



**FDA** U.S. FOOD & DRUG  
ADMINISTRATION

# **FY 2020**

## ***PERFORMANCE REPORT TO CONGRESS***

*for the*

## ***Animal Generic Drug User Fee Act***



## ***Commissioner's Report***

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I am pleased to present to Congress the Food and Drug Administration's (FDA's or the Agency's) fiscal year (FY) 2020 Animal Generic Drug User Fee Act (AGDUFA) performance report. FY 2020 marks the 12<sup>th</sup> year of AGDUFA, and this report covers the second year of the second reauthorization of AGDUFA, referred to as AGDUFA III (which authorized animal generic drug user fees from FY 2019 through FY 2023).

This report details FDA's preliminary performance for FY 2020 and finalizes FDA's performance results for FY 2019. It is my pleasure to report that FDA exceeded all performance goals for FY 2019. The Agency also met performance goals for all FY 2020 cohort submissions reviewed or due for review by September 30, 2020. With some reviews still pending, FDA has the potential to exceed all performance goals for FY 2020.

FDA is committed to improving the efficiency, quality, and predictability of the generic new animal drug review process. The timely approval of generic animal drugs continues to be a critical component of animal health because it provides access to additional sources of animal drugs for ranchers, farmers, and pet owners. Since AGDUFA was enacted, FDA has dramatically reduced average review times from 700 days to less than 270 days. Under the leadership of the President, and in collaboration with Congress and industry, FDA looks forward to continued success in the generic new animal drug review program.

Janet Woodcock, M.D.  
Acting Commissioner of Food and Drugs

## ***Acronyms***

**AGDUFA** – Animal Generic Drug User Fee Act  
**ANADA** – Abbreviated New Animal Drug Application  
**CVM** – Center for Veterinary Medicine  
**FDA** – Food and Drug Administration  
**FD&C Act** – Federal Food, Drug, and Cosmetic Act  
**FY** – Fiscal Year (October 1 to September 30)  
**JINAD** – Generic Investigational New Animal Drug  
**ONADE** – Office of New Animal Drug Evaluation  
**PAI** – Pre-Approval Inspection

## ***Executive Summary***

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On August 14, 2018, the second reauthorization of the Animal Generic Drug User Fee Act (AGDUFA), referred to as AGDUFA III, was signed into law, providing an additional 5 years (from fiscal year (FY) 2019 to FY 2023) of user fees for the generic new animal drug review program. The AGDUFA III program includes a comprehensive set of Food and Drug Administration (FDA) review performance goals and commitments designed to improve the timeliness and predictability of the review of abbreviated new animal drug applications (ANADAs) and reactivations, manufacturing supplemental ANADAs and reactivations, and generic investigational new animal drug submissions. The reauthorization also dramatically reduces review time goals across all submission types.

More information on the history of AGDUFA is available on FDA's AGDUFA website.<sup>1</sup>

### **Information Included in this Report**

This report summarizes FDA's performance results in meeting AGDUFA goals and commitments for FY 2019 and FY 2020. Specifically, it updates and finalizes performance data initially reported in the FY 2019 AGDUFA Performance Report and presents preliminary data on FDA's progress in meeting FY 2020 review goals, implementation activities, and accomplishments.

### **Review Performance**

FDA met or exceeded the expectations of the review performance goals in the first year of AGDUFA III and continued to meet or exceed the expectations of the review performance goals for FY 2020. Key activities and accomplishments during FY 2020 included the following:

- FDA met review-time goals for almost all (540 of 542) FY 2019 submissions. FDA exceeded all (5 of 5) AGDUFA performance goals for the FY 2019 cohort. Please see Appendix A for more details on the submission types and related performance goals.
- Preliminary performance results indicate that FDA met review-time goals for almost all (291 of 299) FY 2020 cohort submissions reviewed and acted on as of September 30, 2020. With 268 additional reviews pending that may yet be completed on time, FDA has the potential to exceed all five AGDUFA performance goals for the FY 2020 cohort. Please see Appendix A for more details on the submission types and related performance goals.

