Five-Year Financial Plan
Fiscal Years

2023-2024-2025-2026-2027

FY 2023 Version

FOR THE

Generic Drug User Fee Amendments Program

FOOD AND DRUG ADMINISTRATION
DEPARTMENT OF HEALTH AND HUMAN SERVICES
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Five-Year Plan Overview

A. Scope

The purpose of the Five-Year Financial Plan is to communicate the anticipated financial position of the Generic Drug User Fee Amendments (GDUFA) program over the current five-year authorization period (GDUFA III). This document addresses the plan for implementation and use of generic drug user fees by the Food and Drug Administration (FDA or the Agency) during the period of October 1, 2022, through September 30, 2027.

B. Five-Year Plan Commitments

In accordance with the GDUFA Reauthorization Performance Goals and Program Enhancements Fiscal Years 2023 Through 2027 (GDUFA III Commitment Letter), section VIII.D.2, FDA will publish a GDUFA five-year financial plan no later than the second quarter of fiscal year (FY) 2023. FDA will publish updates to the five-year financial plan no later than the second quarter of each subsequent fiscal year. The purpose of this document is to meet these commitments.

C. Updates to the Five-Year Plan

All estimates in the plan are subject to review and reassessment each fiscal year as the actual amounts for appropriations, obligations, and collections for the previous year become available. The five-year financial plan provides the baseline from which future changes will be made. Updates to the five-year financial plan will occur on an annual basis and cover the five years in the current reauthorization period.

Management Discussion

D. Organization Background

FDA is responsible for protecting public health by helping to ensure the safety, efficacy, and security of human and veterinary drugs, biological products, and medical devices and by ensuring the safety of our nation’s food supply, cosmetics, and products that emit radiation. FDA also has responsibility for regulating the manufacturing, marketing, and distribution of tobacco products and advancing the public’s health by helping to speed innovations that make medical products more effective, safe, and affordable and by helping the public get accurate, science-based information needed to use medical products and consume foods to maintain and improve their health. FDA similarly plays a significant role in the nation’s counterterrorism capability. FDA fulfills this responsibility by ensuring the security of the food supply and by fostering the development of medical products to respond to deliberate and naturally emerging public health threats.
Program Organization

There are four major FDA components that support the GDUFA program: the Center for Drug Evaluation and Research (CDER), the Center for Biologics Evaluation and Research (CBER), the Office of Regulatory Affairs (ORA), and Headquarters (HQ).

Exhibit 1 provides an overview of the mission for each of these components.

**Exhibit 1: User Fee Program Components**

<table>
<thead>
<tr>
<th>Component</th>
<th>Mission</th>
</tr>
</thead>
<tbody>
<tr>
<td>CDER</td>
<td>Protects and promotes public health by helping to ensure that human drugs are safe and effective, meet established quality standards, and are available to patients.</td>
</tr>
<tr>
<td>CBER</td>
<td>Ensures the safety, purity, potency, and effectiveness of biological products including vaccines, allergens, blood and blood products, and cells, tissues, and gene therapies for the prevention, diagnosis, and treatment of human diseases, conditions, or injury.</td>
</tr>
<tr>
<td>ORA</td>
<td>Protects consumers and enhances public health by maximizing compliance of FDA-regulated products and by minimizing the risk(s) associated with those products. ORA inspects regulated products and manufacturers, conducts sample analyses of regulated products and reviews imported products offered for entry into the United States.</td>
</tr>
<tr>
<td>HQ</td>
<td>Provides FDA-wide program direction and administrative services to ensure FDA’s consumer and patient safety programs are effectively and efficiently managed.</td>
</tr>
</tbody>
</table>

**User Fee Governance**

The Agency’s expanding level of user fees, the reporting of the Agency’s performance commitments associated with these fees, and the need for FDA to convey how these fees are executed calls for strong financial governance. This governance includes an understanding of the design of these programs, clear financial plans, data-driven decisions on resource allocation, consistency and transparency about assumptions, reliable financial forecasting, and accountability for resources spent.

FDA leverages the User Fee Financial Management Committee (UFFMC) for user fee governance. The UFFMC consists of senior financial, business operations, and program experts across the Agency that evaluate user fee resource needs, develop financial allocation plans, and forecast resource requirements – both programmatic and administrative – to support user fee financial decisions. The UFFMC is responsible for providing oversight and support of appropriate standards and policies to ensure FDA’s compliance with sound financial management practices as well as ensuring FDA’s compliance with statutory provisions that authorize FDA to collect and spend user fees. The UFFMC receives policy guidance and strategic direction directly from FDA’s Executive Committee relative to how the Agency will forecast and react to industry trends, plan and manage its research agenda in support of the user fee programs, and
forecast its user fee workload. The UFFMC advises the Executive Committee and other Center and Office-level bodies on a variety of financial and performance-related topics. The Executive Committee is the FDA council comprised of the FDA Commissioner, the Deputy Commissioners, and the Center Directors.

Working Capital Fund/Cost Allocation

FDA utilizes a Cost Allocation and Recovery framework as part of the financial management of user fee resources. Congress authorized FDA to establish a Working Capital Fund (WCF) to finance centralized services (see section 722 of Division A of P.L. 115-141). The WCF benefits the financial management of Agency funds by:

- Increasing transparency through defining administrative activities performed for Centers and Offices and allocating costs based on Agency usage.
- Strengthening accountability by improving the tracking and management of administrative costs, including costs charged to user fees for administrative services.
- Promoting efficiency by optimizing customer usage and improving the management of user fee administrative costs over time.
- Leveraging the WCF governance structure to ensure FDA leadership engagement in decision making relative to administrative costs, efficiency opportunities, recapitalization, and burden on all funding sources – including user fees.

Internal Controls

The Department of Health and Human Services (HHS) provides guidance to its operating divisions (OpDivs) to implement the Federal Managers’ Financial Integrity Act of 1982 (FMFIA) through its FMFIA Guidelines, which is intended to strengthen internal controls and accounting systems. OpDivs, including FDA, are responsible for developing and maintaining internal control and compliance programs that include programmatic and operational controls, as well as reporting controls to support sound financial management. Additionally, FDA established an Enterprise Risk Management (ERM) Program, with an ERM Council as the governance body responsible for providing overall oversight and accountability. For further information regarding the Internal Controls and Enterprise Risk Management, please refer to the GDUFA Financial Report.¹

¹ GDUFA Financial Reports https://www.fda.gov/about-fda/user-fee-financial-reports/gdufa-financial-reports
E. User Fee Background and Structure

Under GDUFA, FDA assesses and collects fees from human generic drug manufacturers to help fund human generic drug activities. The Federal Food, Drug, and Cosmetic Act (FD&C Act), as amended by the FDA User Fee Reauthorization Act of 2022, authorizes FDA to assess and collect fees from industry to supplement non-user fee appropriations that the Agency spends on human generic drug activities.

Originally authorized in 2012, GDUFA was reauthorized by Congress in 2017 (GDUFA II) and most recently in 2022. The FDA User Fee Reauthorization Act of 2022 included the Generic Drug User Fee Amendments of 2022, also known as GDUFA III, which extended the program from October 1, 2022, through September 30, 2027. This five-year reauthorization helps ensure continued funding for FDA from FY 2023 through FY 2027 to support program innovation, evaluation, and improvement. GDUFA III continues FDA’s authority to assess user fees to help fund critical and measurable enhancements to the performance of FDA’s generic drugs program, bringing greater predictability and timeliness to the review of generic drug applications. This delivers tremendous public health benefits by helping to provide the public access to safe, affordable, effective, and high-quality generic drugs.

