if finalized, would replace the Deemed Final Order in its entirety with new conditions under which nonprescription sunscreen drug products would be determined to be GRASE under section 201(p)(1) of the FD&C Act (21 U.S.C. 321(p)(1)). It would also set forth certain characteristics establishing that a sunscreen drug product is not GRASE under section 201(p)(1) of the FD&C Act.

The original close of the public comment period for this Proposed Order was November 12, 2021. On November 2, 2021, the Agency received a request to extend this comment period by a minimum of 45 days, conveying concern that the original comment period did not provide sufficient time for review of the Proposed Order or for submission of needed updates related to sunscreen active ingredients about which FDA had requested additional data. FDA considered the request and extended the public comment period for the Proposed Order for an additional 45 days, until December 27, 2021.2 This extension will allow additional time for the public to submit information related to these active ingredients (and other proposed sunscreen conditions) that has become available since the closure of the comment period on the 2019 Proposed Rule "Sunscreen Drug Products for Over-the-Counter Human Use" (2019 Proposed Rule).

The Agency reiterates that, as stated in the notice of availability of the Proposed Order published in the Federal Register on September 27, 2021, and in the Proposed Order itself, the Agency will consider all comments that were submitted to the public docket for the 2019 Proposed Rule within its comment period to be constructively submitted as comments on the Proposed Order. The Agency requests that commenters do not resubmit comments on this Proposed Order previously submitted on the 2019 Proposed Rule.

Dated: November 16, 2021.

Lauren K. Roth,

Associate Commissioner for Policy. [FR Doc. 2021–25371 Filed 11–19–21; 8:45 am]

BILLING CODE 4164-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

[Document Identifier: OS-0990-0475]

Agency Information Collection Request. 30-Day Public Comment Request

AGENCY: Office of the Secretary, HHS.

ACTION: Notice.

SUMMARY: In compliance with the requirement of the Paperwork Reduction Act of 1995, the Office of the Secretary (OS), Department of Health and Human Services, is publishing the following summary of a proposed collection for public comment.

DATES: Comments on the ICR must be received on or before December 22, 2021.

ADDRESSES: Submit your comments to *Sherrette.Funn@hhs.gov* or by calling (202) 795–7714.

FOR FURTHER INFORMATION CONTACT:

When submitting comments or requesting information, please include the document identifier 0990–0475, and project title for reference, to Sherrette Funn, the Reports Clearance Officer, Sherrette.funn@hhs.gov, or call 202–795–7714.

SUPPLEMENTARY INFORMATION: Interested persons are invited to send comments regarding this burden estimate or any other aspect of this collection of information, including any of the following subjects: (1) The necessity and utility of the proposed information collection for the proper performance of the agency's functions; (2) the accuracy of the estimated burden; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) the use of automated collection techniques or other forms of information technology to minimize the information collection burden.

Title of the Collection: ASPA COVID— 19 Public Education Campaign Evaluation Surveys.

Type of Collection: Extension. *OMB No.:* 0990–0475.

Abstract: The Office of the Assistant Secretary for Public Affairs (ASPA), U.S. Department of Health and Human Services (HHS) is requesting an extension on a currently approved collection including two components: 1. COVID-19 Attitudes and Beliefs Survey (CABS), and 2. Monthly Outcome Survey (MOS). Throughout execution of the campaign, this information will primarily be used by ASPA to determine whether the campaign is having the intended impact on target audiences' (e.g., parents, young adults, 65+) knowledge, attitudes, and beliefs as they relate to COVID-19, COVID-19 vaccination, and adherence to preventative behaviors. It will also keep key stakeholders informed of the Campaign's progress. Ultimately, the data will inform a thorough evaluation of the efficacy of the campaign and its impact on vaccine uptake.

COVID-19 Attitudes and Beliefs Survey (CABS)

The CABS is a longitudinal survey that will be fielded tri-annually to 4,000 U.S. adults for the duration of the Campaign via NORC at the University of Chicago's AmeriSpeak Panel. The survey will be fielded online, and each fielding period will last between 3 and 6 weeks. Those that respond to wave 1 of the survey will be recontacted in each wave, facilitating a comparison of COVID-19 behavior change over time for a representative sample and evaluation of U.S. adults. Panel members selected to participate in the study will receive one pre-invitation postcard in the mail, one email invitation, and three email reminders to complete the survey in each wave.

Monthly Outcome Survey (MOS)

The MOS is a shorter, cross-sectional survey that will be fielded monthly to 5,000 U.S. adults for the duration of the Campaign via the Ipsos KnowledgePanel 5K Omnibus Survey. The survey will be fielded online, and each fielding period will last between 7 and 10 days.

ANNUALIZED BURDEN HOUR TABLE

	CABS	MOS
Hours to complete survey	0.58	0.17
Participants (per wave)	4,000	5,000
(per year)	3	12
year Total burden hours	12,000	60,000
per year	6,960	10,200

Sum of Both Studies

Total respondents per year: 72,000. Total burden hours per year: 17,160.

Sherrette A. Funn,

Paperwork Reduction Act Reports Clearance Officer, Office of the Secretary.

[FR Doc. 2021–25370 Filed 11–19–21; 8:45 am]

BILLING CODE 4150-25-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Proposed Collection; 60-Day Comment Request; NIH COVID-19 Vaccination Status Form Extension

AGENCY: National Institutes of Health, HHS.

ACTION: Notice.

SUMMARY: In compliance with the requirement of the Paperwork

² See https://www.regulations.gov/document/FDA-1978-N-0018-15828.