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Dated: May 27, 2016.

Leslie Kux,

Associate Commissioner for Policy.

[FR Doc. 2016–12950 Filed 6–1–16; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2013–D–1543]

Agency Information Collection Activities; Submission for Office of Management and Budget Review; Comment Request; Guidance for Industry on Nonproprietary Naming of Biological Products

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: Fax written comments on the collection of information by July 5, 2016.

ADDRESSES: To ensure that comments on the information collection are received, OMB recommends that written comments be faxed to the Office of Information and Regulatory Affairs, OMB, Attn: FDA Desk Officer, FAX: 202–395–7285, or emailed to oir_submission@omb.eop.gov. All comments should be identified with the OMB control number 0910–NEW and title “Nonproprietary Naming of

Biological Products.” Also include the FDA docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT: FDA PRA Staff, Office of Operations, Food and Drug Administration, 8455 Colesville Rd., COLE–14526, Silver Spring, MD 20993–0002, 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.

Guidance for Industry on Nonproprietary Naming of Biological Products OMB Control Number 0910–NEW

The guidance entitled “Guidance for Industry on Nonproprietary Naming of Biological Products” describes FDA’s current thinking on the need for biological products licensed under the Public Health Service Act (PHS Act) to bear a nonproprietary name that includes an FDA-designated suffix. There is a need to clearly identify biological products to facilitate pharmacovigilance and, for the purposes of safe use, to minimize inadvertent substitution. Accordingly, for biological products licensed under the PHS Act, FDA intends to designate a nonproprietary name that includes a core name and a distinguishing suffix. This naming convention is applicable to biological products previously licensed and newly licensed under section 351(a) or 351(k) of the PHS Act (42 U.S.C. 262(a) or 262(k)).

The guidance includes information collection by requesting that applicants propose a suffix composed of four lowercase letters for use as the distinguishing identifier included in the proper name designated by FDA at the time of licensure for biological products licensed under the PHS Act. The suffix will be incorporated in the nonproprietary name of the product. The guidance recommends that applicants should submit up to 10 proposed suffixes, in the order of the applicant’s preference. We also recommend including supporting analyses demonstrating that the proposed suffixes meet the factors described in the guidance for FDA’s consideration.

As indicated in table 1, we estimate that we will receive a total of 40 requests annually for the proposed proper name for biological products submitted under section 351(a) of the PHS Act, and 6 requests annually for the proposed proper name for biosimilar products and interchangeable products

submitted under section 351(k) of the PHS Act. The average burden per response (hours) is based on our experience with similar information collection requirements for applicants to create and submit suffix proposals to FDA and in consideration of comments received in response to our 60-day notice.

In the **Federal Register** of August 28, 2015 (80 FR 52296), we published a 60-day notice requesting public comment on the proposed collection of information. Most comments supported our proposal to designate a suffix. Many comments suggested that a meaningful, distinguishable suffix may help to improve pharmacovigilance, enhance safety, and facilitate identification between biological products. Some comments supported use of a random suffix to avoid creating an unfair advantage for specific manufacturers. Several comments stated that the current practices of FDA and non-FDA entities for identifying biosimilar and interchangeable products is sufficient for the purpose of pharmacovigilance, and designation of a suffix is not needed. One comment stated that FDA’s estimate of 6 hours to submit proposed suffixes is based only on the time needed to prepare the submission itself after the multiple suffixes have been selected. The comment further stated that because FDA suggests that each respondent submit three suggested suffixes for consideration, the time needed to do an analysis of each suffix would exceed 720 hours per suffix (based on their own company experience) or 2,160 hours total for the three suffixes.

In response to the comments we note that our estimated annual reporting burden results from information that would be submitted to us by applicants in order to facilitate Agency designation of a suffix as part of the proper name of a biological product. We estimated that sponsors would spend 2 hours completing the submission for each of the three suffixes, resulting in 6 hours as the average burden. This estimate is an annualized figure based on the average number of responses per respondent and the average burden per response over a 3-year period. We understand that there is a certain amount of research and other costs that an applicant might encounter in analyzing any proposed name for a biological product. We also recognize that the burden may be higher for some applicants and lower for other applicants based on a variety of factors specific to the applicant.

