



Medicare Part B Average Sales Price (ASP) Module

Submitter User Guide

Version 2.1

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1. Purpose

The purpose of this user guide is to provide guidance and instructions to representatives of drug manufacturing companies as they submit federally required Medicare Part B drug Average Sales Price (ASP) data to the Centers for Medicare & Medicaid Services (CMS).

CMS uses the Fee-for-Service Data Collection System (FFSDCS) to house various Fee-for-Service modules.

The ASP Data Collection System, referred to within this user guide as the ASP Module, is one of the modules under the FFSDCS system, and offers the following:

- Provides users with an online-based software application for automating the collection, editing, and processing of drug product pricing data drug manufacturers submit on a quarterly basis.
- Establishes a relationship between the manufacturers' reported data and the billing codes Medicare providers use to calculate a weighted average sales price for each billing code.
- Establishes prices for billing codes to determine payment limits of Part B drugs on certain Medicare claims.
- Eliminates data entry errors, data formatting errors, and incomplete submitted data, and greatly reduces the process cycle and resource time needed to provide the pricing to contractors through automation of the manually intensive processes.
- Accepts, stores, validates, and calculates drug pricing on Medicare Part B drug data received for the Center for Medicare Management (CMM) stakeholders.

Section 303 (b) and (c) of the [Medicare Modernization Act \(MMA\) of 2003](#) revised the payment methodology for the majority of Part B covered drugs and biologicals that are not priced on a cost or prospective payment basis (hereafter referred to as drugs).

CMS applies the ASP methodology to the data drug manufacturers have submitted to the ASP Module. Per the MMA, ASP methodology determines the payment limit for these drugs. Local contractors calculate pricing for compounded drugs.

2. Logging in Using MFA

First time users must register and create an account in the [CMS Enterprise Portal](#) before logging in to the ASP Module. Refer to the Resource Library on the [Education and Outreach page](#) to view the ASP Module Registration User Guide for registration steps.

Once registration is complete, follow these steps to log in to the ASP Module as a Submitter using Multi-Factor Authentication (MFA):

1. Navigate to the [CMS Enterprise Portal](#) main page.

The ASP Module Login Page opens. Refer to *Figure 1*.

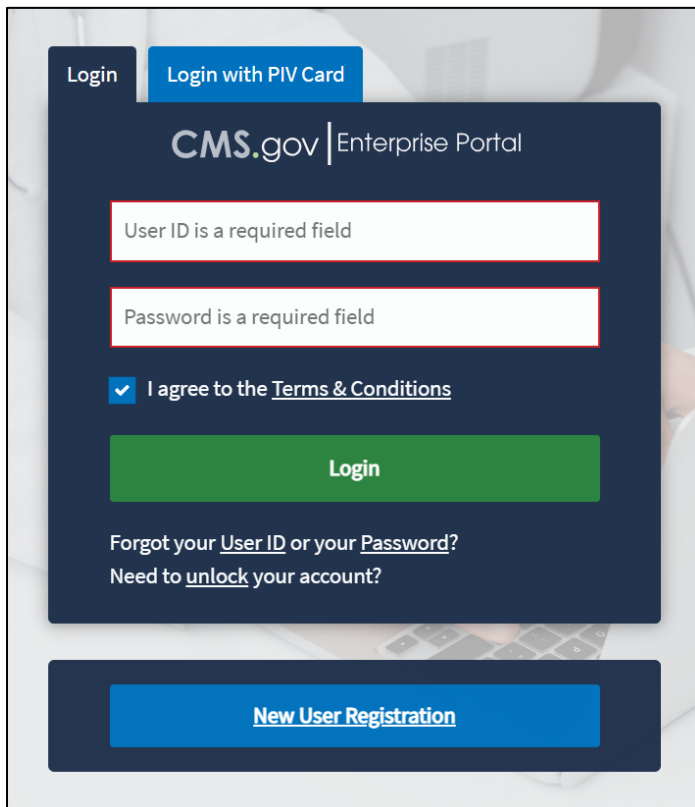


Figure 1: Logging in Using MFA - ASP Module Login

2. Enter your login information in the required **User ID** and **Password** fields.
3. Click the **Terms & Conditions** hyperlink and review the text in the pop-up window; close the window to move on to the next step.
4. Review the terms and conditions and select the **I agree to the Terms & Conditions** checkbox.