FDA spends appropriated GDUFA user fee collections and non-user fee appropriations to hire, support, and maintain personnel for the review of generic drug product submissions to help ensure that safe, effective, and high-quality generic drug products are available to the American public.

The fee types remain unchanged from GDUFA II and are described below.

Exhibit 2 outlines the GDUFA III user fee structure.
### Exhibit 2: GDUFA III Fee Structure

<table>
<thead>
<tr>
<th>Fee Type</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Abbreviated New Drug Application (ANDA)</strong></td>
<td>An ANDA filing fee is incurred upon submission of an abbreviated new drug application.</td>
</tr>
<tr>
<td><strong>Type II Domestic and Foreign Active Pharmaceutical Ingredients (API) Drug Master File (DMF)</strong></td>
<td>The one-time DMF fee is incurred on whichever of the following dates occurs earlier: (1) the first time a generic drug submission references that DMF by an initial letter of authorization on or after October 1, 2012, or (2) the date the DMF holder requests the initial completeness assessment.</td>
</tr>
<tr>
<td><strong>Program:</strong> Small, Medium, Large</td>
<td>Each person (including its affiliates) will be assessed an annual fee depending on the number of approved ANDAs in the person’s portfolio.</td>
</tr>
<tr>
<td><strong>Facility:</strong> Domestic and Foreign (API)</td>
<td>An API facility fee is owed by each person who owns a facility that is identified in (1) at least one approved generic drug submission in which the facility is approved to produce one or more APIs or (2) in a Type II API drug master file referenced in at least one approved generic drug submission. An additional $15,000 is assessed for a facility located outside the United States and its territories and possessions.</td>
</tr>
<tr>
<td><strong>Facility:</strong> Domestic and Foreign Finished Dosage Form (FDF)</td>
<td>An FDF facility fee is owed by each person who owns a facility that is identified in at least one generic drug submission that is approved to produce one or more FDFs of a human generic drug. An additional $15,000 is assessed for a facility located outside the United States and its territories and possessions.</td>
</tr>
<tr>
<td><strong>Facility:</strong> Domestic and Foreign Contract Manufacturing Organization (CMO)</td>
<td>An annual CMO facility fee is owed by each person who owns an FDF facility that is identified in at least one approved ANDA, where the facility is not identified in an approved ANDA held by the owner of that facility or its affiliates. An additional $15,000 is assessed for a facility located outside the United States and its territories and possessions.</td>
</tr>
</tbody>
</table>

The statute specifies how the fees must be calculated each fiscal year, including annual adjustments that must be made for inflation. Starting with FY 2024, a capacity planning adjustment (CPA) calculation will be made and the fee revenues and fees will be adjusted, as needed, to reflect changes in resource capacity needs for human generic drug activities. In addition, adjustments may be made for the operating reserve, including a required decrease as applicable. These changes will be discussed in the
following section. The fee amounts are to be published in the *Federal Register* each year, typically at the beginning of August.\(^2\)

GDUFA user fees collected are not a fee-for-service. The user fees that are collected are pooled and may be used for the allowable activities as defined in the FD&C Act. Refer to Appendix A for a detailed list of allowable and excluded activities.

**F. Forward View**

GDUFA III helps ensure continuity for FDA’s generic drug review program by providing for stable and consistent funding during fiscal years 2023 to 2027 to support FDA’s mission to provide the American public timely access to high-quality, affordable generic drugs. Specifically, these funds will enable FDA to implement important program enhancements related to the assessment of ANDAs and to hire and retain the necessary scientific and technical talent needed to deliver GDUFA performance commitments and help achieve related public health priorities.

*Highlights of the GDUFA III Commitment Letter*

The GDUFA III Commitment Letter describes program enhancements agreed to by FDA and industry designed to improve the predictability and transparency of ANDA assessments and to minimize the number of assessment cycles necessary for approval. For example, FDA’s discretion to take imminent actions (i.e., FDA continuing the assessment of an ANDA past the goal date if it may be possible to approve or tentatively approve an ANDA within 60 days after the goal date), or to extend certain goal dates during its assessment of an ANDA (e.g., Information Requests and Discipline Review Letters classified as “major” may extend the goal date), increases opportunities for first cycle or current cycle approvals under GDUFA III. GDUFA III enhancements under the Commitment Letter related to the content, timing, and assessment of a pre-submission facility correspondence support FDA’s ability to meet related priority review goals.

The GDUFA III Commitment Letter continues focus on the development of complex generic products which, because of their unique scientific and regulatory considerations, are harder to develop. Certain new program enhancements are specifically designed to facilitate the development, assessment, and approval of complex generic products. For example, the GDUFA III Commitment Letter includes enhanced pathways for discussions between FDA and prospective applicants before an ANDA for a complex product is submitted, and between FDA and applicants while an ANDA for a complex product is under assessment or after a complete response letter is issued. GDUFA III also continues to promote and advance scientific research around complex generic drug development. This research helps to ensure that regulatory standards, recommendations, and decisions are based on the most current scientific evidence and

\(^2\) See the GDUFA user fee rates archive at [GDUFA User Fee Rates Archive](https://gdufauserfees.fda.gov).
directly supports the FDA’s ability to meet new goal dates for issuing product specific guidances for complex products.

The GDUFA III Commitment Letter also provides more opportunities for timely regulatory and/or scientific advice on a specific element of generic drug product development or certain post-approval submission requirements through program enhancements to the controlled correspondence program. Similarly, GDUFA III’s new commitments to enhance FDA’s processes for reviewing and responding to suitability petitions will facilitate more timely responses to these submissions.

**Changes to Fee Structure and Fee-Setting Mechanisms in GDUFA III**

Although the fee types remain the same as under GDUFA II, under amendments made to the FD&C Act as part of GDUFA III, several changes were made to the fee structure:

1. The proportion of fee revenues derived from API Facility fees will decrease from seven percent in GDUFA II to six percent in GDUFA III.

2. Under the GDUFA II fee structure, CMOs paid one-third the annual fee paid by firms that manufacture under ANDAs which they or their affiliates own. Under GDUFA III, CMOs will pay 24 percent of the annual fee paid by firms that manufacture under ANDAs which they or their affiliates own.

3. The proportion of fee revenues derived from the Generic Drug Applicant Program Fee will increase from 35 percent in GDUFA II to 36 percent in GDUFA III.

There were several changes to the fee-setting mechanisms under GDUFA III amendments to the FD&C Act:

1. The base revenue amount for each fiscal year in GDUFA III will be set using the general approach used in GDUFA II with some refinements. The total target revenue for FY 2023 is $582,500,000. The base revenue amount for subsequent fiscal years will be based on the total target revenue amount for the prior fiscal year, excluding any operating reserve adjustment for that prior fiscal year.

2. Congress made a technical fix to the inflation adjustment used in GDUFA. The previous inflation adjustment referenced a Consumer Price Index (CPI) that the U.S. Bureau of Labor Statistics had discontinued (i.e., Washington-Baltimore, DC-MD-VA-WV; not seasonally adjusted; all items; annual index) and that it replaced with two separate indices (i.e., "Washington-Arlington-Alexandria, DC-VA-MD-WV" and "Baltimore-Columbia-Towson, MD"). In order to apply a CPI which best reflects the geographic region in which FDA is headquartered and which provides the most current data available, the Washington-Arlington-Alexandria index will be used in GDUFA III to calculate the inflation factor.

3. A capacity planning adjustment (CPA) was added, beginning with FY 2024, to increase annual revenue as needed to account for changes in program workload. The CPA has an annual cap of three percent of inflation-adjusted revenue except
when certain circumstances are met (in which case the cap is increased to four percent). FDA will describe its application of the CPA methodology in the Federal Register notice publishing GDUFA fees each year.

