupational Safety and Health (NIOSH) certification program for respiratory protective devices is found in the Mine Safety and Health Amendments Act of 1977 (30 U.S.C. 577a, 651 et seq., and 657(g)) and the Occupational Safety and Health Act of 1970 (30 U.S.C. 3, 5, 7, 811, 842(h), 844). These regulations provide the basis for the performance tests and the criteria to respirator manufacturers who submit respirators for testing and certification to be NIOSH-approved. Respirators are used by millions of American construction workers, miners, painters, asbestos removal workers, fabric mill workers, and fire fighters. Improved testing requirements have