Health Assessment and Consultation, Agency for Toxic Substances and Disease Registry, 1600 Clifton Road, NE., Mailstop E–32, Atlanta, Georgia 30333, telephone (404) 498–0007.

SUPPLEMENTARY INFORMATION: The most recent list of completed public health assessments was published in the Federal Register on May 9, 2002 [67 FR] 31308]. This announcement is the responsibility of ATSDR under the regulation, Public Health Assessments and Health Effects Studies of Hazardous Substances Releases and Facilities [42] CFR Part 90]. This rule sets forth ATSDR's procedures for the conduct of public health assessments under section 104(i) of the Comprehensive Environmental Response, Compensation, and Liability Act (CERCLA), as amended by the Superfund Amendments and Reauthorization Act (SARA) [42 U.S.C. 9604(i)].

Availability

The completed public health assessments and addenda are available for public inspection at the Division of Health Assessment and Consultation, Agency for Toxic Substances and Disease Registry, Building 33, Executive Park Drive, Atlanta, Georgia (not a mailing address), between 8 a.m. and 4:30 p.m., Monday through Friday except legal holidays. The completed public health assessments are also available by mail through the U.S. Department of Commerce, National Technical Information Service (NTIS), 5285 Port Royal Road, Springfield, Virginia 22161, or by telephone at (703) 605-6000. NTIS charges for copies of public health assessments and addenda. The NTIS order numbers are listed in parentheses following the site names.

Public Health Assessments Completed or Issued

Between April 1, 2002 and June 30, 2002, public health assessments were issued for the sites listed below:

NPL Sites

District of Columbia

Washington Navy Yard (PB2002–104375).

New Jersey

Emmell's Septic Landfill (PB2002–104386).

Texas

Malone Service Company—Swan Lake Plant (PB2002–103158).

Washington

Cenex Supply and Marketing, Incorporated (a/k/a Western Farmers, Incorporated) (PB2002–104385).

Non NPL Petitioned Sites

None.

Dated: July 18, 2002.

Georgi Jones,

Director, Office of Policy and External Affairs, Agency for Toxic Substances and Disease Registry.

[FR Doc. 02–18779 Filed 7–24–02; 8:45 am]

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day-02-70]

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 the CDC Reports Clearance Officer on (404) 498–1210.

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. Send comments to Anne O'Connor, CDC Assistant Reports Clearance Officer, 1600 Clifton Road, MS-D24, Atlanta, GA 30333. Written comments should be received within 60 days of this notice.

Proposed Project: Antineoplastic Drug Exposure: Effectiveness of Guidelines— New—The National Institute for Occupational Safety and Health (NIOSH), Centers for Disease Control and Prevention (CDC). Antineoplastic, chemotherapeutic, or cytostatic drugs

are widely used in the treatment of cancer. These drugs possess mutagenic, teratogenic, and carcinogenic properties, cause organ damage, and affect reproductive function. Healthcare workers such as pharmacists and nurses who handle, prepare, and administer these drugs are at increased risk of adverse health effects from these agents, if exposed. The Occupational Safety and Health Administration (OSHA) developed guidelines for healthcare workers for the safe handling of antineoplastic drugs in 1986 and revised those guidelines again in 1995. However, recent studies suggest that the guidelines have not been effective in preventing exposure. A 1999 industrial hygiene evaluation of six cancer centers in the U.S. and Canada reported that 75% of the wipe test samples in the pharmacy were found to have detectable levels of antineoplastic drugs. Similar findings were reported in the Netherlands, which has similar guidelines. In addition, healthcare workers may assume that gloves designed for bloodborne pathogen protection will also prevent drug exposure which is often not the case. Since air concentrations of antineoplastic drugs in many of the studies have been low to non-detectable, it appears that the dermal route may be an important consideration for internal absorption.

Numerous studies, including those after the OSHA guidelines were revised in 1995, have demonstrated adverse health effects from healthcare workers' exposure to antineoplastic agents. The most common endpoints have been either markers of exposure, such as metabolites in the urine, or genotoxic markers, such as micronuclei, sister chromatid exchange, and chromosomal aberrations. Female reproductive adverse effects have also been shown to occur with healthcare workers' exposure to antineoplastic drugs. Not only have spontaneous abortion and miscarriage been reported, but changes in the menstrual cycle have been demonstrated as well. Based upon animal and human data, one study estimated that exposure to cyclophosphamide by healthcare workers increases the risk of leukemia cases by 17-100 new cases/million workers/10 years.

