**Appropriations Language**

**Salaries and Expenses**
For necessary expenses of the Food and Drug Administration, including hire and purchase of passenger motor vehicles; for payment of space rental and related costs pursuant to Public Law 92–313 for programs and activities of the Food and Drug Administration which are included in this Act; for rental of special purpose space in the District of Columbia or elsewhere; for miscellaneous and emergency expenses of enforcement activities, authorized and approved by the Secretary and to be accounted for solely on the Secretary’s certificate, not to exceed $25,000; and notwithstanding section 521 of Public Law 107–188; [$4,681,392,000 \[^1\] $2,527,960,000: Provided, That of the amount provided under this heading, [$851,481,000 shall be derived from prescription drug user fees authorized by 21 U.S.C. 379h, and shall be credited to this account and remain available until expended; $137,677,000 shall be derived from medical device user fees authorized by 21 U.S.C. 379j, and shall be credited to this account and remain available until expended; $318,363,000 shall be derived from human generic drug user fees authorized by 21 U.S.C. 379j-42, and shall be credited to this account and remain available until expended; $21,540,000 shall be derived from biosimilar biological product user fees authorized by 21 U.S.C. 379j-52, and shall be credited to this account and remain available until expended; $22,818,000]\[^1\]$24,142,000 shall be derived from animal drug user fees authorized by 21 U.S.C. 379j-12, and shall be credited to this account and remain available until expended; [$9,705,000]\[^1\]$12,100,000 shall be derived from animal generic drug user fees authorized by 21 U.S.C. 379j-21, and shall be credited to this account and remain available until expended; [$599,000,000]\[^1\]$672,000,000 shall be derived from tobacco product user fees authorized by 21 U.S.C. 387s, and shall be credited to this account and remain available until expended: Provided further, That in addition to and notwithstanding any other provision under this heading, amounts collected for [prescription drug user fees, medical device user fees, human generic drug user fees, biosimilar biological product user fees, animal drug user fees, and animal generic drug user fees that exceed the respective fiscal year [2016][2018 limitations are appropriated and shall be credited to this account and remain available until expended: Provided further, That fees derived from [prescription drug, medical device, human generic drug, biosimilar biological product, animal drug[,] and animal generic drug assessments for fiscal year [2016][2018, including any such fees collected prior to fiscal year [2016][2018 but credited for fiscal year [2016][2018, shall be subject to the fiscal year [2016][2018 limitations: Provided further, That the Secretary may accept payment during fiscal year [2016][2018 of user fees specified under this heading and authorized for fiscal year [2017][2019, prior to the due date for such fees, and that amounts of such fees assessed for fiscal year [2017][2019 for which the Secretary accepts payment in fiscal year [2016][2018 shall not be included in amounts under this heading: Provided further, That none of these funds shall be used to develop, establish, or operate any program of user fees authorized by 31 U.S.C. 9701: Provided further, That of the total amount appropriated: (1) $987,328,000 shall be for the Center for Food Safety and Applied Nutrition and related field activities in the Office of Regulatory Affairs; (2) $1,394,136,000 shall be for the Center for Drug Evaluation and Research and related field activities in the Office of Regulatory Affairs; (3) $354,901,000 shall be for the Center for Biologics Evaluation and Research and for related field

[^1]: Please note that brackets indicate deleted text and italics indicate new text.
activities in the Office of Regulatory Affairs; (4) $187,825,000 shall be for the Center for Veterinary Medicine and for related field activities in the Office of Regulatory Affairs; (5) $430,443,000 shall be for the Center for Devices and Radiological Health and for related field activities in the Office of Regulatory Affairs; (6) $63,331,000 shall be for the National Center for Toxicological Research; (7) $564,117,000 shall be for the Center for Tobacco Products and for related field activities in the Office of Regulatory Affairs; (8) not to exceed $171,418,000 shall be for Rent and Related activities, of which $52,346,000 is for White Oak Consolidation, other than the amounts paid to the General Services Administration for rent; (9) not to exceed $238,274,000 shall be for payments to the General Services Administration for rent; and (10) $289,619,000 shall be for other activities, including the Office of the Commissioner of Food and Drugs, the Office of Foods and Veterinary Medicine, the Office of Medical and Tobacco Products, the Office of Global and Regulatory Policy, the Office of Operations, the Office of the Chief Scientist, and central services for these offices:]

[Provided further, That not to exceed $25,000 of this amount shall be for official reception and representation expenses, not otherwise provided for, as determined by the Commissioner: [Provided further, That any transfer of funds pursuant to section 770(n) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 379dd(n)) shall only be from amounts made available under this heading for other activities: Provided further, That of the amounts that are made available under this heading for "other activities", and that are not derived from user fees, $1,500,000 shall be transferred to and merged with the appropriation for "Department of Health and Human Services—Office of Inspector General" for oversight of the programs and operations of the Food and Drug Administration and shall be in addition to funds otherwise made available for oversight of the Food and Drug Administration:]

[Provided further, That funds may be transferred from one specified activity to another with the prior [approval] notification of the Committees on Appropriations of both Houses of Congress.


