

**FDA Staff Manual Guides, Volume I - Organizations and Functions**

**Department of Health and Human Services**

**Food and Drug Administration**

**Center for Drug Evaluation and Research**

**Office of New Drugs**

**Office of Oncologic Diseases**

**Division of Oncology I**

Effective Date: September 25, 2019

**1. Division of Oncology I (DCDGFA).**

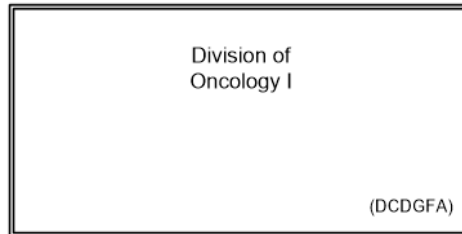
- A. Reviews Investigational New Drug applications and requests for claimed investigational exemption regulated by this Division and decides on appropriate action, including approval or disapproval of research plans and protocols, modifications, and restrictions. Develops policies and procedures pertinent to particular aspects of drug and biologics investigations.
- B. Evaluates New Drug Applications (NDAs) and Biological License Applications (BLAs) for safety and effectiveness and formulates decisions or recommendations regarding approvability in accord with applicable delegations of authority. Also evaluates supplements that propose changes in the conditions upon which NDA/BLA approvals are based. Develops policies and procedures applicable to the review and evaluation of drugs and biologics related to oncologic diseases.
- C. Evaluates adequacy of information in proposed labeling for products related to oncologic diseases.
- D. Evaluates and takes appropriate action on recommendations concerning withdrawal of approval of NDAs and BLAs for drugs and biologics related to oncologic diseases.
- E. Develops, in coordination with other Food and Drug Administration (FDA) components, guidance for staff, sponsor and the public that describes the FDA's interpretation of or policy on regulatory issues. Participates in FDA sponsored consumer and professional educational programs on drug standards.

F. Performs medical and scientific evaluations of submissions on generic drugs, drugs under monograph, and over-the-counter drug products regulated by other offices in the Center, as applicable.

**2. Authority and Effective Date.**

The functional statements for Division of Oncology I were approved by the Secretary of the Health and Human Services on September 25, 2019.

**Department of Health and Human Services  
Food and Drug Administration  
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The following is the Department of Health and Human Services, Food and Drug Administration, Center for Drug Evaluation and Research, Office of New Drugs, Office of Oncologic Diseases, Division of Oncology I organization structure depicting all the organizational structures reporting to the Director.

Division of Oncologic I (DCDGFA).