4. The final year adjustment was replaced with an operating reserve adjustment that allows FDA to adjust fees for FY 2024 or subsequently during GDUFA III to maintain sufficient operating reserves of carryover user fees. FDA may increase fees to maintain up to eight weeks of reserve in FY 2024, nine weeks of reserve in FY 2025, and 10 weeks for FY 2026 and FY 2027. If the estimated carryover balance is in excess of 12 weeks of operating reserves, FDA is required to decrease fees for that fiscal year to reduce the operating reserve to not more than 12 weeks. FDA will provide the rationale for adjustments to the operating reserve in the annual Federal Register notice publishing fee rates for that fiscal year.

**Efforts to Enhance Financial Management**

Under the GDUFA III Commitment Letter, FDA continued its commitment to mature the Agency’s resource capacity planning function, including utilization of modernized time reporting, to support enhanced management of GDUFA resources in GDUFA III and help ensure alignment of user fee resources to staff workload.

To further these efforts, a second assessment of the resource capacity planning capability, including the CPA, will be conducted during GDUFA III that will include examining the ability of the CPA to forecast appropriate resource needs for the GDUFA Program, including an assessment of the scope of the workload drivers in the CPA and their ability to represent the overall workload of the GDUFA Program. The resulting report will be published for public comment and discussed at the FY 2026 GDUFA five-year financial planning meeting. The findings and recommendations of the evaluation may inform future reauthorizations.

FDA also made commitments in GDUFA III to enhance efficiency and transparency in the administration of GDUFA’s financial resources. This includes publishing a five-year plan (this plan), to be updated annually. FDA will also hold an annual public meeting to discuss this five-year financial plan, along with the Agency’s progress in implementing resource capacity planning, including the continual improvement of the capacity planning adjustment and time reporting, and the integration of resource capacity planning in resource and operational decision-making processes.

**Other Financial Impacts**

Section 905(b) of the FDA Reauthorization Act of 2017 (FDARA) amended the FD&C Act to provide that the types of fee-coverable costs under the Prescription Drug User Fee Act (PDUFA) program, the GDUFA program, the Medical Devices User Fee Amendments (MDUFA) program, and the Biosimilar User Fee Act (BsUFA) program will narrow on October 1, 2023. The statutory definition of allowable cost categories, i.e., the “resources allocated for human generic drug activities” under GDUFA, is what
determines the type of expenses related to human generic drug activities that GDUFA fees can be used for.

Due to a later provision in the Food and Drug Omnibus Reform Act (as included in the Consolidated Appropriations Act, 2023), section 744B of the FD&C Act was amended to clarify that while user fees may no longer be used to pay for certain costs, these costs as funded by budget authority will count toward the non-user fee spending trigger. For further information, see Note 5.

This change is not expected to have an impact on the non-user fee spending trigger. The systems supporting these programs; however, are complex and multi-faceted. As such, FDA will continue to plan for and monitor the impacts of these changes to ensure minimal disruption to its user fee commitments and public health mission.

Financial Information

This section provides an overview of the financial outlook for GDUFA for the FY 2023 through FY 2027 reauthorization period including budgetary resources, obligations, carryover, non-user fee appropriations requirements, and planned hiring. The forecasts included in this section are driven by the initiatives and goals as outlined in the Forward View section of this plan.

G. User Fee Program Financial Summaries

Table 1 represents a summary of the estimated GDUFA financial position, as it relates to user fee budgetary resources. This table also provides an overview of estimated obligations for which the user fee resources would be used. Annual updates to this plan will provide actual amounts for the prior fiscal years. The financial notes referenced in this table can be found in Appendix B.

Table 1: Human Generic Drug User Fee Collections, Obligations, and Carryover for Fiscal Year 2023 through Fiscal Year 2027

<table>
<thead>
<tr>
<th>Budgetary Resources</th>
<th>Notes</th>
<th>FY2023 Estimate</th>
<th>FY2024 Estimate</th>
<th>FY2025 Estimate</th>
<th>FY2026 Estimate</th>
<th>FY2027 Estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Target Revenue</td>
<td>Note 1</td>
<td>$582,500,000</td>
<td>$595,034,000</td>
<td>$606,934,000</td>
<td>$619,073,000</td>
<td>$631,454,000</td>
</tr>
<tr>
<td>Cash Collections</td>
<td></td>
<td>$582,500,000</td>
<td>$595,034,000</td>
<td>$606,934,000</td>
<td>$619,073,000</td>
<td>$631,454,000</td>
</tr>
<tr>
<td>Recoveries</td>
<td>Note 2</td>
<td>$10,000,000</td>
<td>$10,000,000</td>
<td>$10,000,000</td>
<td>$10,000,000</td>
<td>$10,000,000</td>
</tr>
<tr>
<td>Carryover Available for Use, Beginning of Year</td>
<td>$131,211,761</td>
<td>$134,626,041</td>
<td>$134,486,874</td>
<td>$135,624,692</td>
<td>$128,428,564</td>
<td></td>
</tr>
<tr>
<td>Total Budgetary Resources</td>
<td></td>
<td>$723,711,761</td>
<td>$739,660,041</td>
<td>$751,420,874</td>
<td>$764,697,692</td>
<td>$769,882,564</td>
</tr>
<tr>
<td>Obligations</td>
<td>Notes</td>
<td>FY2023 Estimate</td>
<td>FY2024 Estimate</td>
<td>FY2025 Estimate</td>
<td>FY2026 Estimate</td>
<td>FY2027 Estimate</td>
</tr>
<tr>
<td>-------------------------------------</td>
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<td>----------------</td>
<td>----------------</td>
<td>----------------</td>
<td>----------------</td>
</tr>
<tr>
<td>Total Payroll &amp; Operating</td>
<td>Note 3</td>
<td>$495,261,771</td>
<td>$520,565,927</td>
<td>$530,342,870</td>
<td>$549,961,283</td>
<td>$561,120,910</td>
</tr>
<tr>
<td>Total Rent</td>
<td>Note 5</td>
<td>$21,595,013</td>
<td>$12,242,365</td>
<td>$12,364,788</td>
<td>$12,488,436</td>
<td>$12,613,320</td>
</tr>
<tr>
<td>Total Shared Services</td>
<td>Note 6</td>
<td>$72,228,936</td>
<td>$72,364,876</td>
<td>$73,088,524</td>
<td>$73,819,410</td>
<td>$74,557,604</td>
</tr>
<tr>
<td><strong>Total Obligations</strong></td>
<td></td>
<td><strong>$589,085,720</strong></td>
<td><strong>$605,173,167</strong></td>
<td><strong>$615,796,182</strong></td>
<td><strong>$636,269,129</strong></td>
<td><strong>$648,291,835</strong></td>
</tr>
<tr>
<td>Carryover</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total Carryover, End of Year</td>
<td></td>
<td>$134,626,041</td>
<td>$134,486,874</td>
<td>$135,624,692</td>
<td>$128,428,564</td>
<td>$121,590,729</td>
</tr>
<tr>
<td>Future Year Refunds Allowance, Set Aside</td>
<td></td>
<td>($4,000,000)</td>
<td>($4,000,000)</td>
<td>($4,000,000)</td>
<td>($4,000,000)</td>
<td>($4,000,000)</td>
</tr>
<tr>
<td>Carryover Net of Set Aside, End of Year</td>
<td></td>
<td>$130,626,041</td>
<td>$130,486,874</td>
<td>$131,624,692</td>
<td>$124,428,564</td>
<td>$117,590,729</td>
</tr>
</tbody>
</table>

Target Revenue has been rounded to the nearest thousand dollars
All other numbers have been rounded to the nearest dollar

**Budgetary Resources:** The Total Budgetary Resources component of Table 1 illustrates the total user fee funding estimates for FY 2023 through FY 2027. Budgetary resources include net collections, recoveries, and carryover amounts.

