

under this provision of the FD&C Act for statutorily regulated products.

Section 2 of Table 1 estimates that 77 respondents (43 cigarette filler and RYO tobacco and 34 smokeless manufacturers, importers, or their agents) will test quantities of HPHCs in an average of 44 products annually. This section addresses the time required for manufacturers and importers (or their agents) who must test HPHC quantities in products. The burden estimates include the burden to test the tobacco products, draft testing reports, and submit the report to FDA. The total expected burden for this section is 481 hours.

Section 3 of Table 1 addresses the time required for manufacturers and importers to test quantities for HPHCs in cigarette smoke. The burden estimates include: the burden to test the number of replicate measurements; test date range; manufacture date range; extraction method; separation method; detection method; and mean quantity and standard deviation of HPHCs and includes the burden to test the tobacco products, draft testing reports, and submit the report to FDA. The annual burden reflects our estimate of the time it takes to test the tobacco products (*i.e.*, carry out laboratory work). The burden estimate assumes that manufacturers and importers report HPHC quantities in cigarette mainstream smoke according to both the ISO and Health Canada smoking regimens. The total expected burden is 17,996 hours for this section.

The total estimated burden for this information collection is expected to be 19,193 hours and 424 annual responses.

Our estimated burden for the information collection reflects an overall increase of 354 annual responses and a corresponding increase of 16,677 hours. We attribute this adjustment to updated methodology in which the current estimates are derived from historical statutory tobacco product applications submitted and authorized by FDA in the past 4 years as: (1) manufacturers and importers (or their agents) of authorized products are required to submit HPHC reports at least 90 days prior to delivery for introduction into interstate commerce for all new products; and (2) initial reporting under section 904(a)(3) of the FD&C Act for statutory products was completed in 2012.

Dated: July 1, 2022.
Lauren K. Roth,
Associate Commissioner for Policy.
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DEPARTMENT OF HEALTH AND HUMAN SERVICES

[Document Identifier: OS–0990–0002]

Agency Information Collection Request. 60-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 September 12, 2022.

ADDRESSES: Submit your comments to *Sherrette.Funn@hhs.gov* or by calling (202) 264–0041.

FOR FURTHER INFORMATION CONTACT: When submitting comments or requesting information, please include the document identifier 0945–0002–60D and project title for reference, to Sherrette A. Funn, email: *Sherrette.Funn@hhs.gov*, or call (202) 264–0041 the Reports Clearance Officer.

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: Civil Rights and Conscience Complaint and Health Information Privacy & Security Complaint

Type of Collection: Office of Civil Rights (OCR)—Extension

OMB No. 0945–0002

Abstract:

ESTIMATED ANNUALIZED BURDEN TABLE

Written forms/electronic forms	Type of respondent	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden hours
Civil Rights/Conscience Discrimination Complaint. Health Information Privacy Complaint.	Individuals or households, Not-for-profit institutions.	15,446	1	45/60	11,585
	Individuals or households, Not-for-profit institutions.	30,392	1	45/60	22,794
Total	45,838	45/60	34,379

Sherrette A. Funn,
Paperwork Reduction Act Reports Clearance Officer, Office of the Secretary.
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DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Institute on Deafness and Other Communication Disorders; Notice of Meeting

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended, notice is hereby given of a meeting of the National Deafness and

Other Communication Disorders Advisory Council.
The meeting will be open to the public as indicated below, with attendance limited to space available. Individuals who plan to attend and need special assistance, such as sign language interpretation or other reasonable accommodations, should notify the contact person listed below in advance of the meeting.
The meeting will be closed to the public in accordance with the