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<sup>1</sup> [www.fda.gov/ForIndustry/UserFees/AnimalGenericDrugUserFeeActAGDUFA/default.htm](http://www.fda.gov/ForIndustry/UserFees/AnimalGenericDrugUserFeeActAGDUFA/default.htm).

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## Introduction

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The Animal Generic Drug User Fee Act (AGDUFA) requires the Secretary of Health and Human Services to submit two annual reports to Congress: (1) a performance report and (2) a financial report. The fiscal year (FY) 2020 report is the Food and Drug Administration's (FDA's or the Agency's) second annual performance report to Congress under the second reauthorization of AGDUFA, referred to as AGDUFA III. Under AGDUFA III, FDA agreed to meet review performance goals for certain submissions over a 5-year period (FY 2019 through FY 2023). Further details on FDA's commitments under AGDUFA III can be found in the AGDUFA III Performance Goals Letter on FDA's website.<sup>2</sup>

AGDUFA is designed to bring greater predictability in review times for the generic animal drug industry by providing FDA with supplemental funding for the review of generic new animal drug submissions. AGDUFA accelerates the availability of safe and effective generic new animal drug products. The guidelines and definitions below apply to the information provided in the FY 2020 report.

### Information Presented in This Report

In any given year, FDA's performance includes reviews of applications and submissions pending from previous fiscal years, along with submissions received during the current fiscal year. This report presents FDA's final performance results for the FY 2019 cohort and presents FDA's preliminary performance results with respect to performance goals for the FY 2020 cohort submissions that were received early enough to be reviewed or due for review by September 30, 2020.

The following information refers to FDA's performance results presented in this report:

- The term *submission* is used to refer to abbreviated new animal drug applications (ANADAs) and reactivations, supplemental ANADAs and reactivations, generic investigational new animal drug (JINAD) studies, and JINAD protocols when referencing the fiscal year cohort.
- *Review-time goal* is the targeted time period, identified in number of calendar days, within which individual submissions are to be acted on by FDA. AGDUFA review-time goals range from 60 days to 270 days. An *on-time review* indicates that FDA completed action within the number of calendar days specified by the review-time goal.
- *Percent on time* refers to the percentage of reviews where FDA met a review-time goal for a given type of submission. FDA's percent on time for a given type of submission is used to determine whether FDA met or exceeded the AGDUFA III performance goals.
- *Performance goal* refers to the percentage of total submissions, agreed to under AGDUFA III, where FDA is expected to meet the review-time goal for a given type of submission. The AGDUFA III performance goals call for FDA to meet the review-time goals 90 percent of the time for the defined fiscal year cohort.

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<sup>2</sup> [www.fda.gov/ForIndustry/UserFees/AnimalGenericDrugUserFeeActAGDUFA/default.htm](https://www.fda.gov/ForIndustry/UserFees/AnimalGenericDrugUserFeeActAGDUFA/default.htm).

- The performance statistics in this report are based on submissions received during a fiscal year (known as a receipt cohort). This methodology calculates performance statistics for submissions according to the fiscal year FDA received them, regardless of the year in which FDA ultimately acted on the submissions. A result of this methodology is that the statistics shown for a particular fiscal year may change from one report to the next. As more submissions are completed, the statistics for that year of receipt are adjusted to reflect the new completions. Therefore, until all submissions in a cohort are acted on or have passed the due date, whichever comes first, only a preliminary performance assessment is provided for that fiscal year cohort.
- For submission types with a longer review-time goal (for example, 270 days), review performance data are usually limited. For those submissions with a shorter review-time goal (for example, 60 days), review performance data for submissions received early in the fiscal year are available at the time the report is prepared, and thus the report may provide an early indicator of review performance.
- Performance goal tables indicate the total number of submissions filed as well as whether the submission was reviewed on time, was overdue, or is still pending and not past its due date.
- The workload count presented in this report for FY 2020 includes all submissions received in FY 2020. For AGDUFA review times, FDA calculates from the original receipt of the application or submission.
- When determining performance, FDA-calculated percentages are rounded to the nearest whole number up to 99 percent. Percentages above 99 percent, but below 100 percent, are rounded down to 99 percent.