Budgetary resources are discussed in more detail in Section H.

**Obligations:** The Obligations component of Table 1 shows the planned annual expenditure for FY 2023 through FY 2027 of GDUFA fee funds broken out into major expense categories. GDUFA fees may be expended only for certain costs to support “human generic drug activities,” as defined in the statute. For more information on the allowable and excluded costs, see Appendix A.

Obligations are discussed in more detail in Section I.

**Carryover:** GDUFA fees collected, appropriated, and not obligated at the end of the fiscal year remain available to support the GDUFA program in future fiscal years. In this report, such fee funds are referred to as the “total carryover” or “GDUFA carryover.”

Carryover is discussed in more detail in Section J.

**H. Budgetary Resources**

Budgetary resources include net collections, recoveries, and carryover amounts. Net collections are a function of the annual target revenue amount and the number of fees paid. This section describes the process for the setting of the annual target revenue amount and then describes the estimated total budgetary resources.
Annual updates to this plan will update the actual revenue amount for the current fiscal year and the actual collections amount from the preceding fiscal years.

Table 2 outlines the annual target revenue amounts for each fiscal year. The financial notes referenced in this table can be found in Appendix B.

Table 2: Human Generic Drug User Fee Target Revenue for Fiscal Year 2023 through Fiscal Year 2027

<table>
<thead>
<tr>
<th>Budgetary Resources</th>
<th>Notes</th>
<th>FY2023 Actual</th>
<th>FY2024 Estimate</th>
<th>FY2025 Estimate</th>
<th>FY2026 Estimate</th>
<th>FY2027 Estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Base Amount</td>
<td></td>
<td>$582,500,000†</td>
<td>$582,500,000</td>
<td>$595,033,653</td>
<td>$606,934,326</td>
<td>$619,073,013</td>
</tr>
<tr>
<td>Inflation Adjustment</td>
<td>Note 4</td>
<td>$0</td>
<td>$12,533,653</td>
<td>$11,900,673</td>
<td>$12,138,687</td>
<td>$12,381,460</td>
</tr>
<tr>
<td>Capacity Planning</td>
<td>Note 10</td>
<td>N/A</td>
<td>TBD</td>
<td>TBD</td>
<td>TBD</td>
<td>TBD</td>
</tr>
<tr>
<td>Operating Reserve</td>
<td>Note 7</td>
<td>N/A</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
</tr>
<tr>
<td>Target Revenue Total</td>
<td>Note 1</td>
<td>$582,500,000†</td>
<td>$595,033,653</td>
<td>$606,934,326</td>
<td>$619,073,013</td>
<td>$631,454,473</td>
</tr>
</tbody>
</table>

Numbers have been rounded to the nearest dollar (note: The Target Revenue amount in Table 1 has been rounded to the nearest thousand dollars.)

†Indicates an actual amount
N/A = Not Applicable, TBD = To Be Determined

Target Revenue: The process for setting the annual target revenue is defined in the statute and is described below.

- **Statutory Base:** The base amount for FY 2023 is specified in the statute ($582,500,000). There are no adjustments to this amount for FY 2023. Starting with FY 2024, the base amount is the total revenue amount (target revenue) for the prior fiscal year, not including any operating reserve adjustment for that prior year. This base amount is adjusted for inflation and by the CPA (as needed), and may be further adjusted for operating reserve, as described below. See Note 1 for a diagram of this process.

- **Inflation Adjustment:** The inflation adjustment, which begins with FY 2024, adjusts the base amount to maintain the purchasing power of fee funds in consideration of inflation. The adjustment is a composite measure that weights operating expenses by changes in the applicable CPI and payroll-related expenses by changes in FDA’s average personnel compensation and benefits amounts.

Per the statute, there was no inflation adjustment for FY 2023. The inflation adjustment for future years, for purposes of this plan, is estimated to be a two percent increase, per the Federal Reserve’s inflation targeting policy. For more information, see Note 4.
• **Capacity Planning Adjustment:** Beginning with FY 2024, FDA shall use the capacity planning adjustment to further adjust, as needed, the fee revenue and fees to reflect changes in the resource capacity needs of FDA for human generic drug activities. FDA does not have an estimated amount for the CPA at this time.

**Operating Reserve Adjustment:** The operating reserve adjustment was established to provide a mechanism to support the management of the carryover balance from year to year. The operating reserve adjustment would increase or decrease, as applicable, the fee revenue amount to set fees. Should FDA make an operating reserve adjustment, either up or down, FDA must explain its rationale in the annual GDUFA fee-setting Federal Register Notice. For more information, see Note 7.

Table 3 connects the target revenue to the net collections while showing the estimated total budgetary resources for each fiscal year. The financial notes referenced in this table can be found in Appendix B.

**Table 3: Generic Drug User Fee Budgetary Resources FY 2023 through FY 2027**

<table>
<thead>
<tr>
<th>Budgetary Resources</th>
<th>Notes</th>
<th>FY2023 Estimate</th>
<th>FY2024 Estimate</th>
<th>FY2025 Estimate</th>
<th>FY2026 Estimate</th>
<th>FY2027 Estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Target Revenue</td>
<td>Note 1</td>
<td>$582,500,000</td>
<td>$595,034,000</td>
<td>$606,934,000</td>
<td>$619,073,000</td>
<td>$631,454,000</td>
</tr>
<tr>
<td>Net Collections</td>
<td></td>
<td>$582,500,000</td>
<td>$595,034,000</td>
<td>$606,934,000</td>
<td>$619,073,000</td>
<td>$631,454,000</td>
</tr>
<tr>
<td>Recoveries</td>
<td>Note 2</td>
<td>$10,000,000</td>
<td>$10,000,000</td>
<td>$10,000,000</td>
<td>$10,000,000</td>
<td>$10,000,000</td>
</tr>
<tr>
<td>Total Carryover, Beginning of Year</td>
<td></td>
<td>$131,211,761</td>
<td>$134,626,041</td>
<td>$134,486,874</td>
<td>$135,624,692</td>
<td>$128,428,564</td>
</tr>
<tr>
<td>Total Budgetary Resources</td>
<td></td>
<td>$723,711,761</td>
<td>$739,660,041</td>
<td>$751,420,874</td>
<td>$764,697,692</td>
<td>$769,882,564</td>
</tr>
</tbody>
</table>

*Numbers have been rounded to the nearest dollar*

**Budgetary Resources:** Budgetary resources include net collections, recoveries, and carryover amounts.

- **Net Collections:** FDA assumes, for planning purposes, that net collections will equal the target revenue amount. In practice, net collections may differ from the annual target revenue amount if the actual number of fee-paying units differs from the number of fee-paying units estimated when fees are set each year. The target revenue amount includes any applicable adjustments made for the year, including inflation adjustment, CPA, and the operating reserve adjustment.

- **Recoveries:** For the purposes of this plan, future year recoveries are estimated to be $10 million annually. This value is based on the actual amount recovered from the prior years plus the current actual recoveries rate for FY 2023. Additional details on recoveries are included in Note 2.
• **Total Carryover, beginning of year**: Total carryover represents the balance of unspent GDUFA fee funds at the beginning of the fiscal year. The total year carryover at the end of one fiscal year equals the total carryover at the beginning of the subsequent fiscal year. Carryover is discussed in more detail in **Section J**.

