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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60-Day-20-20AZ; Docket No. CDC-2019-
0099]

Proposed Data Collection Submitted for Public Comment and Recommendations

AGENCY: Centers for Disease Control and
Prevention (CDC), Department of Health
and Human Services (HHS).

ACTION: Notice with comment period.

SUMMARY: The Centers for Disease
Control and Prevention (CDC), as part of
its continuing effort to reduce public
burden and maximize the utility of
government information, invites the
general public and other Federal
agencies the opportunity to comment on
a proposed and/or continuing
information collection, as required by
the Paperwork Reduction Act of 1995.
This notice invites comment on a
proposed information collection project
titled Evaluation of the Effectiveness of
the Training and Education Modules in
the North American Fatigue
Management Program, which is an
observational study evaluating 180 long-
haul and regional truck drivers in a
naturalistic driving study over eight
months. Questionnaires, in-vehicle
monitor system, Actigraphy devices,
and smartphones will be used in the
data collection.

DATES: CDC must receive written
comments on or before January 3, 2020.

ADDRESSES: You may submit comments,
identified by Docket No. CDC-2019-
0099 by any of the following methods:

- *Federal eRulemaking Portal:*
Regulations.gov. Follow the instructions
for submitting comments.

- *Mail:* Jeffrey M. Zirger, Information
Collection Review Office, Centers for
Disease Control and Prevention, 1600
Clifton Road NE, MS-D74, Atlanta,
Georgia 30329.

Instructions: All submissions received
must include the agency name and
Docket Number. CDC will post, without
change, all relevant comments to
Regulations.gov.

Please note: Submit all comments through
the Federal eRulemaking portal

(*regulations.gov*) or by U.S. mail to the
address listed above.

FOR FURTHER INFORMATION CONTACT: To
request more information on the
proposed project or to obtain a copy of
the information collection plan and
instruments, contact Jeffrey M. Zirger,
Information Collection Review Office,
Centers for Disease Control and
Prevention, 1600 Clifton Road NE, MS-
D74, Atlanta, Georgia 30329; phone:
404-639-7570; Email: *omb@cdc.gov*.

SUPPLEMENTARY INFORMATION: Under the
Paperwork Reduction Act of 1995 (PRA)
(44 U.S.C. 3501-3520), Federal agencies
must obtain approval from the Office of
Management and Budget (OMB) for each
collection of information they conduct
or sponsor. In addition, the PRA also
requires Federal agencies to provide a
60-day notice in the **Federal Register**
concerning each proposed collection of
information, including each new
proposed collection, each proposed
extension of existing collection of
information, and each reinstatement of
previously approved information
collection before submitting the
collection to the OMB for approval. To
comply with this requirement, we are
publishing this notice of a proposed
data collection as described below.

The OMB is particularly interested in
comments that will help:

1. Evaluate whether the proposed
collection of information is necessary
for the proper performance of the
functions of the agency, including
whether the information will have
practical utility;
2. Evaluate the accuracy of the
agency's estimate of the burden of the
proposed collection of information,
including the validity of the
methodology and assumptions used;
3. Enhance the quality, utility, and
clarity of the information to be
collected; and
4. Minimize the burden of the
collection of information on those who
are to respond, including through the
use of appropriate automated,
electronic, mechanical, or other
technological collection techniques or
other forms of information technology,
e.g., permitting electronic submissions
of responses.
5. Assess information collection costs.

Proposed Project

Evaluation of the Effectiveness of the
Training and Education Modules in the
North American Fatigue Management
Program—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. Reducing fatigue-
related crashes is one of the top 10
changes needed to reduce transportation
accidents and save lives identified by
the National Transportation Safety
Board (NTSB) for 2017-2018 and a
National Occupational Research Agenda
(NORA) priority.

Fatigue is a preventable cause of
crashes. The North American Fatigue
Management Program (NAFMP) was
developed by the Federal Motor Carrier
Safety Administration, Transport
Canada, and other entities to address
commercial motor vehicle (CMV) driver
fatigue through a comprehensive
approach that delivers prevention
information to carriers, dispatchers,
drivers, and family members. In 2015,
the National Academy of Sciences
published the report "Commercial
motor vehicle driver fatigue, long-term
health, and highway safety research
needs" that identified the need for fully
evaluating the NAFMP so that
recommendations for implementation of
NAFMP are supported by scientific
evidence. NIOSH is collaborating with
the FMCSA to ensure the success of the
proposed study.