#### **File Types Included in This Report**

- **ANADA** – An ANADA is an abbreviated new animal drug application including all reactivations and supplements. This report presents original applications and reactivations, administrative applications, and supplemental applications and reactivations as separate goals.
- **JINAD file** – The generic investigational new animal drug file is the investigational file for generic animal drugs. The information submitted to the file may be used to support an ANADA. This report presents study submissions and protocols.

**Source:**

<https://www.fda.gov/animal-veterinary/development-approval-process/new-animal-drug-applications>

## ***AGDUFA Review Workload***

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### **Review Workload: FY 2015 to FY 2020**

In the table below, preliminary review workload numbers from FY 2020 are compared to the previous 5-year averages for all AGDUFA application and submission types filed. The individual years that are included in the 5-year average can also be referenced below. There are no performance goals associated with workload, but the variations in workload over time can provide context for FDA's performance. Workload for four application and submission types showed an increase in FY 2020 from the 5-year average, while one flatlined. Please see Appendix A for more details on the application and submission types included in the table below.

**Review Workload for Applications and Submissions**

<b>Application/Submission Type</b>	<b>FY 15</b>	<b>FY 16</b>	<b>FY 17</b>	<b>FY 18</b>	<b>FY 19</b>	<b>FY 20<sup>†</sup></b>	<b>FY 15 to FY 19 5-Year Average</b>	<b>FY 20 Compared to 5-Year Average</b>
Original ANADAs and Reactivations	22	16	17	19	33	21	21	0%
Administrative ANADAs	1	1	4	3	3	10	2	+400%
Manufacturing Supplemental ANADAs and Reactivations	152	156	173	180	270*	308	186	+66%
JINAD Studies	54	63	66	97	153*	150	87	+72%
JINAD Protocols	12	22	48	40	83	78	41	+90%

\* Numbers were changed to reflect updates to data presented in the FY 2019 AGDUFA Performance Report.

<sup>†</sup> FY 2020 numbers are preliminary and will be updated in the FY 2021 AGDUFA Performance Report.

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## ***FY 2019 and FY 2020 AGDUFA Performance Results***

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The tables that follow present FDA's review performance results for the FY 2019 and FY 2020 AGDUFA cohort submissions.

### **Final FY 2019 Performance Results**

FDA exceeded the 90 percent performance level for all five of the review performance goals for the FY 2019 cohort. Across all submission types, FDA met the review-time goal in 540 of 542 submissions. The entire FY 2019 cohort has closed; therefore, there are no pending submissions. Please see Appendix A for more details on the submission types in the table below and the performance goals.

<b>Submission Type</b>	<b>Filed</b>	<b>On Time</b>	<b>Overdue</b>	<b>Percent on Time</b>
Original ANADAs and Reactivations	33	33	0	100%
Administrative ANADAs	3	3	0	100%
Manufacturing Supplemental ANADAs and Reactivations	270*	268	2	99%
JINAD Studies	153*	153	0	100%
JINAD Protocols	83	83	0	100%

\* Numbers were changed to reflect updates to data presented in the FY 2019 AGDUFA Performance Report

## Preliminary FY 2020 Performance Results

As of September 30, 2020, preliminary performance data was available for 299 of 567 submissions filed in FY 2020. FDA is currently exceeding performance goals for all five submission types. Overall, FDA met review-time goals for 291 of 299 submissions acted on. With 268 submissions pending within goal, FDA has the potential to meet or exceed the 90 percent performance goal for all five submission types. Please see Appendix A for more details on the submission types in the table below and the performance goals.