The comment suggesting that it will take 720 hours to complete an analysis

and submission for each suffix does not provide a basis by which the estimate was calculated or whether it is broadly applicable. We find this figure rather high and retain our original estimate of 6 hours. The latter figure is based on our familiarity with the average amount of time required by similar submissions to

FDA. At the same time, the comment also suggested that we failed to adequately account for the time spent on creating proposed suffixes. In consideration, therefore, we have revised our estimate upward to account for burden associated with creating and

submitting up to 10 proposed suffixes for designation, as reflected in table 1. FDA estimates the information collection burden as follows:
Description of Respondents: Respondents to the collection are sponsors of biological product applications.

TABLE 1—ESTIMATED ANNUAL REPORTING BURDEN ¹

Activity	Number of respondents	Number of responses per respondent	Total annual responses	Average burden per response	Total hours
Information for the Proposed Proper Name for Biological Products Submitted Under Section 351(a) of the PHS Act	20	2	40	420	16,800
Information for the Proposed Proper Name for Biosimilar Products and Interchangeable Products Submitted Under Section 351(k) of the PHS Act	3	2	6	420	2,520
Total	19,320

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

The guidance also refers to previously approved collections of information found in FDA regulations. The collections of information for the submission of a biologics license application (BLA) and changes (supplements) to an approved application under 21 CFR part 601 have been approved under OMB control number 0910–0338. The collections of information for the submission of a BLA under section 351(k) of the PHS Act (biosimilar products and interchangeable products) have been approved under OMB control number 0910–0719.

Dated: May 26, 2016.

Leslie Kux,

Associate Commissioner for Policy.

[FR Doc. 2016–12885 Filed 6–1–16; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Health Resources and Services Administration

National Advisory Council on the National Health Service Corps; Notice of Meeting

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92–463), notice is hereby given of the following meeting:

Name: National Advisory Council on the National Health Service Corps (NACNHSC).

Dates and Times: June 22, 2016 11:00 a.m.–5:00 p.m. EST.

Place: U.S. Department of Health and Human Services, Health Resources and

Services Administration, Conference Room #5E29, 5600 Fishers Lane, Rockville, Maryland 20857, Webinar and Conference Call Format.

Status: This advisory council meeting will be open to the public.

Purpose: The NACNHSC provides advice and recommendations to the Secretary of the U.S. Department of Health and Human Services (HHS) and, by designation, the Administrator of the Health Resources and Services Administration (HRSA) on a range of issues including identifying the priorities for the National Health Service Corps (NHSC) and policy revisions.

Agenda: The NACNHSC will refine the Council’s top priorities for the NHSC and will continue to work on the draft of formal recommendations to submit to the HHS Secretary and the HRSA Administrator. During the March 2016 NACNHSC meeting, the Council identified its priorities for the NHSC. The Council will continue the discussion on how the priorities meet the goals of the 2016 Bureau of Health Workforce Strategic Plan. The priority areas include, but are not limited to, telehealth, Medication Assisted Treatment (MAT) certification, mentorship and NHSC discipline expansion, specifically for mental and behavioral and oral health providers. The content of the agenda is subject to change prior to the meeting. The NACNHSC final agenda and call-in information will be available 3 days in advance of the meeting at <http://nhsc.hrsa.gov/corpsexperience/aboutus/nationaladvisorycouncil/meetingsummaries/index.html>.

FOR FURTHER INFORMATION CONTACT:

Please send requests for information to Dawn Smith, Bureau of Health Workforce, HRSA, in one of two ways: (1) Send a request to the following address: Dawn Smith, Bureau of Health Workforce, Health Resources and Services Administration, Room 14N70B, 5600 Fishers Lane, Rockville, Maryland 20857; or (2) send an email to dsmith3@hrsa.gov.

SUPPLEMENTARY INFORMATION: Further information regarding the NACNHSC, including the roster of members and past meeting summaries, is available at: <http://nhsc.hrsa.gov/corpsexperience/aboutus/nationaladvisorycouncil/meetingsummaries/index.html>.

Members of the public and interested parties may request to participate in the meeting by contacting Dawn Smith via email at dsmith3@hrsa.gov to obtain access information. Access will be granted on a first come, first served basis. Individuals who plan to attend and need special assistance, such as sign language interpretation or other reasonable accommodations, should notify the contact person listed above at least 10 days prior to the meeting. Public participants may submit written statements in advance of the scheduled meeting. If you would like to provide oral public comment during the meeting, please register with Dawn Smith. Public comment will be limited to 3 minutes per speaker. Statements and comments can be addressed to Dawn Smith by emailing her at dsmith@hrsa.gov.

In addition, please be advised that committee members are given copies of all written statements submitted from the public. Any further public