DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2014-D-1351]

Flow Cytometric Devices; Draft Guidance for Industry and Food and Drug Administration Staff; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the availability of the draft guidance entitled "Flow Cytometric Devices." This draft guidance addresses the current major review concerns regarding submissions for flow cytometric devices used as in vitro diagnostic devices for leukocyte immunophenotyping and provides suggestions on the content of submissions for these types of devices. This draft guidance is not final nor is it in effect at this time.

DATES: Although you can comment on any guidance at any time (see 21 CFR 10.115(g)(5)), to ensure that the Agency considers your comment of this draft guidance before it begins work on the final version of the guidance, submit either electronic or written comments on the draft guidance by [INSERT DATE 90 DAYS AFTER DATE OF PUBLICATION IN THE FEDERAL REGISTER].

ADDRESSES: An electronic copy of the guidance document is available for download from the Internet. See the SUPPLEMENTARY INFORMATION section for information on electronic access to the guidance. Submit written requests for a single hard copy of the draft guidance document entitled "Flow Cytometric Devices" to the Office of the Center Director, Guidance and
I. Background

This draft guidance addresses certain issues that arise in premarket submissions for flow cytometric devices used as in vitro diagnostic devices for leukocyte immunophenotyping and provides suggestions on the content of submissions for these types of devices. It is intended to be used in conjunction with the other cited guidance documents referenced therein. In preparing your submission to FDA, we recommend that you contact FDA's Office of In Vitro Diagnostics
and Radiological Health (see FOR FURTHER INFORMATION CONTACT) for additional information regarding your submission. This draft guidance focuses on issues relevant to flow cytometric devices with an expanded scope of review topics that reflect the recognition of a flow cytometric device as an analytical system, which includes processing reagents, processing instrumentation, flow cytometers, and analytical software, in addition to the monoclonal antibody (mAb) component. The information presented in this draft guidance is based on the following: (1) Current basic science, (2) clinical experience, and (3) previous submissions by manufacturers to FDA. As advances are made in science and medicine, the content of this guidance will be re-evaluated and revised as necessary to accommodate new knowledge.

This draft guidance is directed toward immunophenotyping of leukocytes using mAbs. However, the concepts may be applicable to related devices that utilize fluorochromes or fluorogenic substrates to measure ligand binding on solid particles in suspension, with or without mAbs. This draft guidance does not cover microscopy devices utilizing fluorescent or chromogenic enzyme-substrate detection methods (e.g., immunohistochemical stains) nor does it cover the use of flow cytometry for cell enrichment and cell sorting/purification when used in cell therapy product manufacturing.

II. Significance of Guidance

This draft guidance is being issued consistent with FDA's good guidance practices regulation (21 CFR 10.115). The draft guidance, when finalized, will represent the Agency's current thinking on flow cytometric devices. It does not create or confer any rights for or on any person and does not operate to bind FDA or the public. An alternative approach may be used if such approach satisfies the requirements of the applicable statute and regulations.
III. Electronic Access

Persons interested in obtaining a copy of the draft guidance may do so by downloading an electronic copy from the Internet. A search capability for all CDRH guidance documents is available at


IV. Paperwork Reduction Act of 1995

This draft guidance refers to previously approved collections of information found in FDA regulations. These collections of information are subject to review by the Office of Management and Budget (OMB) under the Paperwork Reduction Act of 1995 (44 U.S.C. 3501-3520). The collections of information in 21 CFR part 807, subpart E, have been approved under OMB control number 0910-0120, and the collections of information in 21 CFR part 801 and 21 CFR 809.10 have been approved under OMB control number 0910-0485.

V. Comments

Interested persons may submit either electronic comments regarding this document to http://www.regulations.gov or written comments to the Division of Dockets Management (see ADDRESSES). It is only necessary to send one set of comments. Identify comments with the docket number found in brackets in the heading of this document. Received comments may be
seen in the Division of Dockets Management between 9 a.m. and 4 p.m., Monday through Friday, and will be posted to the docket at http://www.regulations.gov.

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Leslie Kux,
Assistant Commissioner for Policy.

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