Submission Type	Filed	On Time	Overdue	Pending Within Goal	Pending Overdue	Percent on Time
Original ANADAs and Reactivations	21	7	0	14	0	100%
Administrative ANADAs	10	9	0	1	0	100%
Manufacturing Supplemental ANADAs and Reactivations	308	134	1	173	0	99%
JINAD Studies	150	87	4	59	0	96%
JINAD Protocols	78	54	3	21	0	95%

## **FY 2020 Process Improvements and Major Accomplishments**

Under AGDUFA III, FDA committed to a variety of process improvements. FDA agreed to continue to enhance and further improve its review process via the following goals and procedures:

- **Foreign Pre-Approval Inspections (PAIs).** Continuing under AGDUFA III, to improve the timeliness and predictability of foreign PAIs, the regulated industry may voluntarily submit, at the beginning of the calendar year, a list of foreign manufacturing facilities (1) that are specified in an ANADA, supplemental ANADA, or JINAD file and (2) that may be subject to foreign PAIs. Due to staff travel restrictions implemented by FDA in response to the COVID-19 pandemic,<sup>3</sup> fewer foreign inspections than anticipated were completed in FY 2020. The table below shows the number of foreign PAIs conducted and the average time it took to complete a PAI during each fiscal year.

<b>Fiscal Year</b>	<b>Number of Foreign PAIs Conducted</b>	<b>Average Time to Completion (in Days)</b>
<b>2019</b>	5	145
<b>2020</b>	1	135
<b>2021</b>	N/A	N/A
<b>2022</b>	N/A	N/A
<b>2023</b>	N/A	N/A

- **Review Times.** The Agency agreed to develop a shortened review-time process for certain ANADA and JINAD submissions (see Appendix A).
- **Multiple Data Submissions to the Chemistry, Manufacturing, and Controls Technical Section.** The Agency continued to allow two-phased Chemistry, Manufacturing, and Controls technical section submissions under the JINAD process.

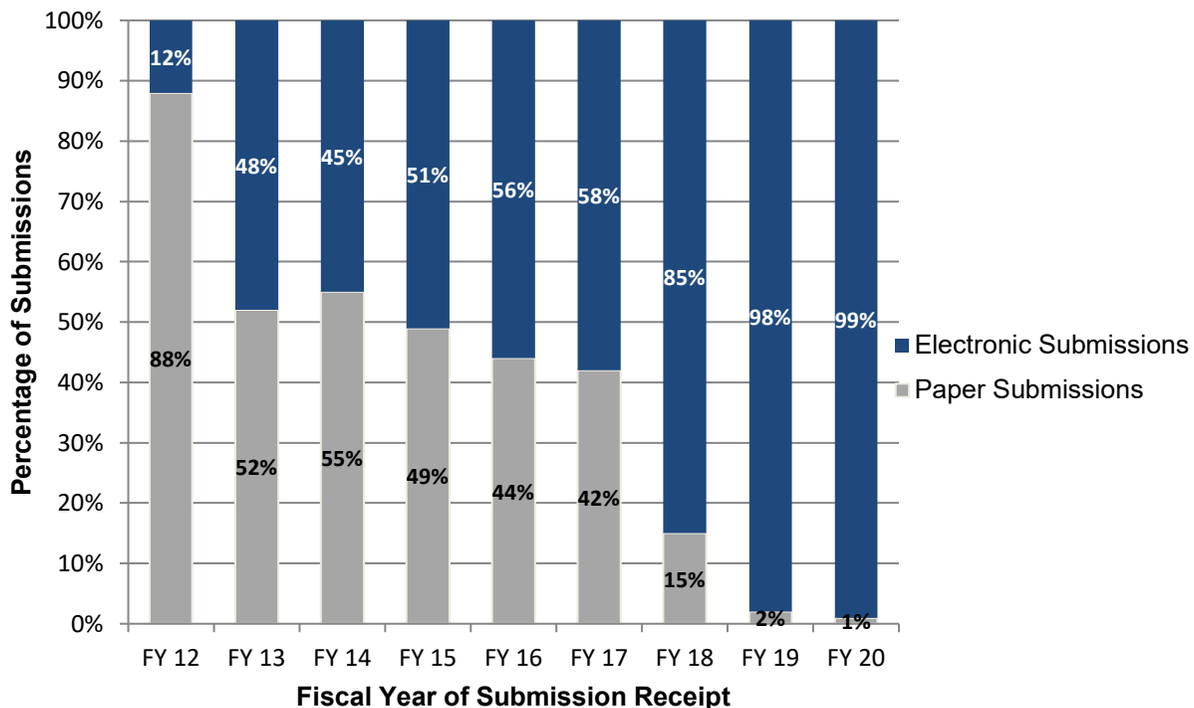
<sup>3</sup> [www.fda.gov/news-events/press-announcements/coronavirus-disease-2019-covid-19-update-foreign-inspections](https://www.fda.gov/news-events/press-announcements/coronavirus-disease-2019-covid-19-update-foreign-inspections).