**Net Collections vs. Cohort Year Collections**: User fee collections are reported in two different ways:

• **Net Collections**: Net collections are the actual dollar amounts collected in a fiscal year, regardless of the fiscal year the fee was due. **Table 1** and **Table 3** report net collections.

• **Cohort Year Collections**: Cohort year collections represent the fiscal year for which the fee was originally due. **Table 4** reports cohort year collections.

Example: Assume a fee was due in FY 2023 but was paid in FY 2024. This would be reported as a net collection in FY 2024 and a cohort year collection in FY 2023.

**Table 4** presents the forecasted total annual collections by fee type and cohort year. Refer to **Section E** for more background and information on the GDUFA III fee structure.

<table>
<thead>
<tr>
<th>Fee Type</th>
<th>Cohort Year 2023 Estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td>ANDA (Application) Fees</td>
<td>$192,225,000</td>
</tr>
<tr>
<td>DMF Fees</td>
<td>$29,125,000</td>
</tr>
<tr>
<td>Facilities Fees (FDF, CMO, and API)</td>
<td>$151,450,000</td>
</tr>
<tr>
<td>GDUFA Program Fees</td>
<td>$209,700,000</td>
</tr>
<tr>
<td><strong>Total Net Collections</strong></td>
<td><strong>$582,500,000</strong></td>
</tr>
</tbody>
</table>

*Estimated Total Net Collections have been rounded to the nearest thousand dollars*

The annual updates to this plan will provide the actual Net Collections amounts by cohort year for the preceding year(s) as well as updated planned amounts for the remaining fiscal years.

**I. User Fee Obligations**

GDUFA fees may be expended only for certain costs necessary to support “human generic drug activities,” as defined in section 744A(9) of the FD&C Act. For more information on the allowable and excluded costs, see **Appendix A**.

**Table 5** provides a breakout of planned user fee obligations by expense category for the 5 years represented in this plan. The annual updates to this plan will provide actual
amounts for the preceding fiscal year, as well as updated planned amounts for the remaining fiscal years. The financial notes can be found in Appendix B.

Table 5: Human Generic Drug User Fee Obligations by Expense Category for Fiscal Year 2023 through Fiscal Year 2027

<table>
<thead>
<tr>
<th>User Fee Obligations</th>
<th>Notes</th>
<th>FY2023 Estimate</th>
<th>FY2024 Estimate</th>
<th>FY2025 Estimate</th>
<th>FY2026 Estimate</th>
<th>FY2027 Estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Payroll &amp; Operating</td>
<td>Note 3</td>
<td>$1,040,390</td>
<td>$1,062,776</td>
<td>$1,084,031</td>
<td>$1,105,712</td>
<td>$1,127,826</td>
</tr>
<tr>
<td>CBER</td>
<td></td>
<td>$403,639,923</td>
<td>$426,626,067</td>
<td>$435,813,589</td>
<td>$453,541,550</td>
<td>$462,760,296</td>
</tr>
<tr>
<td>CDER</td>
<td></td>
<td>$53,494,587</td>
<td>$56,955,304</td>
<td>$58,094,410</td>
<td>$59,256,298</td>
<td>$60,441,424</td>
</tr>
<tr>
<td>ORA</td>
<td></td>
<td>$37,086,872</td>
<td>$35,921,779</td>
<td>$35,350,840</td>
<td>$36,057,723</td>
<td>$36,791,365</td>
</tr>
<tr>
<td>HQ</td>
<td></td>
<td>$21,595,013</td>
<td>$12,242,365</td>
<td>$12,364,788</td>
<td>$12,488,436</td>
<td>$12,613,320</td>
</tr>
<tr>
<td>Total Rent</td>
<td>Note 5</td>
<td>$72,228,936</td>
<td>$72,364,876</td>
<td>$73,088,524</td>
<td>$73,819,410</td>
<td>$74,557,604</td>
</tr>
<tr>
<td>Total Shared Services</td>
<td>Note 6</td>
<td>$589,085,720</td>
<td>$605,173,167</td>
<td>$615,796,182</td>
<td>$636,269,129</td>
<td>$648,291,835</td>
</tr>
</tbody>
</table>

Total obligations include payroll and operating, rent, and shared services costs. Non-user fee funds supporting the GDUFA program are not included here. The details of each component of total obligations are as follows:

- **Payroll and Operating**: These obligations provide for all payroll and operating costs that support the allowable activities for which GDUFA fees may be expended, as set forth in the statute. These allowable activities include, for example, core regulatory review functions, pre-approval inspections, guidance and policy development activities, regulatory science activities, and management and administrative functions that support the GDUFA program.

  Payroll and operating are presented by each major organizational component relevant to the GDUFA program.

- **Rent**: This amount is paid to the General Services Administration (GSA) for the federal buildings that FDA occupies, as well as to non-federal sources for direct leases and services (see Note 5). Rental rates vary based on the type and location of the space provided.

  For FY 2023, rent includes all costs allowable under prior GDUFA authorizations. Due to amendments made by section 905 of FDARA, certain previously allowable costs will be excluded starting in FY 2024 (see Note 5). As a result, rent costs drop from FY 2023 to FY 2024.
This lower rent cost is then adjusted for inflation using the inflation adjustment amount of amount of 2.1517 percent. The exclusion of previously allowable costs exceeds the increase from applying the inflation adjustment, resulting in a lower FY 2024 rent cost than FY 2023.

- **Shared Services**: FDA has several shared service organizations that provide support across the user fee programs, such as human resources and information technology (IT). Shared services are located within the Working Capital Fund (WCF). **Note 6** provides a full list of what is contained in the WCF.

FY 2024 Shared Service amounts use the inflation adjustment amount of 3.4398 percent, offset by some one-time adjustments that occurred in FY 2023. All years also include small, proportionate increases to support the growth of the program.

Rent and Shared Services projections are informed by prior year actuals. For FY 2025 through FY 2027, the Rent and Shared Services future year amounts, for the purposes of this plan, are assumed to have an increase of an inflation amount of one percent yearly. Yearly costs are determined by the Cost Allocation and Recovery framework discussed previously. As stated in **Section H**, Payroll and Operating projections for future year amounts is assumed to have an increase of two percent yearly.

**Exhibit 3** provides an illustration of historical GDUFA II obligations and projected GDUFA III needs.
GDUFA III obligations are expected to continue to grow as the program hires new personnel to deliver on negotiated enhancements and as a result of inflationary pressures.

**J. User Fee Carryover**

GDUFA fees collected, appropriated, and not obligated at the end of the fiscal year remain available to support the GDUFA program in future fiscal years.

Maintaining an appropriate level of carryover enables FDA to mitigate financial risks to the program, including, for example, the risk of under collecting fees and the risk of a lapse in appropriations so that FDA can continue performing activities related to human generic drug activities under such financial constraints. FDA may set aside available user fee funds in the carryover for certain purposes, including, for example, for processing future year refunds.

The net change in carryover each year is equal to net collections minus net obligations. This is demonstrated best in Table 1 above.

**Table 6** provides projections of GDUFA carryover balances at the end of each fiscal year. Forecasted estimates will be updated with actual amounts in future Five-Year Financial Plan annual updates. The financial notes can be found in Appendix B.
Table 6: GDUFA Carryover by Fiscal Year

<table>
<thead>
<tr>
<th>Carryover</th>
<th>FY2023 Estimate</th>
<th>FY2024 Estimate</th>
<th>FY2025 Estimate</th>
<th>FY2026 Estimate</th>
<th>FY2027 Estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Carryover, End of Year</td>
<td>$134,626,041</td>
<td>$134,486,874</td>
<td>$135,624,692</td>
<td>$128,428,564</td>
<td>$121,590,729</td>
</tr>
<tr>
<td>Future Year Refunds Allowance, Set Aside</td>
<td>($4,000,000)</td>
<td>($4,000,000)</td>
<td>($4,000,000)</td>
<td>($4,000,000)</td>
<td>($4,000,000)</td>
</tr>
<tr>
<td>Carryover Net of Set Aside, End of Year</td>
<td>$130,626,041</td>
<td>$130,486,874</td>
<td>$131,624,692</td>
<td>$124,428,564</td>
<td>$117,590,729</td>
</tr>
</tbody>
</table>

Numbers have been rounded to the nearest dollar.

