

trading programs and trading methods could reasonably expect consistent investment returns of \$2,500 to \$3,500 per week; that users of respondents' trading programs and trading methods could reasonably expect to succeed at day trading for a lifetime of profitable and enjoyable trading; and that testimonials appearing in the advertisements for respondents' trading programs and trading methods reflected the typical or ordinary experience of members of the public who use the program. In addition, the complaint alleges that respondents misrepresented that users of respondents' trading programs and trading methods could trade in volatile markets with LOW RISK.

The proposed consent order contains provisions designed to prevent respondents from engaging in similar acts and practices in the future.

Part I of the proposed order requires respondents to have a reasonable basis substantiating any representation that users of respondents' day trading program can reasonably expect to earn large profits: (1) That users of respondents' trading program or trading method can reasonably expect to earn large profits, or as much as \$2,000 to \$5,000 per day on some days; (2) that users of respondents' trading program or trading method can reasonably expect to earn profits of \$500 to \$750 or more per day; (3) that users of respondents' trading program or trading method can reasonably expect to approach trading as a business and earn a consistent living from the markets; and (4) that users of respondents' trading program or trading method can reasonably expect to trade in volatile markets with low risk. Part I also requires respondents to possess a reasonable basis substantiating claims about the amount of earnings, income, or profit that a prospective user of any trading program or trading method could reasonably expect to attain, or about any financial benefit or other benefit from the purchase or use of any such trading program or trading method.

Part II of the proposed order prohibits respondents from misrepresenting that users of any trading program can reasonably expect to trade with little or no financial risk and from misrepresenting the extent of risk to which users of any such program are exposed.

Part III of the proposed order requires respondents to disclose, clearly and conspicuously, "DAYTRADING involves HIGH RISKS and YOU can LOSE a lot of money." in close proximity to any representation they make about the financial benefits of any

trading program. This disclosure is in addition to, and not instead of, any other disclosure that respondents may be required to make.

Part IV of the proposed order prohibits respondents from representing without a reasonable basis that the experience represented by any user, testimonial or endorsement of any trading program represents the typical or ordinary experience of members of the public who use the program; or respondents must disclose either what the generally expected results would be for users of the trading program, or the limited applicability of the endorser's experience to what users may generally expect to achieve, that is, that users should not expect to experience similar results.

Parts V and VI of the proposed order require respondents to keep copies of relevant advertisements and materials substantiating claims made in the advertisements and to provide copies of the order to certain personnel. Part VII requires R.S. of Houston Workshop to notify the Commission of any changes in the corporate structure that might affect compliance with the order. Parts VIII and IX require that individual respondents Ronald J. Schoemmell and Valdimar Thorkeleson, respectively, to notify the Commission of changes in their employment status for a period of seven years. Part X requires respondents to file compliance reports with the Commission. Part XI provides that the order will terminate after twenty (20) years under certain circumstances.

The purpose of this analysis is to facilitate public comment on the proposed order. It is not intended to constitute an official interpretation of the agreement and proposed order or to modify in any way their terms.

By direction of the Commission.

Donald S. Clark,

Secretary.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Proposed Collection; Comment Request Drug Accountability Form and Drug Transfer Form

SUMMARY: In compliance with the requirement of section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995 for public comment on proposed data collection projects, the National Institutes of Health (NIH), National Cancer Institute (NCI) will publish

periodic summaries of proposed projects to be submitted to the Office of Management and Budget (OMB) for review and approval.

Proposed Collection

Title: Drug Accountability Form and Drug Transfer Form.

Type of Information Collection

Request: Revision. (OMB No. 0925-0240, expires 4/30/2002).

Need and use of Information

Collection: The regulations of the Food and Drug Administration (FDA) require investigators to establish a record of the receipt, use, and disposition of all investigational agents. The National Cancer Institute (NCI), as a sponsor of investigational drug trials, has the responsibility for assuring to the FDA that systems for drug accountability are being maintained by investigators in its clinical trials program. In order to fulfill these requirements, we have developed a standardized investigational Drug Accountability Report Form (NIH 2564) designed to account for drug inventories and usage by protocol. The Transfer Investigational Drug Form (NIH-2564-1) permits intra-institutional transfer of agents to NCI approved protocols for use by the investigator or other NCI registered investigators on approved protocols. The data obtained from the drug accountability record is used to track the dispensing of investigational anticancer drugs from receipt from NCI to dispensing or administration to patients. NCI uses the accountability data to ensure that investigational drug supplies are not diverted for inappropriate protocol or patient use. The drug accountability information is used to validate patient protocol reporting forms during site audits conducted at each of the Cooperative Groups. The intent is to ensure the investigational agents are used according to protocol guidelines and to ensure the patient's safety and protection.

Frequency of response: Daily.