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## FY 2020 Additional Activities Toward Compliance with AGDUFA III

- Section 301 of the Animal Drug and Animal Generic Drug User Fee Amendments of 2018. Electronic submissions.** Beginning October 1, 2018, all applications and submissions under sections 512(b) and 571(a) of the Federal Food, Drug, and Cosmetic Act (FD&C Act) must be created using the eSubmitter tool and submitted to the Agency through the Center for Veterinary Medicine’s (CVM’s) Electronic Submission System.
  - Accomplishment:** CVM provided support to industry users to facilitate their transition to using eSubmitter for all submissions to CVM. A small percentage of applications and submissions were submitted in paper.

**Percent of Electronic vs. Paper Submissions Received by FDA  
FY 2012- 2020**



- Section 303. Misbranded drugs and devices.** This section of the reauthorization legislation mandates that, with limited exceptions, pioneer and generic new animal drugs approved under section 512 of the FD&C Act must include a specific statement (i.e., “Approved by FDA under (A)NADA #”), followed by the application number on the sponsor’s labeling, by September 30, 2023, or else such drugs will be considered misbranded under section 502(w) of the FD&C Act.
  - Accomplishment:** CVM continued to remind and encourage sponsors of pioneer and generic approved new animal drugs to update their products’ labeling with the new statement. By the end of FY 2020, the labeling of

approximately one-third of such approved and marketed products were in compliance.

## Appendix

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### Appendix A: AGDUFA Performance Goals

The table below shows the performance goals for AGDUFA II (FY 2014 to FY 2018).

Submission Type	Performance Goal: Act on 90 Percent Within
<b>Original ANADAs and Reactivations</b>	
Original ANADAs	270 days
Original ANADAs Reactivations	270 days
Shortened Review Original ANADA Reactivations	190 days
<b>Administrative ANADAs</b>	100 days
<b>Manufacturing Supplemental ANADAs and Reactivations</b>	
Manufacturing Supplements and Reactivations (Prior Approval)	270 days
Manufacturing Supplements and Reactivations (Changes Being Effected)	270 days
<b>JINAD Study Submissions</b>	
JINAD Data Submissions	270 days
JINAD Data Resubmissions	270 days
Shortened Review JINAD Data Resubmissions	90 days
<b>JINAD Protocol Submissions</b>	100 days

The table below shows the performance goals for AGDUFA III (FY 2019 to FY 2023).

<b>Submission Type</b>	<b>Performance Goal: Act on 90 Percent Within</b>
<b>Original ANADAs and Reactivations</b>	
Original ANADAs	240 days
Original ANADAs Reactivations	240 days
Shortened Review Original ANADAs Reactivations	120 days
<b>Administrative ANADAs</b>	60 days
<b>Manufacturing Supplemental ANADAs and Reactivations</b>	
Manufacturing Supplements and Reactivations (Prior Approval)	180 days
Manufacturing Supplements and Reactivations (Changes Being Effected)	270 days
<b>JINAD Study Submissions</b>	
JINAD Data Submissions	180 days
JINAD Data Resubmissions	180 days
Shortened Review JINAD Data submissions	60 days
<b>JINAD Protocol Submissions</b>	75 days





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Food and Drug Administration**

This report was prepared by FDA's Office of Planning in collaboration with the Center for Veterinary Medicine. For information on obtaining additional copies, contact:

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