Buildings and Facilities
For plans, construction, repair, improvement, extension, alteration, demolition, and purchase of fixed equipment or facilities of or used by the Food and Drug Administration, where not otherwise provided, $11,788,000, to remain available until expended.

Salaries and Expenses (Legislative Proposal)
In addition, contingent upon the enactment of authorizing legislation, the Secretary shall charge a fee for prescription drug review activities, medical device review activities, biosimilar biological products review activities, and human generic drugs review activities: Provided, That fees of $1,262,182,000, for prescription drug reviews, shall be credited to this account and remain
available until expended; $439,001,000 for medical device reviews, shall be credited to this account and remain available until expended; $615,746,000 for human generic drug reviews, shall be credited to this account and remain available until expended; and $86,736,000 for biosimilar biological product reviews, shall be credited to this account and remain available until expended: Provided further, That, in addition and notwithstanding any other provision under this heading, amounts collected for prescription drug user fees, medical device user fees, biosimilar biological product user fees, and human generic drug user fees that exceed the respective fiscal year 2018 limitations are appropriated and shall be credited to this account and remain available until expended: Provided further, That fees derived from prescription drug reviews, medical device reviews, biosimilar biological products reviews, and human generic drugs reviews for fiscal year 2018 received during fiscal year 2018, including any such fees assessed prior to fiscal year 2018 but credited for fiscal year 2018, shall be subject to the fiscal year 2018 limitations: Provided further, That the Secretary may accept payment during fiscal year 2018 of user fees specified in this paragraph and authorized for fiscal year 2019, prior to the due date for such fees, and that amounts of such fees assessed for fiscal year 2019 for which the Secretary accepts payment in fiscal year 2018 shall not be included in amounts in this paragraph.

In addition, contingent upon the enactment of authorizing legislation, the Secretary shall increase the fees for animal drug review activities and animal generic drug review activities: Provided, That additional fees of $46,110,000, for animal drug reviews, shall be credited to this account and remain available until expended; and $6,375,000, for animal generic drug reviews, shall be credited to this account and remain available until expended.
FY 2018 PROPOSED GENERAL PROVISIONS

SEC. __. INCREASE IN EXPORT CERTIFICATION FEES.—

Section 801(c)(4) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 381(c)(4)) is amended—

(1) in subparagraph (B) by striking “but shall not exceed $175 for each certification” and inserting “in an amount specified in subparagraph (c)”; and

(2) by adding at the end the following new subparagraphs:

“(c) The fee for each written export certification issued by the Secretary under this paragraph shall not exceed—

“(i) $600 for fiscal year 2018; and

“(ii) for each subsequent fiscal year, the prior fiscal year maximum amount multiplied by the inflation adjustment under section 738(c)(2)(c), applied without regard to the limitation in clause (ii)(II) of such section.

“(F) The Secretary shall, for each fiscal year, publish in the Federal Register a notice of the export certification fee under this paragraph for such year, not later than 60 days before such fee takes effect.”

"SEC. __. FDA WORKING CAPITAL FUND.

(a) There is hereby established in the Treasury of the United States a Working Capital Fund (the Fund) to be administered by the Food and Drug Administration (FDA), without fiscal year limitation, for the payment of salaries, travel, and other expenses necessary to the maintenance and operation of (1) a supply service for the purchase, storage, handling, issuance, packing, or shipping of stationery, supplies, materials, equipment, and blank forms, for which stocks may be maintained to meet, in whole or in part, the needs of the FDA and requisitions of other Government Offices, and (2) such other services as the Commissioner of the FDA, subject to review by the Secretary of Health and Human Services, determines may be performed more advantageously as central services. The Fund shall be reimbursed from applicable discretionary resources, notwithstanding any otherwise applicable purpose limitations, available when services are performed or stock furnished, or in advance, on a basis of rates which shall include estimated or actual charges for personal services, materials, equipment, information technology, and other expenses. Charges for equipment and information technology shall include costs associated with maintenance, repair, and depreciation (including improvement and replacement).

(b) Of any discretionary resources appropriated in this Act for fiscal year 2018 for “Department of Health and Human Services, Food and Drug Administration, Salaries and Expenses”, not to exceed $5,000,000 of amounts available as of September 30 may be transferred to and merged with the Fund established under subsection (a), notwithstanding any otherwise applicable purpose limitations.

(c) No amounts may be transferred pursuant to this section that are designated by the Congress as an emergency requirement pursuant to a concurrent resolution on the budget or the Balanced Budget and Emergency Deficit Control Act of 1985.
SEC. __. 21ST CENTURY CURES.—

Sec. XXX. For necessary expenses to carry out the purposes described under section 1002(b)(4) of the 21st Century Cures Act, in addition to amounts available for such purposes under the heading "Salaries and Expenses", $60,000,000, to remain available until expended: Provided, That amounts appropriated in this paragraph are appropriated pursuant to section 1002(b)(3) of the 21st Century Cures Act, are to be derived from amounts transferred under section 1002(b)(2)(A) of such Act, and may be transferred by the Secretary of Health and Human Services to other accounts of the Department solely for the purposes provided in such Act: Provided further, That such transfer authority is in addition to any other transfer authority provided by law.