Note: By selecting this checkbox, you certify that you read and consent to monitoring while accessing and using the ASP Module. The terms and conditions link provides additional hyperlinks to the HHS Rules of Behavior and the CMS Privacy Act Statement.

5. Click **Login**.

Note: If you forget your user ID or password, click the **Forgot your User ID or your Password?** hyperlink under the **Login** button and follow the provided instructions. If you still cannot access your account and need to unlock it, click the **Need to unlock your account?** hyperlink under **Login** button.

The **Multi-Factor Authentication** page opens. Refer to *Figure 2*.

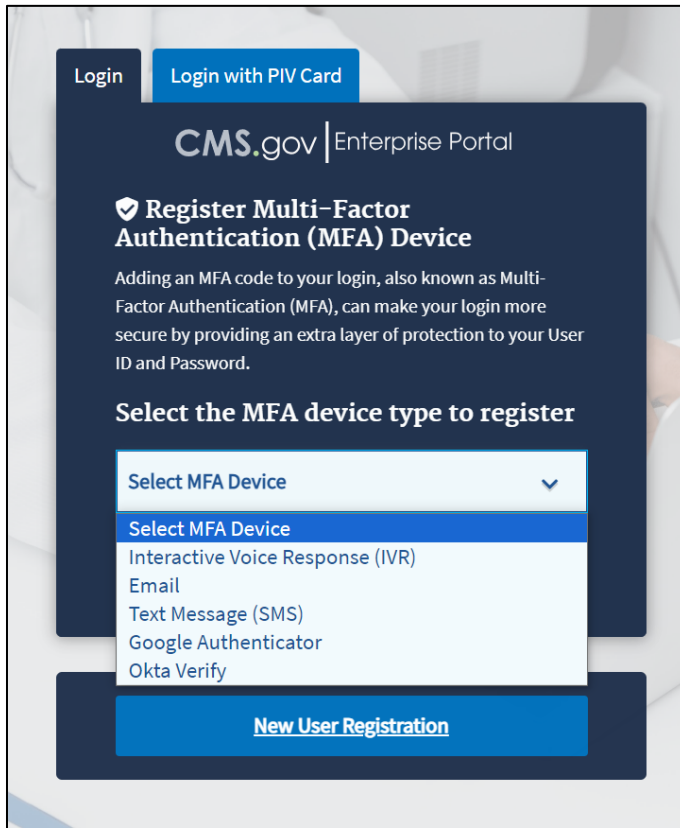


Figure 2: Logging in Using MFA - Select MFA Device Type Drop-Down Menu

To ensure the security of high value data submitted to the ASP Module, you must authenticate your identity using an MFA process. The first time you attempt to log in, you must choose an authentication method. Users have various authentication options, including Interactive Voice Response (IVR), Email, Text Message (Short Message Service (SMS)), Google Authenticator and Okta Verify.

6. Click the **Select MFA Device** drop-down menu; select your preferred MFA device type from the list. Whenever you log back into the ASP Module through this process, your preferred method of MFA reloads automatically.

Note: Figure 3 demonstrates MFA registration using IVR as the selected option.

