

a completed property sale. The possibility of sellers or buyers using the MLS but bypassing brokerage services is already addressed effectively by the Respondent's existing rules that do not distinguish between forms of listing contracts, and does not justify the series of exclusionary rules and policies adopted by MiRealSource. It is possible, of course, that a buyer of an Exclusive Agency Listing may make the purchase without using a selling broker, but this is true for traditional Exclusive Right to Sell Listings as well.

IV. The Proposed Consent Order

The proposed order is designed to ensure that the Respondent does not misuse its market power, while preserving the procompetitive incentives of members to contribute to the MLS.

The proposed order prohibits MiRealSource from adopting or enforcing any rules or policies that deny or limit the ability of MLS members to enter into Exclusive Agency Listings, or any other lawful listing agreements, with sellers of properties. More specifically, the proposed order prohibits MiRealSource from preventing its members from offering or accepting Exclusive Agency Listings or other lawful listing agreements; cooperating with Listing Brokers or agents that offer or accept Exclusive Agency Listings or other lawful listing agreements; publishing Exclusive Agency Listings or other lawful listing agreements on the MLS and approved Web sites; publishing their information concerning listings on public real estate Web sites, including but not limited to <http://www.FSBO.com>; requiring members to have a physical office; and offering unbundled real estate brokerage services, including but not limited to requiring MiRealSource Shareholders to provide a minimum set of real estate brokerage services. The proposed order also prohibits MiRealSource from denying or restricting the services of the MLS to Exclusive Agency Listings or other lawful listings in any way that such services of the MLS are not denied or restricted to Exclusive Right to Sell Listings; or treating Exclusive Agency Listings, or any other lawful listings, in a less advantageous manner than Exclusive Right to Sell Listings, including but not limited to, any policy, rule or practice pertaining to the transmission, downloading, or displaying of information pertaining to such listings.

In addition to these substantive provisions, the proposed order states that, within forty-five days after it becomes final, Respondent shall have

conformed its rules to the substantive provisions of the order. Respondent is further required to notify its members of the applicable order through its usual business communications and its Web site. The proposed order requires notification to the Commission of changes in the respondent's structure, and periodic filings of written reports concerning compliance. The relief in the proposed consent order ensures that the Respondent cannot revert to the old rules or policies, or engage in future variations of the challenged conduct.

The proposed order applies to MiRealSource and entities it owns or controls, including its respective MLS and any affiliated Web site it operates. The order does not prohibit members, or other independent persons or entities that receive listing information from Respondent, from making independent decisions concerning the use or display of such listing information on member or third-party Web sites, consistent with any contractual obligations to Respondent.

The proposed order will expire in 10 years.

By direction of the Commission.

Donald S. Clark,
Secretary.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day-07-0527]

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 or send comments to Joan F. Karr, 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

Human Exposure to Cyanobacterial Toxins in Water (OMB No. 0920-0527)—Reinstatement—National Center for Environmental Health (NCEH), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

Cyanobacteria (blue-green algae) can be found in terrestrial, fresh, brackish, or marine water environments. Some species of cyanobacteria produce toxins that may cause acute or chronic illnesses (including neurotoxicity, hepatotoxicity, and skin irritation) in humans and animals (including other mammals, fish, and birds). A number of human health effects, including gastroenteritis, respiratory effects, skin irritations, allergic responses, and liver damage, are associated with the ingestion of or contact with water containing cyanobacterial blooms. Although the balance of evidence, in conjunction with data from laboratory animal research, suggests that cyanobacterial toxins are responsible for a range of human health effects, there have been few epidemiologic studies of this association.

During August 2006, we conducted our first study to assess exposure to microcystins in recreational waters with a bloom of *Microcystis aeruginosa*. We recruited 104 people who gave informed consent to participate. Ninety seven people did their recreational activities on Lake 1, which had a confirmed *M. aeruginosa* bloom, and 7 others did their activities on Lake 2, which had no bloom. Study participants completed a pre-activity questionnaire, a post-activity questionnaire, provided a 10-ml blood sample, and completed a telephone symptom survey 7-10 days after exposure. The concentrations of microcystins in Lake 1 ranged from 2 to 5 ug/L and in Lake 2 were all below the limit of detection (LOD). When we designed the study, we calculated that a person exposed to recreationally-generated aerosols from water containing 10 ug/L of microcystins should have levels of microcystins in

their blood. However, the microcystin concentrations in Lake 2 were below the LOD and in Lake 1 were actually 2ug/L to 5ug/L, much lower than we anticipated based on data from the previous week. Thus, the recreational exposures were not likely high enough for us to quantify microcystins in blood and the serum samples were all below the LOD for microcystins.

For the new data collection, we will recruit 100 study participants who are at risk for swallowing water or inhaling spray (*i.e.*, water skiers, jet skiers, people sailing small boats) and who

would normally be doing these activities, even in the presence of a bloom. We may recruit people who train for organized swimming events (*e.g.*, triathlons) in lakes. In addition, we will recruit 50 study participants from lakes with no blooms as a comparison group to assess the health effects associated with recreational activities on "clean" lakes. Study participants will be asked to sign a consent form, complete a symptom survey before and after doing their recreational water activities, provide one 10-ml whole blood sample after their recreational activities, and

complete a telephone symptom survey 8–10 days after doing study activities.

The purpose of the new data collection is to continue assessing the public health impact of exposure to the cyanobacterial toxins, microcystins, during recreational activities. We will examine the extent of human exposure to microcystins present in recreational waters and associated aerosols and whether serum levels of microcystins can be used as a biomarker of exposure.

There is no cost to the respondents other than their time.

ESTIMATE OF ANNUALIZED BURDEN HOURS

Forms	Number of respondents	Number of responses per respondent	Average burden per response (in hours)	Total burden hours
Screening Questionnaire	188	1	10/60	31
Pre-exposure Questionnaire	150	1	10/60	25
Post-exposure Questionnaire	150	1	10/60	25
10-day post exposure Questionnaire	150	1	10/60	25
Total				106

Dated: February 6, 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

[60Day-07-0630]

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 Joan Karr, CDC Acting Reports Clearance Officer, 1600 Clifton Road, MS-D74, Atlanta, GA 30333 or send an e-mail 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—Extension—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 both major depressive disorder and forms of minor depression 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 repeated measures, prospective design with data collection at three points (baseline and 1-year and 2-year follow-ups). A 45-minute survey