instruments must be requested in writing.

SUPPLEMENTARY INFORMATION: Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995 requires: written comments and/or suggestions from the public and affected agencies are invited to address one or more of the following points: (1) Whether the proposed collection of information is necessary for the proper performance of the function of the agency, including whether the information will 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.

Proposed Collection Title: The National Institute of Mental Health Data Archive (NDA), REVISION, OMB Control Number 0925–0667, National Institute of Mental Health (NIMH), National Institutes of Health (NIH).

Need and Use of Information Collection: This REVISION request seeks approval of updates to the previously approved National Database for Autism Research Data Access Request and Data Use Certification documents. The NIMH Data Archive (NDA), formerly known as the National Database for Autism Research (NDAR), is an infrastructure that allows for the submission and storage of human subjects data from researchers conducting studies related to many scientific domains, regardless of the source of funding. The NIH and NIMH developed this resource to allow for the public collection of information from: (1) Individuals who seek permission to access data from the NDA for the purpose of scientific investigation, scholarship or teaching, or other forms of research and research development, via the Data Use Certification (DUC),

and (2) individuals who request permission to submit data to the NDA for the purpose of scientific investigation, scholarship or teaching, or other forms of research and research development, via the Data Submission Agreement (DSA). The extensive information stored in the NDA continues to provide a rare and valuable scientific resource to the field, and plays an integral part in fulfilling research objectives in multiple scientific domains. The NIH and the NIMH seek to encourage use of the NDA by investigators in the field of multiple scientific research domains to achieve rapid scientific progress. In order to take full advantage of this resource and maximize its research value, it is important that data are made broadly available, on appropriate terms and conditions, to the largest possible number of investigators.

OMB approval is requested for 3 years. There are no costs to respondents other than their time. The total estimated annualized burden hours are 1000.

ESTIMATED ANNUALIZED BURDEN HOURS

Form name	Type of respondents	Number of respondents	Number of responses per respondent	Average burden per response (in hours)	Total annual burden hours
NDA Data Submission Agreement (DSA).	Researchers submitting data	250	1	1 hour	250
NDA Data Use Certification (DUC)	Researchers requesting access to data.	750	1	1 hour	750
Total		1000	1000		1000

Melba Rojas,

Project Clearance Liaison, National Institute of Mental Health, National Institutes of Health.

[FR Doc. 2017–14451 Filed 7–10–17; 8:45 am] BILLING CODE 4140–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Institute of Allergy and Infectious Diseases; Notice of Closed Meeting

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is hereby given of the following meeting.

The meeting will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title 5 U.S.C., as amended. The grant applications and

the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: National Institute of Allergy and Infectious Diseases Special Emphasis Panel; T Cell Reagent Resource for the Study of Allergic Diseases (U19).

Date: August 1–2, 2017. Time: 9:00 a.m. to 3:00 p.m. Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, 5601 Fishers Lane, Rockville, MD 20892 (Telephone Conference Call).

Contact Person: Thomas F. Conway, Ph.D., Scientific Review Officer, Scientific Review Program, Division of Extramural Activities, Room 3G51, National Institutes of Health, NIAID, 5601 Fishers Lane, MSC 9823, Bethesda, MD 20892–9823, 240–507–9685, thomas.conway@nih.gov.

(Catalogue of Federal Domestic Assistance Program Nos. 93.855, Allergy, Immunology, and Transplantation Research; 93.856, Microbiology and Infectious Diseases Research, National Institutes of Health, HHS)

Dated: July 5, 2017.

Natasha M. Copeland,

Program Analyst, Office of Federal Advisory Committee Policy.

[FR Doc. 2017-14433 Filed 7-10-17; 8:45 am]

BILLING CODE 4140-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Substance Abuse and Mental Health Services Administration

Agency Information Collection Activities: Submission for OMB Review; Comment Request

Periodically, the Substance Abuse and Mental Health Services Administration (SAMHSA) will publish a summary of information collection requests under OMB review, in compliance with the Paperwork Reduction Act (44 U.S.C. Chapter 35). To request a copy of these documents, call the SAMHSA Reports Clearance Officer on (240) 276–1243.