This project addresses the continuing concern of healthcare workers' exposure to antineoplastic agents. This is a multifaceted project that involves environmental sampling of the workplace and the collection of biological samples to determine how much of the agent is absorbed and if there are any early biological effects

from that exposure. Biological measurements or biomarkers can detect effects of exposure long before a disease can be diagnosed. A questionnaire will be administered to determine confounders and other conditions that might affect exposure such as work history and work practices. This project

will recruit oncology nurses, pharmacists, and pharmacy technicians and will be conducted in collaboration with the University of Maryland, the University of North Carolina, and the M.D. Anderson Cancer Center.

By utilizing a battery of sensitive biomarkers, the effects of low-level chronic exposure to antineoplastic agents can be determined. Using the results of the proposed study, exposures can be minimized or eliminated before adverse health effects occur. Ultimately, the study will contribute to the prevention of occupational disease from antineoplastic drug exposure. There are no costs to respondents.

Survey	Number of respondents	Number of re- sponses/re- spondent	Average bur- den/response (in hours)	Total burden (in hours)
Genotoxicity Immunotocixicity Study*	150 150	1 1	1 225/60	150 562.5
Total				714.50

This part of the study involves the participant, after informed consent, voluntarily providing blood and urine samples and responding to a questionnaire concerning medical history, work history, and work practices to identify study eligibility, past exposures, and confounders.

Dated: July 18, 2002.

Nancy E. Cheal,

Acting Associate Director for Policy, Planning and Evaluation, Centers for Disease Control and Prevention.

[FR Doc. 02–18781 Filed 7–24–02; 8:45 am] **BILLING CODE 4163–18–P**

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day-02-71]

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 the CDC Reports Clearance Officer on (404)498–1210.

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. Send comments to Anne O'Connor, CDC Assistant Reports Clearance Officer, 1600 Clifton Road, MS–D24, Atlanta, GA 30333. Written comments should be received within 60 days of this notice.

Proposed Project: Adult and Pediatric HIV/AIDS Confidential Case Reports (CDC 50.42A, 50.42B)—New—National Center for HIV, STD, and TB Prevention (NCHSTP), Centers for Disease Control and Prevention (CDC). This data collection system was formerly included and approved under the National Disease Surveillance Program, OMB No. 0920-0009, National Center for Infectious Disease (NCID), CDC. CDC is seeking a 3-year OMB approval to continue data collection of the HIV/ AIDS case reports, with revisions of the report forms to collect race and ethnicity data in adherence to OMB Statistical Policy Directive 15, Race and Ethnic Standards for Federal Statistics and Administrative Reporting.

The National Adult and Pediatric HIV/AIDS Confidential Case Reports are collected as part of the HIV/AIDS Surveillance System. CDC in collaboration with health departments in the states, territories, and the District of Columbia, conducts national surveillance for cases of human immunodeficiency virus (HIV) infection and the acquired immunodeficiency syndrome (AIDS), the end-stage of disease caused by infection with HIV. HIV/AIDS surveillance data collection by CDC is authorized under sections 301 and 306 of the Public Health Service Act (42 U.S.C. 241 and 242k).

Currently, 55 states (areas/territories) mandate and collect AIDS surveillance

data. In addition, 35 areas mandate and collect surveillance data on HIV cases which have not progressed to AIDS in adults/adolescents and/or children using the HIV/AIDS case report forms. The purpose of HIV/AIDS surveillance data is to monitor trends in HIV/AIDS and describe the characteristics of infected persons (e.g., demographics, modes of exposure to HIV, manifestations of severe HIV disease, and deaths due to AIDS). Because HIV infection results in untimely death and most often infects younger adults in the prime years of life, large amounts of federal, state, and local government funding have been allocated to address all aspects of HIV infection, including prevention and treatment. HIV/AIDS surveillance data are widely used at all government levels to assess the impact of HIV infection on morbidity and mortality, to allocate medical care resources and services, and to guide prevention and disease control activities.

HIV/AIDS reports are sent to state/local health departments by laboratories, physicians, hospitals, clinics, and other health care providers using standard adult and pediatric case report forms. Areas use a microcomputer system developed by CDC (the HIV/AIDS Reporting System, HARS) to store and analyze data, as well as transmit encrypted data to CDC. An HIV program area module (PAM) for the National Electronic Disease Surveillance System (NEDSS) is in the early development stage and will replace HARS when it is complete.

In order to adhere to OMB Directive 15, the proposed data collection form will collect race and ethnicity separately, collect multiple races, and

[†] In the reproductive health part of the study and after informed consent, women are being asked to voluntarily give a daily urine sample for approximately 45 days and keep track of their menstrual cycle by entries into a diary. In addition, a short questionnaire is given to each participant to determine eligibility for inclusion into the study and confounders of hormone analysis.