These terms are defined below:

- **Total Carryover, End of Year**: This is the total amount of unobligated fee funds at the end of the fiscal year.

- **Future Year Refunds Allowance, Set Aside**: FDA maintains a limited amount to provide for any refunds, as a matter of prudent operations. For that purpose, a total of $4,000,000 in fee funds available for obligation is being set aside annually. See Note 8 for additional details.

- **Carryover Net of Set Aside, End of Year**: This is the total carryover less any carryover funds subject to set asides.

Exhibit 4 below shows the historic trend of carryover in GDUFA II and the forecasted carryover in GDUFA III.
Looking forward into GDUFA III, the operating reserve adjustment will be a mitigation tool to ensure carryover remains below the 12-week threshold. FDA will monitor the operating reserve levels and will apply appropriate discretion within the statutory framework in the utilization of any increase each year, or as applicable implement a required decrease. See Table 7 below for the operating reserve threshold amounts.

Table 7: Operating Reserve Thresholds

<table>
<thead>
<tr>
<th>Operating Reserve</th>
<th>FY2023 Estimate</th>
<th>FY2024 Estimate</th>
<th>FY2025 Estimate</th>
<th>FY2026 Estimate</th>
<th>FY2027 Estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-Week Operating Amount</td>
<td>N/A</td>
<td>$11,442,955</td>
<td>$11,671,814</td>
<td>$11,905,250</td>
<td>$12,143,355</td>
</tr>
<tr>
<td>Discretionary Operating Reserve Statutory Increase Threshold (weeks)</td>
<td>N/A</td>
<td>8</td>
<td>9</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Discretionary Operating Reserve Statutory Increase Threshold ($)</td>
<td>N/A</td>
<td>$91,543,639</td>
<td>$105,046,326</td>
<td>$119,052,503</td>
<td>$121,433,553</td>
</tr>
<tr>
<td>Total Carryover, End of Year</td>
<td>$134,626,041</td>
<td>$134,486,874</td>
<td>$135,624,692</td>
<td>$128,428,564</td>
<td>$121,590,729</td>
</tr>
<tr>
<td>Operating Reserve Statutory Decrease Threshold (weeks)</td>
<td>N/A</td>
<td>12</td>
<td>12</td>
<td>12</td>
<td>12</td>
</tr>
<tr>
<td>Operating Reserve Statutory Decrease Threshold ($)</td>
<td>N/A</td>
<td>$137,315,458</td>
<td>$140,061,768</td>
<td>$142,863,003</td>
<td>$145,720,263</td>
</tr>
</tbody>
</table>

Numbers have been rounded to the nearest dollar.
Beginning with FY 2024, should the operating reserves of carryover user fees drop below a statutorily-specified, week-based level, FDA may use the operating reserve adjustment to further increase the fee revenue and fees to provide operating reserves of carryover user fees for human generic drug activities for not more than the number of weeks so specified: 8 weeks in FY 2024; 9 weeks in FY 2025; and 10 weeks in FY 2026 and FY 2027.

For any fiscal year FDA has carryover balances for human generic drug activities in excess of 12 weeks of operating reserves, FDA shall decrease the fee revenue and fees to provide for not more than 12 weeks of such operating reserves.

To calculate the dollar amounts of these week-based thresholds, applicable adjustments for inflation and capacity planning are applied to the base revenue. This amount is then divided by 52 to generate the one-week operating amount. The one-week operating amount is then multiplied by the threshold number of weeks.

**K. Non-User Fee Appropriations**

For FDA to obligate user fees collected under GDUFA, a certain amount of non-user fee appropriations must be spent on human generic drug activities during that fiscal year. This is often referred to as a “non-user fee spending trigger.” Table 8 presents the forecasted non-user fee spending triggers for FY 2023 through FY 2027.

**Table 8: Minimum Allocation of GDUFA Non-User Fee Appropriations by Fiscal Year**

<table>
<thead>
<tr>
<th>FY2023 Estimate</th>
<th>FY2024 Estimate</th>
<th>FY2025 Estimate</th>
<th>FY2026 Estimate</th>
<th>FY2027 Estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td>$118,492,290</td>
<td>$127,669,945</td>
<td>$130,223,344</td>
<td>$132,827,811</td>
<td>$135,484,367</td>
</tr>
</tbody>
</table>

Numbers have been rounded to the nearest dollar

The non-user fee spending trigger amount is determined by multiplying the base amount of non-user fee appropriations spent on human generic drug activities ($97 million) times the adjustment factor for the fiscal year. See Note 9 for more details on the adjustment factor.

As a result of amendments under section 905(b) of FDARA, starting in FY 2024, certain costs will be shifted from user fee spending to non-user fee appropriations spending. Even though these costs will be shifted, these costs will still be counted towards the spending trigger. See Note 5 for more information.

FDA is committed to spend at least the required minimum from non-user fee appropriations each fiscal year. In years when FDA programs do not receive appropriations to cover costs of inflation and mandatory pay increases, FDA activities other than human generic drug activities may be reduced to assure that the allocation of non-user fee appropriations for human generic drug activities meets the requirements of this trigger.
L. Planned Hiring

Under the GDUFA III Commitment Letter, FDA agreed to the hiring of 128 staff in FY 2023 to support the workload associated with initiatives established or expanded by GDUFA III (see section VIII.E.2. of the Commitment Letter). FDA will provide additional information on the progress of the hiring of GDUFA III staff in the GDUFA annual reports.
Challenges, Risk and Mitigation

As is the case with all financial programs, there are certain financial risks and challenges that exist with FDA’s user fee programs. These risks and challenges can vary from program to program, with some in FDA’s control and some out of FDA’s control. An example of a financial risk shared across all user fee programs is that FDA cannot obligate funds in advance of receiving the funds, either through appropriated user fee collections or non-user fee appropriations. FDA can only estimate what the Agency’s total appropriated resources will be in any given year. As a result, FDA has some risk of missing important performance goals, or failing to meet the non-user fee spending trigger for the fiscal year if that total amount is considerably lower than anticipated. Below is a list of foreseeable risks associated with the collections and obligations of funds for which FDA has identified contingency plans; these contingency plans help ensure FDA is able to move forward in the best interests of the program.

- **Under-Executing Planned Spend:** During GDUFA I and to a lesser extent in GDUFA II, budgetary resources had been under spent due to the uncertainty around the timing of budgetary resources availability, non-user fee spending trigger requirements, and difficulties with hiring. To minimize this risk, FDA continues to enhance its planning and execution around the hiring of new staff and contract actions.

- **Uncertainty of Non-User Fee Appropriations Levels:** It is difficult to predict the amount of non-user fee appropriations that will be approved by Congress, which creates planning challenges. With Continuing Resolutions (CR) becoming more prevalent, non-user fee fund levels are often uncertain for a good portion of the fiscal year. Accordingly, FDA has needed to spend at or slightly below levels from the prior authorized fiscal year during the CR period, thus limiting its ability to fully utilize the non-user fee appropriations from the onset.

