Sherette.funncoleman@hhs.gov, or call the Reports Clearance Officer on (202) 690–5683. Send written comments and recommendations for the proposed information collections within 30 days of this notice directly to the OS OMB Desk Officer; faxed to OMB at 202–395–5806.

FOR FURTHER INFORMATION CONTACT: To request additional information, please contact *Sherette.funncoleman@hhs.gov*, or call the Reports Clearance Office on (202) 690–6162.

SUPPLEMENTARY INFORMATION:

Title: FedStrive Employee Wellness Program Social Media Survey.

Abstract: The information collection activity will garner qualitative customer and stakeholder feedback in an efficient, timely manner, in accordance with the Administration's commitment to improving service delivery. By qualitative feedback we mean information that provides useful insights on perceptions and opinions, but are not statistical surveys that yield quantitative results that can be generalized to the population of study. This feedback will provide insights into customer or stakeholder perceptions, experiences and expectations, provide an early warning of issues with service, or focus attention on areas where communication, training or changes in operations might improve delivery of products or services. These collections will allow for ongoing, collaborative and actionable communications between the Agency and its customers and stakeholders. It will also allow feedback to contribute directly to the improvement of program management.

Feedback collected under this generic clearance will provide useful information, but it will not yield data that can be generalized to the overall population. This type of generic clearance for qualitative information will not be used for quantitative information collections that are designed to yield reliably actionable results, such as monitoring trends over time or documenting program performance. Such data uses require more rigorous designs that address: the target population to which generalizations will be made, the sampling frame, the sample design (including stratification and clustering), the precision requirements or power calculations that justify the proposed sample size, the expected response rate, methods for assessing potential nonresponse bias, the protocols for data collection, and any testing procedures that were or will be undertaken prior fielding the study. Depending on the degree of influence the results are likely

to have, such collections may still be eligible for submission for other generic mechanisms that are designed to yield quantitative results.

The Agency received 0 comments were received in response to the 60-day notice published in the **Federal Register** of December 22, 2010 (75 FR 80542).

Below we provide the Department of Health and Human Services, projected average estimates for the next three years:¹

Current Actions: New collection of information.

Type of Review: New Collection. Affected Public: Individuals and Households, Businesses and Organizations, State, Local or Tribal Government.

Average Expected Annual Number of Activities

Respondents: 3000.
Annual Responses: 3000.
Frequency of Response: Once per Request.

Average Minutes per Response: 5. *Burden Hours:* 250 total.

An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid Office of Management and Budget control number.

Keith Tucker,

Office of the Secretary, Paperwork Reduction Act Clearance Officer.

[FR Doc. 2011–25143 Filed 10–3–11; 8:45 am] **BILLING CODE P**

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Office of the Secretary

Findings of Research Misconduct

AGENCY: Office of the Secretary, HHS. **ACTION:** Notice.

SUMMARY: Notice is hereby given that the Office of Research Integrity (ORI) has taken final action in the following case:

Scott Weber, Ed.D., MSN, University of Pittsburgh: Based on the letters from the Research Integrity Officer at the University of Pittsburgh (UP), ORI's

oversight review, and an admission by the Respondent, ORI found that Dr. Scott Weber, former Assistant Professor, Health and Community Systems, School of Nursing, UP, engaged in research misconduct by (1) plagiarizing text and falsifying data from two publications supported by U.S. Public Health Service (PHS) funding (P30 MH60570; HS5 SM52671; PHS employee generated article) in two unpublished manuscripts, and (2) including significant portions of that plagiarized text in two grant applications to the National Institutes of Health (NIH) (1 L30 NR010444-01; 1 R03 HD062761-

ORI found that the Respondent engaged in research misconduct by plagiarizing text, falsifying data and references, and fabricating data from two publications (Mufson, L., Dorta, K.P., Wickramaratne, P., Nomura, Y., Olfson, M., Weissman, M.M. "A randomized effectiveness trial of interpersonal psychotherapy for depressed adolescents." Arch Gen Psychiatry 61(6):577-84, 2004 June; hereafter referred to as "Mufson et al. 2004;" and Cho, M.J., Mościcki, E.K., Narrow, W.E., Rae, D.S., Locke, B.Z., Regier, D.A. "Concordance between two measures of depression in the Hispanic Health and Nutrition Examination Survey." Soc Psychiatry Psychiatr Epidemiol. 28(4):156-63, 1993 August; hereafter referred to as "Cho et al... 1993") supported by PHS in two journal article submissions. Specifically, ORI found that the Respondent plagiarized more than 90 percent of the text from Mufson *et al.* 2004 in a manuscript entitled "A randomized effectiveness trial of psychiatric-mental health nurse practitioner-administered interpersonal psychotherapy for sexual minority adolescents with depression in primary care clinics" and submitted to the Journal of the American Academy of Nurse Practitioners (JAANP MS). Furthermore, the Respondent plagiarized approximately 66 percent of the text from Cho et al. 1993 in a manuscript entitled "Assessing the diagnostic predictive power of a screening tool for depression: Concordance between the CES-D and DIS in the Parent Identity Survey" and submitted to the Journal of GLBT Family Studies (IGMS MS).