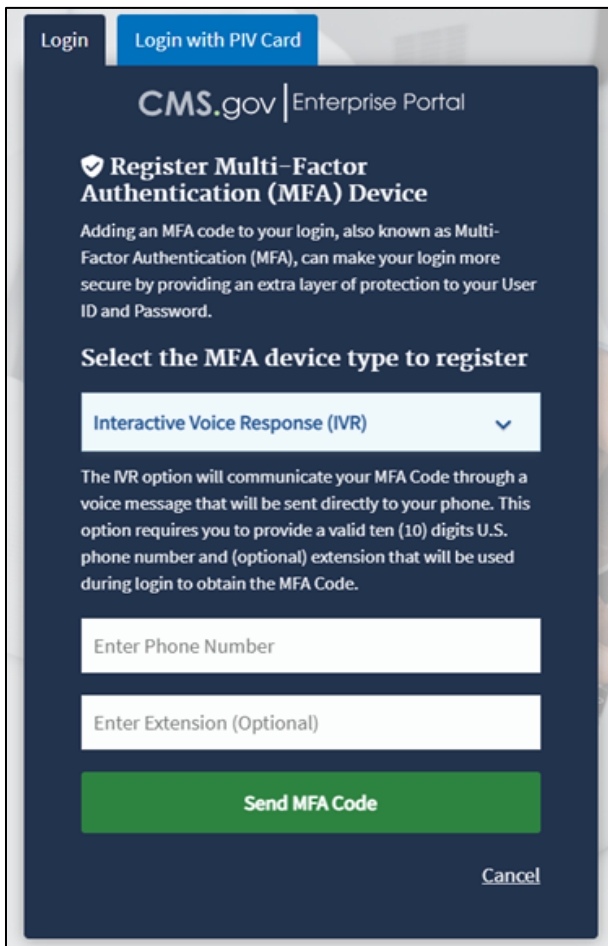


Figure 3: Logging in Using MFA - Multi-Factor Authentication - (IVR) Example

7. Enter your phone number in the **Phone Number** field; enter your extension in the **Extension** field, if necessary.
8. Click the **Send MFA Code** button to receive a six-digit code via your chosen contact method.
9. Record and enter the six-digit code you received into the **Enter MFA Code** field. Refer to *Figure 4*.

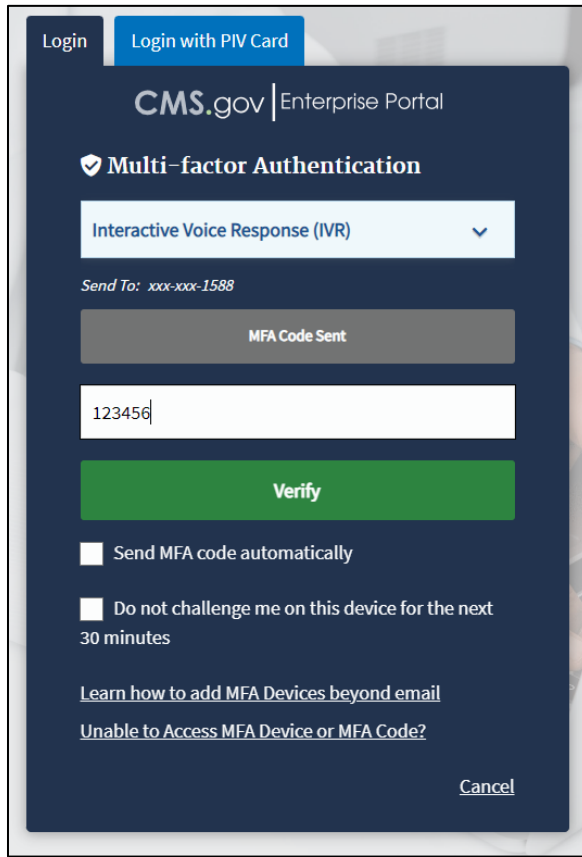


Figure 4: Logging in Using MFA - Multi-Factor Authentication - Verify MFA Code

10. Check the **Send MFA code automatically** and **Do not challenge me on this device for the next 30 minutes** checkboxes depending on your preference.

Note: If you need help, click the [Learn how to add MFA Devices beyond email](#) and [Unable to Access MFA Devices or MFA Code?](#) hyperlinks.

11. Click the **Verify** button to confirm your identity and enter the ASP Module homepage.

The **My Portal** landing page opens. Refer to *Figure 5*.



Figure 5: Logging in Using MFA - My Portal Landing Page

Note: Other CMS applications you have access to may display on the **My Portal** landing page.

12. Click the **Fee For Service Data Collection System (FFSDCS)** box.

A Fee for Service Data Collection System (FFSDCS) drop-down menu displays. Refer to *Figure 6*.