Project: National Mental Health Study Field Test—NEW

The Substance Abuse and Mental Health Services Administration (SAMHSA) plans to conduct a methodological field test for a potential national mental health study, provisionally named the National Mental Health Study (NMHS). The NMHS will use mental disorder assessments similar to studies last conducted over a decade ago in the National Comorbidity Survey-Replication among adults in 2001–2003 and the National Comorbidity Survey-Adolescent supplement among adolescents in 2001-2002. SAMHSA is collaborating with the National Institute of Mental Health (NIMH) to implement this field test.

The purpose of the NMHS Field Test is to test the procedures for a potential NMHS. The field test consists of three general components. The first component is sample selection using a household screener. The household screener will be used to determine eligibility of individuals and to make selections of individuals to recruit for participation in the second component. The second component consists of an in-person survey of the selected adult and adolescent respondents. The NMHS procedures vary somewhat between adults (aged 18 or older) and adolescents (aged 13 to 17). For all respondents, the in-person assessment (using either the adult or adolescent instrument) will be conducted primarily using audio computer-assisted selfinterviewing (ACASI), with an emphasis on respondents completing the interview in a single session. In addition to the adolescent in-person assessment, parents/legal guardians of adolescent respondents will receive an additional web or phone interviews (the parent instrument). The final component

consists of a telephone clinical reappraisal of a selected subgroup of adult and adolescent respondents, with an additional parent/guardian reporting for adolescents.

The NMHS field test will include 1,200 English speaking respondents— 900 adults and 300 adolescents in the United States excluding Alaska and Hawaii. Approximately 210 parents/ legal guardians of adolescent respondents will complete an additional parent interview. A subsample of approximately 150 adult and adolescent respondents and 50 parent respondents will complete a telephone-based clinical reappraisal follow-up interview. In addition, a subsample of completed screening and interview cases will be recontacted for a brief telephone interview to verify that interviewers followed proper protocols when collecting data. The sample size supports testing of field procedures, sampling algorithms, and data processing steps. The total annual burden estimate is shown in the table helow

ANNUALIZED ESTIMATED BURDEN FOR THE NATIONAL MENTAL HEALTH STUDY FIELD TEST

Instrument	Number of respondents	Responses per respondent	Total number of responses	Hours per response	Total burden hours
Household Screening Interview (including interviews with Adults and Adoles-	2,331	1	2,331	0.083	193
cents)	1,200	1	1,200	1.083	1,300
Parent Interview	210	1	210	0.500	105
Clinical Interview	150	1	150	1.000	150
Clinical Parent Interview	50	1	50	0.500	25
Screening Verification	142	1	142	0.067	10
Interview Verification	180	1	180	0.067	12
Total	4,263		4,263		1,795

Written comments and recommendations concerning the proposed information collection should be sent by August 10, 2017 to the SAMHSA Desk Officer at the Office of Information and Regulatory Affairs, Office of Management and Budget (OMB). To ensure timely receipt of comments, and to avoid potential delays in OMB's receipt and processing of mail sent through the U.S. Postal Service, commenters are encouraged to submit their comments to OMB via email to: OIRA Submission@omb.eop.gov. Although commenters are encouraged to send their comments via email, commenters may also fax their comments to: 202-395-7285. Commenters may also mail them to: Office of Management and Budget, Office of Information and Regulatory

Affairs, New Executive Office Building, Room 10102, Washington, DC 20503.

Summer King,

Statistician.

[FR Doc. 2017–14374 Filed 7–10–17; 8:45 am]

BILLING CODE 4162-20-P

DEPARTMENT OF HOUSING AND URBAN DEVELOPMENT

[Docket No. FR-5997-N-32]

30-Day Notice of Proposed Information Collection: FHA-Insured Mortgage Loan Servicing Involving the Claims and Conveyance Process, Property Inspection/Preservation

AGENCY: Office of the Chief Information

Officer, HUD.

ACTION: Notice.

SUMMARY: HUD is seeking approval from the Office of Management and Budget (OMB) for the information collection described below. In accordance with the Paperwork Reduction Act, HUD is requesting comment from all interested parties on the proposed collection of information. The purpose of this notice is to allow for 30 days of public comment.

DATES: Comments Due Date: August 10, 2017.

ADDRESSES: Interested persons are invited to submit comments regarding this proposal. Comments should refer to the proposal by name and/or OMB Control Number and should be sent to: HUD Desk Officer, Office of Management and Budget, New Executive Office Building, Washington, DC 20503; fax: 202–395–5806, Email: OIRA Submission@omb.eop.gov.