In both manuscripts, the Respondent falsified and fabricated tables and figures by using all or nearly all of the data in tables and graphs from the plagiarized articles while altering numbers and changing text to represent data as if from another subject population; he also copied most of the original bibliographic references but

¹The 60-day notice included the following estimate of the aggregate burden hours for this generic clearance federal-wide:

 $[\]label{lem:average-expected-expected} Annual \ Number\ of\ Activities: 25,000.$

Average Number of Respondents per Activity: 200.

Annual Responses: 5,000,000. Frequency of Response: Once per Request. Average Minutes per Response: 30. Burden Hours: 2,500,000.

falsified 35% of the copied references from *JAANP MS* and 25% of the copied references from *JGMS MS*, by changing volume numbers and/or publication years, apparently to hinder detection of the plagiarism. The data fabrication occurred when the Respondent altered or added values to Table 2 in each manuscript describing the demographic characteristics of the study population that was never studied.

ORI also finds that the Respondent engaged in research misconduct by plagiarizing text from Cho *et al.* 1993 in two NIH grant applications (1 L30 NR010444–01 and 1 R03 HD062761–01) by copying substantial word-for-word portions of the text describing the test instrument to be used in the proposed study without citing the Cho *et al.* 1993 paper.

Dr. Weber has voluntarily agreed for a period of three (3) years, beginning on September 7, 2011:

- (1) To exclude himself from any contracting or subcontracting with any agency of the United States Government and from eligibility or involvement in nonprocurement programs of the United States Government referred to as "covered transactions" pursuant to HHS' Implementation (2 CFR part 376 et seq.) of OMB Guidelines to Agencies on Governmentwide Debarment and Suspension, 2 CFR part 180 (collectively the "Debarment Regulations"); and
- (2) to exclude himself from serving in any advisory capacity to PHS including, but not limited to, service on any PHS advisory committee, board, and/or peer review committee, or as a consultant.

FOR FURTHER INFORMATION CONTACT:

Director, Division of Investigative Oversight, Office of Research Integrity, 1101 Wootton Parkway, Suite 750, Rockville, MD 20852, (240) 453–8800.

John Dahlberg, Ph.D.,

Director, Division of Investigative Oversight, Office of Research Integrity.

[FR Doc. 2011–25537 Filed 10–3–11; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day-11-0213]

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 project or to obtain a copy of data collection plans and instruments, call the CDC Reports Clearance Officer on 404-639-5960 or send comments to Daniel Holcomb, CDC 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

National Vital Statistics Report Forms (OMB No. 0920–0213, Expiration Date April 30, 2012)—Extension—National Center for Health Statistics (NCHS), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

The compilation of national vital statistics dates back to the beginning of the 20th century and has been conducted since 1960 by the Division of Vital Statistics of the National Center for Health Statistics, CDC. The collection of the data is authorized by 42 U.S.C. 242k. This submission requests approval to collect the monthly and annually summary statistics for three years.

The Monthly Vital Statistics Report forms provide counts of monthly occurrences of births, deaths, infant deaths, marriages, and divorces. Similar data have been published since 1937 and are the sole source of these data at the National level. The data are used by the Department of Health and Human Services and by other government, academic, and private research and commercial organizations in tracking changes in trends of vital events. Respondents for the Monthly Vital Statistics Report Form are registration officials in each State and Territory, the District of Columbia, and New York City. In addition, local (county) officials in New Mexico who record marriages occurring and divorces and annulments granted in each county of New Mexico will use this form. This form is also designed to collect counts of monthly occurrences of births, deaths, infant deaths, marriages, and divorces immediately following the month of occurrence.

The Annual Vital Statistics Occurrence Report Form collects final annual counts of marriages and divorces by month for the United States and for each State. The statistical counts requested on this form differ from provisional estimates obtained on the Monthly Vital Statistics Report Form in that they represent complete counts of marriages, divorces, and annulments occurring during the months of the prior year. These final counts are usually available from State or county officials about eight months after the end of the data year. The data are widely used by government, academic, private research, and commercial organizations in tracking changes in trends of family formation and dissolution. Respondents for the Annual Vital Statistics Occurrence Report Form are registration officials in each State and Territory, the District of Columbia, and New York City.

There are no costs to respondents other than their time.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondent	Form name	Number of respondents	Number of responses per respondent	Average burden per response (in hours)	Total burden hours
State, Territory and New Mexico County officials.	Monthly Vital Statistics Report	91	12	10/60	182