- **Lapse in Non-User Fee Appropriations:** FDA cannot control this risk; however, under GDUFA III, FDA has the authority to maintain up to a specified week-based level of an operating reserve of appropriated carryover fees, which can be utilized to continue program operations in the event of a lapse in appropriations.

- **Undercollecting and Overcollecting:** If FDA does not receive the estimated number of industry submissions, there may be an excess or deficit in collections relative to the targeted revenue. When FDA undercollects user fees, it leverages its carryover to maintain continuity in operations. When FDA overcollects, the carryover may increase without additional planned expenditures being identified towards which to obligate those funds. The operating reserve adjustment mitigates these risks in GDUFA III. Resource capacity planning helps improve fee setting and allows FDA to adjust for sustained increases in workload. In addition, FDA monitors collections throughout the fiscal year, and the UFFMC
and other FDA senior leaders determine how to mitigate any instances when user fee revenue deviates from forecasted estimates.

- **FDARA section 905**: Starting with FY 2024, FDA cannot use user fees on certain previously allowable types of expenses. This change will have an impact on the finances of the program. FDARA section 905 amended the FD&C Act to narrow the definition of allowable cost categories, i.e., the “resources allocated for human generic drug activities” under GDUFA, which is what determines the types of expenses related to human generic drug activities that GDUFA fees can be used for. FDA is monitoring the impacts to the shifts in funding.
Appendices

A. Allowable and Excluded Costs for the GDUFA Program

Section 744A(9) of the FD&C Act defines the term “human generic drug activities,” in general, as the activities associated with generic drugs and inspection of facilities associated with generic drugs. In summary, costs related to the following have been attributed to human generic drug activities:

<table>
<thead>
<tr>
<th>Included Activities</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The activities necessary for the review of generic drug submissions, including review of DMFs referenced in such submissions.</td>
</tr>
<tr>
<td>2. The issuance of:</td>
</tr>
<tr>
<td>a. Approval letters that approve ANDAs or prior approval supplements to such applications.</td>
</tr>
<tr>
<td>b. Complete response letters that set forth in detail the specific deficiencies in such applications and, where appropriate, the actions necessary to place such applications in condition for approval.</td>
</tr>
<tr>
<td>3. The issuance of letters related to Type II API DMFs that:</td>
</tr>
<tr>
<td>a. Set forth in detail the specific deficiencies in such submissions, and where appropriate, the actions necessary to resolve those deficiencies; or</td>
</tr>
<tr>
<td>b. Document that no deficiencies need to be addressed.</td>
</tr>
<tr>
<td>4. Inspections related to generic drugs.</td>
</tr>
<tr>
<td>5. Monitoring of research conducted in connection with the review of generic drug submissions and DMFs.</td>
</tr>
<tr>
<td>6. Post-market safety activities with respect to drugs approved under ANDAs or supplements, including the following activities:</td>
</tr>
<tr>
<td>a. Collecting, developing, and reviewing safety information on approved drugs including adverse event reports.</td>
</tr>
<tr>
<td>b. Developing and using improved adverse-event data collection systems, including IT systems.</td>
</tr>
<tr>
<td>c. Developing and using improved analytical tools to assess potential safety problems including access to external databases.</td>
</tr>
<tr>
<td>d. Implementing and enforcing section 505(o) (relating to post-approval studies and clinical trials and labeling changes) and section 505(p) (relating to risk evaluation and mitigation strategies) insofar as those activities relate to ANDAs.</td>
</tr>
<tr>
<td>e. Carrying out section 505(k)(5) (relating to adverse-event reports and post-market safety activities).</td>
</tr>
<tr>
<td>7. Regulatory science activities related to generic drugs.</td>
</tr>
</tbody>
</table>

Section 744A(12) of the FD&C Act defines the term “resources allocated for human generic drug activities” as expenses for the following:
Included Expenses

1. Officers and employees of FDA, contractors of FDA, advisory committees, and the costs related to such officers, employees, and committees, and to contracts with such contractors.
2. Management of information and the acquisition, maintenance, and repair of computer resources.
3. Leasing, maintenance, renovation, and repair of facilities and acquisition, maintenance and repair of fixtures, furniture, scientific equipment, and other necessary materials and supplies.
4. Collecting fees under section 744B and accounting for resources allocated for the review of ANDAs and supplements and inspections related to generic drugs.

The GDUFA program excludes costs related to the following:

Excluded Activities

1. All activities necessary for the review of new drug applications, biologic license applications, and investigational new drugs for drugs that will not be approved under ANDAs.
2. The issuance of controlled correspondence unrelated to abbreviated new drug submissions or prior approval supplements.
3. Inspections unrelated to human generic drugs.
4. Monitoring of research unrelated to human generic drug submissions and DMFs.
5. Post-market safety activities apart from those drugs approved under ANDAs or supplements.

Section 744B of the FD&C Act was amended by section 905 of FDARA to limit the definition of certain “resources allocated for human generic drug activities” under section 744A(12)(C), i.e., the types of expenses related to human generic drug activities that fees can cover, to include only expenditures for leasing and necessary scientific equipment starting with FY 2024:

Included Expenses

1. Officers and employees of FDA, contractors of FDA, advisory committees, and the costs related to such officers, employees, and committees, and to contracts with such contractors.
2. Management of information and the acquisition, maintenance, and repair of computer resources.
3. Leasing and necessary scientific equipment.
4. Collecting fees under section 744B and accounting for resources allocated for the review of ANDAs and supplements and inspection related to generic drugs.

Beginning in FY 2024, in addition to costs excluded under the FDARA section 905 amendments, the GDUFA program will continue to exclude costs as outlined above.
B. Financial Notes

Note 1. Annual Target Revenue Methodology

Exhibit 5 is a flowchart that outlines GDUFA III’s Annual Target Revenue Methodology.

Exhibit 5: GDUFA III Annualized Base and Target Revenue Methodology

Note 2. Recoveries

Recoveries account for funds returned to the Agency in the form of deobligations of prior year obligations. For example, recoveries could include funding from a contract that ended in a prior year and was not expended.

Note 3. Pay and Operating Costs

For payroll, employees are required to report their time in an activity-based reporting system, which allows FDA to identify activities that user fees can be used to support. See Appendix A for a listing of those activities. For operating activities (e.g., contracting services), funds are allocated based on the proportion to which those activities support the GDUFA program. If an operating activity solely supports GDUFA, it will be fully funded by the program. If the operating activity is shared, GDUFA will fund the activity in proportion to how it is used by the program as compared to other programs.

Note 4. Inflation Adjustment

The inflation adjustment adjusts the base amount to maintain the purchasing power of fee funds in consideration of inflation. The adjustment is a composite measure that weights operating expenses by changes in the CPI and payroll-related expenses by changes in FDA’s average personnel compensation and benefits amounts.
For FY 2023, the first program year of the current five-year GDUFA authorization (under the FDA User Fee Reauthorization Act of 2022), there is no inflation adjustment, as the target revenue is set by the statute. For FY 2024, the inflation adjustment is estimated to be 2.1517 percent for GDUFA. This is subject to change during the annual fee setting process.

For purposes of this report, inflation for future years is an estimated 2 percent increase, per the Federal Reserve’s inflation targeting policy. The Federal Reserve believes that 2 percent inflation maximizes employment and price stability for the US economy.³

Inflation Rates:

- FY 2023: Not applicable.
- FY 2024: 2.1517 percent (estimated).
- FY 2025: 2 percent (estimated).
- FY 2026: 2 percent (estimated).
- FY 2027: 2 percent (estimated).