Data will be collected from CMV
drivers (hereafter referred to as "driver")
during their application to participate in
the study, briefing session, study
participation, and debriefing session.
Data collection will primarily focus on
driving performance, sleep, and
sleepiness. These outcomes will be
compared between pre-rollout of the
NAFMP (in which drivers will operate
as they did before their participation in
the study) and after the rollout of the
NAFMP training and education modules
(in which drivers and managers will
operate with increased knowledge,
strategies, and techniques to reduce
their fatigue). All drivers interested in
participating in the study may complete
the application. A briefing session will
be scheduled with drivers who are
found eligible for the study. During the
briefing session, drivers who provide
informed consent will be enrolled in the
study. Drivers will have a debriefing
session if a driver chooses to withdraw
from the study early or upon completion
of the eight-month participation period.

The sample of drivers in the study
will include those employed as drivers
at the participating carriers. Drivers who
have a valid Class-A commercial
driver's license (CDL) and work at the
participating company in regional and

long-haul operations for at least one year will be eligible for the study. A convenience sample of 180 eligible drivers over a two-year period will be recruited to participate in the study. The study sample will include approximately 90 regional and 90 long-haul drivers. There will be no required minimum number of female or minority drivers to be included in the study.

Data will be collected during each phase: (1) In the application, drivers will be asked to provide their name and contact information (home address, telephone number, and email address) to allow contact from the research team regarding their eligibility for the study. (2) In the briefing session, drivers will be asked to complete the Background Questionnaire. (3) During the study, information collection will occur through several streams: (a) A real-time

fatigue monitoring system installed in the participating driver's vehicle; (b) Smart phone apps to collect psychomotor vigilance test, Karolinska Sleepiness Scale, sleep log, difficulty of drive scale, degree of drive hazards scale, a fatigue scale, and a stress scale; (c) an electronic logging device to collect data on the driver's duty and driving; (d) a wrist actigraphy to collect data on driver sleep and wake times. Drivers will be asked to sync the actigraph with a smartphone app daily; (e) smartphone or web-based questionnaires including Exercise and Food Consumption Questionnaire, the quality of life short form 36 version-2 questionnaire (SF-36v2), Family Interactions Questionnaire, and Job Descriptive Index. These will be completed by drivers at four different intervals, including the beginning (first

week) and middle (second month) of the baseline phase, and the middle (fifth month) and end (eighth month) of the intervention phase; (f) A questionnaire to assess corporate practices and corporate safety climate will be given to managers at the participating carriers. These will be completed by managers at the beginning (first week) of the study and end (eighth month) of the intervention phase; and (g) during the field study, carriers will be asked to provide information concerning crashes and roadside violations occurring during each driver's period of study participation. Administrative cost information (e.g., equipment, labor, etc.) will also be collected from the carrier to evaluate cost-benefit of the intervention. The total annualized burden hours requested is 5,139.

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 (in hours)
Carrier Management	Participation Agreements	1	1	1	1
	Monthly Roadside Violations, ELD, Crash Reports, Administrative Costs.	1	16	30/60	8
Drivers	Corporate Practices Questionnaire	10	1	45/60	8
	Application to Participate	150	1	12/60	30
	Actigraph Training	90	1	10/60	15
	Background Questionnaire	90	1	45/60	68
	Daily Smartphone Questions	90	720	1/60	1,037
	PVT	90	720	3/60	3,240
	Exercise and Food Consumption Questionnaire.	90	4	20/60	120
	SF-36v2	90	4	30/60	180
	Family Interactions Questionnaire	90	4	15/60	90
	Job Descriptive Index	90	4	30/60	180
Post-Study Questionnaire	90	1	1	90	
Phone Briefings	90	8	6/60	72	
Total					5,139

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day-20-19DO]

Agency Forms Undergoing Paperwork Reduction Act Review

In accordance with the Paperwork Reduction Act of 1995, the Centers for Disease Control and Prevention (CDC)

has submitted the information collection request titled National Surveillance of Community Water Systems and Corresponding Populations with the Recommended Fluoridation Level to the Office of Management and Budget (OMB) for review and approval. CDC previously published a "Proposed Data Collection Submitted for Public Comment and Recommendations" notice on December 6, 2018 to obtain comments from the public and affected agencies. CDC received two comments related to the previous notice. This notice serves to allow an additional 30 days for public and affected agency comments.

CDC will accept all comments for this proposed information collection project. The Office of Management and Budget

is particularly interested in comments that:

(a) Evaluate whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information will have practical utility;

(b) Evaluate the accuracy of the agency's estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used;

(c) Enhance the quality, utility, and clarity of the information to be collected;

(d) Minimize the burden of the collection of information on those who are to respond, including, through the use of appropriate automated, electronic, mechanical, or other