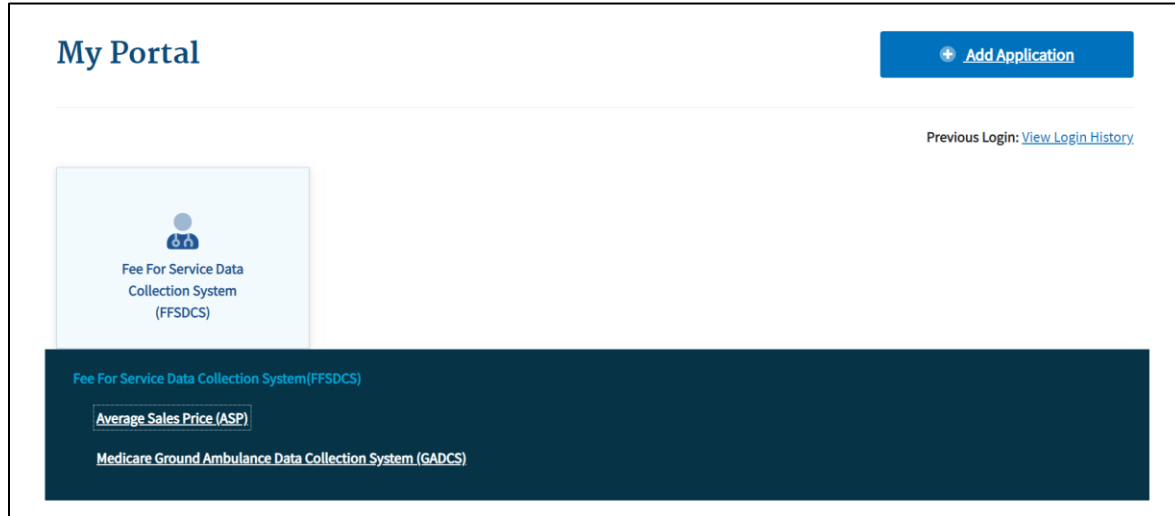


Figure 6: Logging in Using MFA - My Portal Landing Page - FFSDCS Drop-down Menu

13. Click the **Average Sales Price (ASP)** hyperlink.

A full-page statement displays, titled **ASP Data for Drugs and Biologics Covered Under Medicare Part B**. The statement details recent statutory requirements stated in the Social Security Act (the Act), and the Consolidated Appropriations Act (CAA), 2021. These requirements hold that manufacturers must report their ASP data to CMS with precision on a quarterly basis without errors or miscalculations. Refer to *Figure 7*.

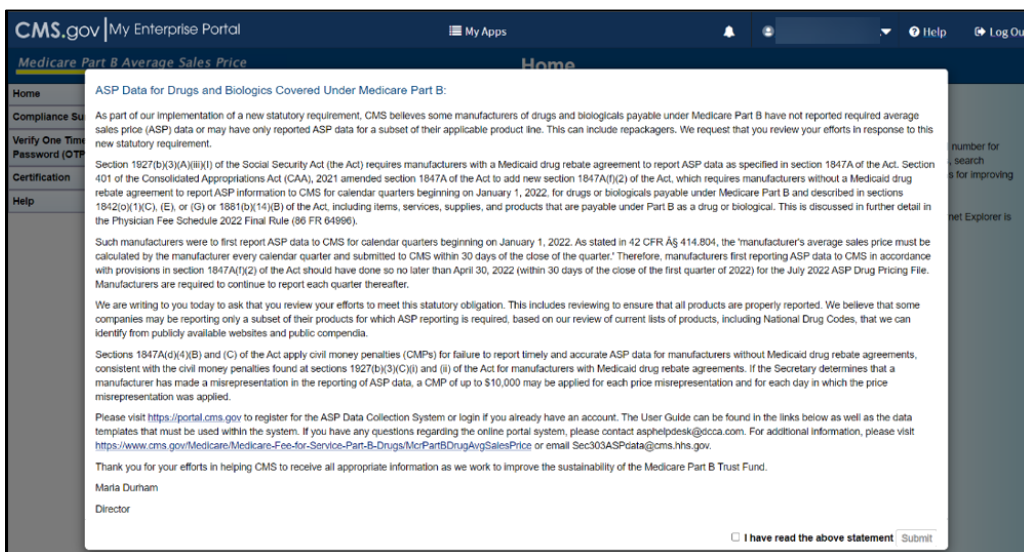


Figure 7: Logging in Using MFA - ASP Data for Drugs and Biologics Under Medicare Part B

14. Read the statement; select the **I have read the above statement** checkbox and click **Submit**.

The Medicare Part B Average Sales Price homepage opens. Refer to *Figure 8*.