Note 5. Rent Costs

GSA charges rent to FDA for the federal buildings that FDA occupies. This rent is charged at different rates depending on the type and location of the space provided. Because rent is an essential support cost for human generic drug activities, a portion of those charges is paid from non-user fee appropriations and a portion is paid from GDUFA fees.

Also included in this account are recurring costs that FDA pays directly to non-federal sources under the delegation of direct lease and service authority. These services include rental of space, and all recurring services for building operations such as overtime utilities, janitorial services, guards, and ground maintenance. The amount of rent and rent related costs each Center pays is directly related to the square footage occupied by that Center.

Section 905(b) of FDARA amended the FD&C Act to provide that the types of fee-coverable costs for the PDUFA, GDUFA, MDUFA, and BSUFA user fee programs will change on October 1, 2023. The statutory definition of allowable cost categories, i.e., the “resources allocated for human generic drug activities” under GDUFA, is what determines the type of expenses related to human generic drug activities that GDUFA fees can be spent on.

Specifically, section 744B of the FD&C Act was amended by FDARA section 905(b) to narrow the “resources allocated for human generic drug activities”, as detailed in section 744A(12)(C) of the FD&C Act, to include only expenditures for leasing and necessary scientific equipment. The impact of the change means that certain costs related to

³ See the following for the Federal Reserve’s 2 percent inflation policy: The Fed - Why does the Federal Reserve aim for inflation of 2 percent over the longer run?
facilities, equipment, materials, and supplies will no longer be able to be funded by GDUFA user fee funds.

Note 6. Shared Service Costs

FDA has several shared service organizations, located within the WCF, that provide support across the user fee programs. The shared service organizations in FY 2022 include:

- **FDA Central**: Provides for Center-wide and Agency-wide services such as telecommunications, training, printing, mail and document management, IT systems, employee health units, and other support and miscellaneous services.
- **Office of Digital Transformation**: Provides the vision and leadership in IT, data, and cybersecurity needed to advance FDA’s mission and strategic priorities.
- **Office of Acquisitions and Grants Services**: Manages contracts, grants, and other agreements.
- **Office of Equal Employment Opportunity**: Promotes an inclusive work environment that ensures equal employment opportunity and fosters a culture that values diversity and empowers individuals.
- **Office of Facilities, Engineering, and Mission Support Services**: Provides FDA employees with office and laboratory facilities.
- **Office of Financial Management**: Provides financial managerial services and policy guidance.
- **Division of Budget Execution and Control**: Initiates, monitors, and analyzes FDA’s budget resources.
- **Office of Finance, Budget, Acquisitions, and Planning**: Leads FDA’s budget, acquisitions, and financial management functions while ensuring the financial integrity of FDA’s resources.
- **Office of Security Operations**: Develops and implements the Agency-wide security policies and programs by providing leadership and guidance to managers and staff on all aspects of security. Administers vital security functions that contribute to the Agency’s mission of protecting public health by enhancing the safety and security of all personnel, facilities, and information.
- **Office of Laboratory Safety**: Reinforces FDA’s expectations for safety and laboratory security, enhances communications among FDA’s safety staff, and provides program support.
- **Office of Ethics and Integrity**: Protects the integrity of FDA’s programs and operations by promoting an ethical culture and ensuring compliance with applicable federal ethics laws.
- **Office of Enterprise Management Services**: Provides strategic and tactical enterprise-wide services through the development and implementation of administrative policies, programs, and initiatives.
- **Office of Human Capital Management**: Provides human resource services that promote collaboration and a work environment that is characterized by diversity, fairness, open communication, personal accountability, trust, and mutual respect.
- **Office of Talent Solutions**: Provides high-quality and efficient human resource solutions that enable FDA to hire a talented and qualified workforce.
- **Office of Planning, Evaluation, and Risk Management**: Partners with FDA’s leaders to achieve organizational excellence by improving program performance, governance, operational efficiency, and risk management.

### Note 7. Operating Reserve Adjustment

GDUFA III amendments to the FD&C Act established an operating reserve adjustment that authorizes FDA to adjust fees for FY 2024 or subsequently during GDUFA III to maintain sufficient operating reserves of carryover user fees. To determine the dollar amounts for the operating reserve thresholds, adjustments for inflation and capacity planning are applied to the base revenue. This amount is then divided by 52 to generate the one-week operating amount. The one-week operating amount is then multiplied by threshold amounts (For FY 2024, the increase threshold is 8 weeks and the decrease threshold is 12 weeks).

FDA may use the operating reserve adjustment to further increase the fee revenue and fees to provide operating reserves of carryover user fees for human generic drug activities for not more than the number of weeks specified: 8 weeks in FY 2024; 9 weeks in FY 2025; and 10 weeks in FY 2026 and FY 2027.

If the estimated carryover balance is in excess of 12 weeks of operating reserves, FDA is required to decrease fees for that fiscal year to reduce the operating reserve to not more than 12 weeks.

Should FDA make an operating reserve adjustment, either up or down, FDA must explain its rationale in the annual fee-setting Federal Register notices.

To calculate what the dollar amounts these estimated thresholds would be, adjustments for inflation and capacity planning are applied to the base revenue. This estimated amount is then divided by 52 to generate the estimated 1-week operating amount. The one-week operating amount is then multiplied by the threshold amount.

- For FY 2024, the base revenue amount is $582,500,000, plus the inflation adjustment estimate of $12,533,653, plus the CPA of an estimated $0, results in an estimated target revenue amount of $595,033,653. This amount is divided by 52 to generate the estimated 1-week operating amount of $11,442,955.

  Taking the 1-week operating amount and multiplying it by 8 produces the estimated 8-week operating reserve discretionary increase threshold amount of $91,543,639. The estimated FY 2024 total carryover amount, end of year is $134,486,874, which exceeds this 8-week increase threshold. As a result, FDA cannot increase GDUFA fee rates using the operating reserve adjustment.

  Multiplying the 1-week operating amount by 12 results in a 12-week decrease threshold amount of $137,315,458. The FY 2024 total carryover amount, end of
year is below this threshold; therefore, FDA is not required to use the operating reserve adjustment to decrease GDUFA fee rates. These values are for estimation purposes only. These values are subject to change during the annual fee setting process.

Note 8. Future Year Refunds Allowance, Set Aside

If an ANDA is considered not to have been received within the meaning of section 505(j)(5)(A) of the FD&C Act for a cause other than failure to pay user fees, or if the ANDA is withdrawn prior to being received within the meaning of section 505(j)(5)(A), the applicant is eligible for a 75 percent refund of the ANDA filing fee. If an ANDA is initially received under section 505(j)(5)(A), but FDA subsequently determines that the exclusivity period for a listed drug should have prevented the ANDA from being received, the ANDA is no longer considered received under section 505(j)(5)(A), and the applicant is eligible for a full refund of the ANDA filing fee paid.

Refunds impact net fee collections for each fiscal year. Net collections reflect the amount of fees collected net any refunds or adjustments that occurred during that fiscal year.

Note 9. Minimum Non-User Fee Appropriations Adjustment Factor

FDA must calculate and incorporate an adjustment factor (defined in section 744A(3) of the FD&C Act. This FD&C Act provision states, “the term ‘adjustment factor’ means a factor applicable to a fiscal year that is the Consumer Price Index for all urban consumers (all items; United States city average) for October of the preceding fiscal year divided by such Index for October 2011.”

Note 10. Capacity Planning Adjustment

GDUFA III amendments to the FD&C Act established a capacity planning adjustment that authorizes FDA to adjust the annual target revenue amount, within certain parameters, to account for sustained increases in workload. This adjustment helps ensure that FDA can expand its review capacity to meet additional workload demands and maintain performance on its review timelines.
This report was prepared by FDA’s Office of Financial Management.
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