

Dated: April 24, 2025.

Grace R. Graham,

*Deputy Commissioner for Policy, Legislation,
and International Affairs.*

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2024–N–4754]

Agency Information Collection Activities; Submission for Office of Management and Budget Review; Comment Request; Financial Disclosure by Clinical Investigators

AGENCY: Food and Drug Administration,
HHS.

ACTION: Notice.

SUMMARY: The Food and Drug
Administration (FDA) is announcing
that a proposed collection of

information has been submitted to the
Office of Management and Budget
(OMB) for review and clearance under
the Paperwork Reduction Act of 1995.

DATES: Submit written comments
(including recommendations) on the
collection of information by June 2,
2025.

ADDRESSES: To ensure that comments on
the information collection are received,
OMB recommends that written
comments be submitted to [https://
www.reginfo.gov/public/do/PRAMain](https://www.reginfo.gov/public/do/PRAMain).
Find this particular information
collection by selecting “Currently under
Review—Open for Public Comments” or
by using the search function. The OMB
control number for this information
collection is 0910–0396. Also include
the FDA docket number found in
brackets in the heading of this
document.

FOR FURTHER INFORMATION CONTACT:
Amber Sanford, Office of Operations,
Food and Drug Administration, Three
White Flint North, 10A–12M, 11601

Landsdown St., North Bethesda, MD
20852, 301–796–8867, [PRAStaff@
fda.hhs.gov](mailto:PRAStaff@fda.hhs.gov).

SUPPLEMENTARY INFORMATION: In
compliance with 44 U.S.C. 3507, FDA
has submitted the following proposed
collection of information to OMB for
review and clearance.

Financial Disclosure by Clinical Investigators

*OMB Control Number 0910–0396—
Extension*

Respondents to this collection are
sponsors of marketing applications that
contain clinical data from studies
covered by the regulations. These
sponsors represent pharmaceutical,
biologic, and medical device firms.
Respondents are also clinical
investigators who provide financial
information to the sponsors of
marketing applications.

Table 1 shows information that is the
basis of the estimated number of
respondents in tables 2 through 4.

TABLE 1—ESTIMATED NUMBER OF APPLICATIONS, CLINICAL TRIALS, AND INVESTIGATORS SUBJECT TO THE REGULATION
BY TYPE OF APPLICATION ¹

Application type	Total number of applications	Number of applications affected	Number of trials	Number of investigators
Drugs:				
New drug application (NDA), new molecular entity (NME)	35	35	3 to 10	3 to 100.
NDA non-NME	94	44	3 to 10	3 to 100.
NDA efficacy supplement	171	100	1 to 3	10 to 30.
Abbreviated new drug application (ANDA)	685	1	1.1	2.
ANDA supplement	10,366	1	1	2.
CBER Biologics:				
Biologics license application (BLA)	26	26	3 to 10	3 to 100.
BLA efficacy supplement	26	26	1 to 3	10 to 30.
CDER Biologics:				
BLAs	19	19	3 to 10	3 to 100.
BLA efficacy supplements	64	50	1 to 3	10 to 30.
Medical Devices:				
Premarket approval (PMA)	43	50	1 to 31	10 to 20.
PMA supplement	28	30	to 3	3 to 10
Reclassification devices	0	0	0	0.
510(k)	3,401	254	1	3 to 10.
De Novo requests	84	76	1 to 3	10 to 20.

¹ Source: Agency estimates.

FDA estimates the burden of this
collection of information as follows:

Reporting Burden

Under § 54.4(a) (21 CFR 54.4(a)),
applicants submitting an application
that relies on clinical studies must
submit a complete list of clinical
investigators who participated in a
covered clinical study, and must either
certify to the absence of certain financial
arrangements with clinical investigators
(Form FDA 3454) or, under § 54.4(a)(3),
disclose to FDA the nature of those

arrangements and the steps taken by the
applicant or sponsor to minimize the
potential for bias (Form FDA 3455).

FDA estimates that almost all
applicants submit a certification
statement under § 54.4(a)(1) and (2).
Preparation of the statement using Form
FDA 3454 should require no more than
1 hour per study. The number of
respondents is based on the estimated
number of affected applications.

When certification is not possible and
disclosure is made using Form FDA
3455, the applicant must describe,

under § 54.4(a)(3), the financial
arrangements or interests and the steps
that were taken to minimize the
potential for bias in the affected study.
As the applicant would be fully aware
of those arrangements and the steps
taken to address them, describing them
will be straightforward. The Agency
estimates that it will take about 5 hours
to prepare this narrative. Based on our
experience with this collection, FDA
estimates that approximately 10 percent
of the respondents with affected

applications will submit disclosure statements.
In the **Federal Register** of November 29, 2024 (89 FR 94735), FDA published

a 60-day notice requesting public comment on the proposed collection of information. One comment was received but was not PRA related.

FDA estimates the burden of this collection of information as follows:

TABLE 2—ESTIMATED ANNUAL REPORTING BURDEN ¹

21 CFR section	Number of respondents	Number of responses per respondent	Total annual responses	Average burden per response	Total hours
Certification—54.4(a)(1) and (2)—Form FDA 3454	712	1	712	1	712
Disclosure—54.4(a)(3)—Form FDA 3455	71	1	71	5	355
Total					1,067

¹ There are no capital costs or operating and maintenance costs associated with this collection of information.

Recordkeeping Burden

Under § 54.6 (21 CFR 54.6), the sponsors of covered studies must maintain complete records of compensation agreements with any compensation paid to nonemployee

clinical investigators, including information showing any financial interests held by the clinical investigator, for 2 years after the date of approval of the applications. Sponsors of covered studies maintain many records regarding clinical investigators,

including protocol agreements and investigator resumes or curriculum vitae. FDA estimates an average of 15 minutes will be required for each recordkeeper to add this record to the clinical investigators' file.

TABLE 3—ESTIMATED ANNUAL RECORDKEEPING BURDEN ¹

21 CFR section	Number of recordkeepers	Number of records per recordkeeper	Total annual records	Average burden per recordkeeping	Total hours ²
Recordkeeping—54.6	712	1	712	0.25	178

¹ There are no capital costs or operating and maintenance costs associated with this collection of information.

² Numbers have been rounded.

Third-Party Disclosure Burden

Under § 54.4(b), clinical investigators supply to the sponsor of a covered study financial information sufficient to allow the sponsor to submit complete and accurate certification or disclosure statements. Clinical investigators are accustomed to supplying such information when applying for research

grants. Also, most people know the financial holdings of their immediate family and records of such interests are generally accessible because they are needed for preparing tax records. For these reasons, FDA estimates that the time required for this task may range from 5 to 15 minutes; we used the median, 10 minutes, for the average burden per disclosure (see table 4). To

estimate the number of respondents for each FDA Center, we took the median number of investigators for each application type, multiplied each median number of investigators by the number of affected applications for that application type, then summed those products to get the total number of respondents for the Center.

TABLE 4—ESTIMATED ANNUAL THIRD-PARTY DISCLOSURE BURDEN ¹

21 CFR section	Number of respondents	Number of disclosures per respondent	Total annual disclosures	Average burden per disclosure	Total hours ²
54.4(b)—Clinical Investigators	13,646	1	13,646	0.17	2,320

¹ There are no capital costs or operating and maintenance costs associated with this collection of information.

² Numbers have been rounded.

The burden for this information collection request has changed since the last OMB approval. We have adjusted our estimated burden for the information collection to reflect the

number of submissions we received in the last few years. These adjustments result in an increase of 557 total annual responses and a corresponding increase of 87 total hours.

Dated: April 24, 2025.
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