Affected public: State or local governments, businesses or other for profit, Federal agencies or employees, non-profit institutions, and small business or organizations. *Types of Respondents:* Investigators and their designees, pharmacists, nurses, pharmacy technicians, data managers. The annual reporting burden is divided into two major areas. These are the audits of Drug Accountability Forms by Government and its contractors and the use of the forms by clinical research sites. The burden is as follows: The annualized respondents' burden for record keeping is estimated to require 3,648 hours for drug accountability and

120 hours for drug transfer. The reporting burden is the average time (4 minutes or 0.1 hours) required to complete the transfer investigational drug form multiplied by the number of forms completed annually. The record

keeping burden represents an average time required for multiple entries (4 minutes or 0.1 hour per entry) on the drug accountability form, the average number of forms maintained by each record keeper and the number of record

keepers. These estimates are based on the items shipped by the PMB and the number of transfer approvals in the calendar year 1999.

Type of respondents	Est. number of respondents	Est. number of responses-respondents	Ave. burden hrs per response	Ave. burden hours	Est. total annual burden hours requested
Drug Transfer Form	1,200	1	0.1	120	120
Drug Accountability Form	4,560	8	0.1	3,648	3,648
Total	5,760				3,768

There are no Capital Costs to report. There are no Operating or Maintenance Costs to report.

Request for Comments

Written comments and/or suggestions from the public and affected agencies are invited on one or more of the following points: (1) Whether the proposed collection of information is necessary for the proposed performance of the functions of the agency, including whether the information shall have practical utility; (2) 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) Ways to enhance the quality, utility, and clarity of the information to be collected; and (4) Ways to minimize the burden of the collection of information on those who are to respond, including the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology.

FOR FURTHER INFORMATION CONTACT: To request more information on the proposed project or to obtain a copy of the data collection plans and instruments, contact Carl Huntley, Head Drug Management and Authorization Section, Pharmaceutical Management Branch, Cancer Therapy Evaluation Program, Division of Cancer Therapy and Diagnosis, National Cancer Institute, Executive Plaza North, Room 7112, 9000 Rockville Pike, Bethesda, Maryland 20892. Or call non-toll-free number 301-496-5725 or e-mail your request, include your address to HuntleyC&ctep.nci.nih.gov.

Comments Due Date: Comments regarding this information collection are best assured of having their full effect if received on or before February 12, 2001.

Dated: December 5, 2000.

Reesa Nichols,
NCI Project Clearance Liaison.
 [FR Doc. 00-31829 Filed 12-13-00; 8:45 am]
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DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Proposed Collection; Comment Request; Tobacco Use Supplement to the 2001-2002 Current Population Survey

SUMMARY: In compliance with the requirement of section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, for public comment on proposed data collection projects, the National Cancer Institute (NCI), the National Institutes of Health (NIH) will publish periodic summaries of proposed projects to be submitted to the Office of Management and Budget (OMB) for review and approval.

Proposed Collection

Title: Tobacco Use Supplement to the 2001-2002 Current Population Survey.

Type of information request: REVISION, OMB No. 0925-0368, Expiration 01/31/2003.

Need and Use of Information Collection: The 2001-2002 Tobacco Use Supplement to the Current Population Survey conducted by the Bureau of the Census will collect data from the civilian non-institutionalized population on tobacco use and smoking prevalence, workplace smoking policies, medical and dental advice to stop smoking, and changes in smoking norms and attitudes. This survey will provide invaluable information to government agencies, other scientists and the general public necessary for tobacco control surveillance and research, as well as measure progress toward tobacco control as part of the National

Cancer Institute's Extraordinary Opportunities in Tobacco Research. This survey is part of a continuing series of surveys that were sponsored by NCI and fielded periodically over the 1990's by the Census Bureau as part of the American Stop Smoking Intervention Study for Cancer Prevention (ASSIST) project and made available for general public use. The Tobacco Use Supplements will be continuing over the next decade alternating between a standard or core tobacco use survey (such as the 2001-2002 survey) and a special topic survey focusing on emerging adult tobacco control issues. The survey will allow state specific estimates to be made. Data will be collected in June 2001, November 2001 and February 2002 from approximately 293,000 respondents. The National Cancer Institute is co-sponsoring this survey with the Centers for Disease Control and Prevention.

Frequency of Response: One-time study.

Affected Public: Individuals or households.

Type of respondents: Persons 15 years of age or older.

The total annual reporting burden is as follows:

Estimated Number of Respondents: 97,666;

Estimated Number of Responses per Respondent: 1;

Average Burden Hours per Response: 0.1169; and

Estimated Total Annual Burden Hours Requested: 11,417.

There are no Capital Costs, Operating Costs, and/or Maintenance Costs to report.

Request for Comments

Written comments and/or suggestions from the public and affected agencies are invited on one or more of the following points: (1) Whether the proposed collection of information is necessary for the proper performance of