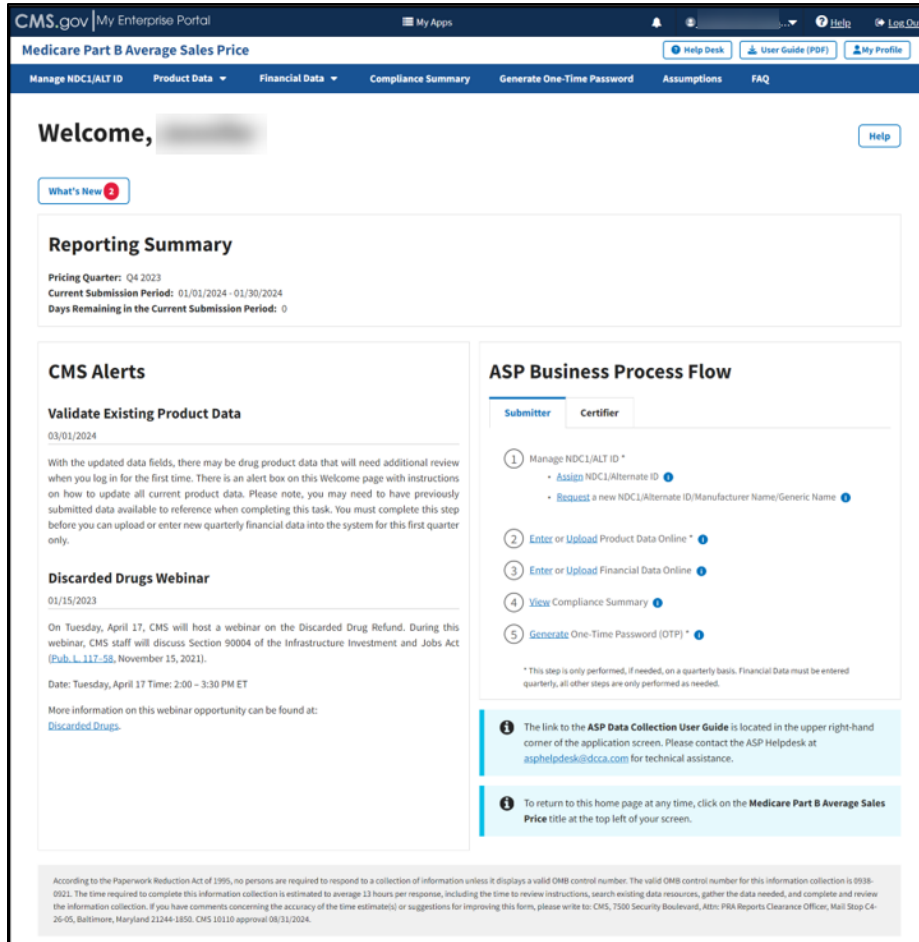


Figure 8: Medicare Part B Average Sales Price Homepage

3. ASP Homepage Menu Tabs

The following sections describe the functionality of each menu tab on the ASP homepage, including **Manage NDC1/ALT ID**, **Product Data**, **Financial Data**, **Compliance Summary**, **Generate One-Time Password**, **Assumptions** and **FAQ**.

Note: This user guide is written in order of the system menu tabs and the respective tasks completed on that page and not necessarily in chronological order of steps to follow for quarterly data submission.

3.1 Manage NDC1/ALT ID

The following sections describe how to assign NDC1s and Alternate IDs, as well as how to request a new NDC1, Alternate ID, and manufacturer or generic name.

Note: To add a new product, users must first request to add an NDC1/ALT ID and Manufacturer/Generic Name, if needed. (Refer to *Section 3.1.3 - Request New NDC1/ALT ID/Manufacturer/Generic Name*). Once the new product has been approved into the system, users can establish the relationship between the manufacturer and the product by assigning the NDC1/ALT ID to the manufacturer. (Refer to *Section 3.1.1 - Assign by NDC1* and *Section 3.1.2 - Assign by Alternate ID*.)

3.1.1 Assign by NDC1

Follow these steps to assign NDC1s:

1. From the Medicare Part B Average Sales Price homepage, click the **Manage NDC1/ALT ID** tab.

The **Manage NDC1/ALT ID** page opens and displays the **Assign NDC1/ALT ID** tab by default. Refer to *Figure 9*.

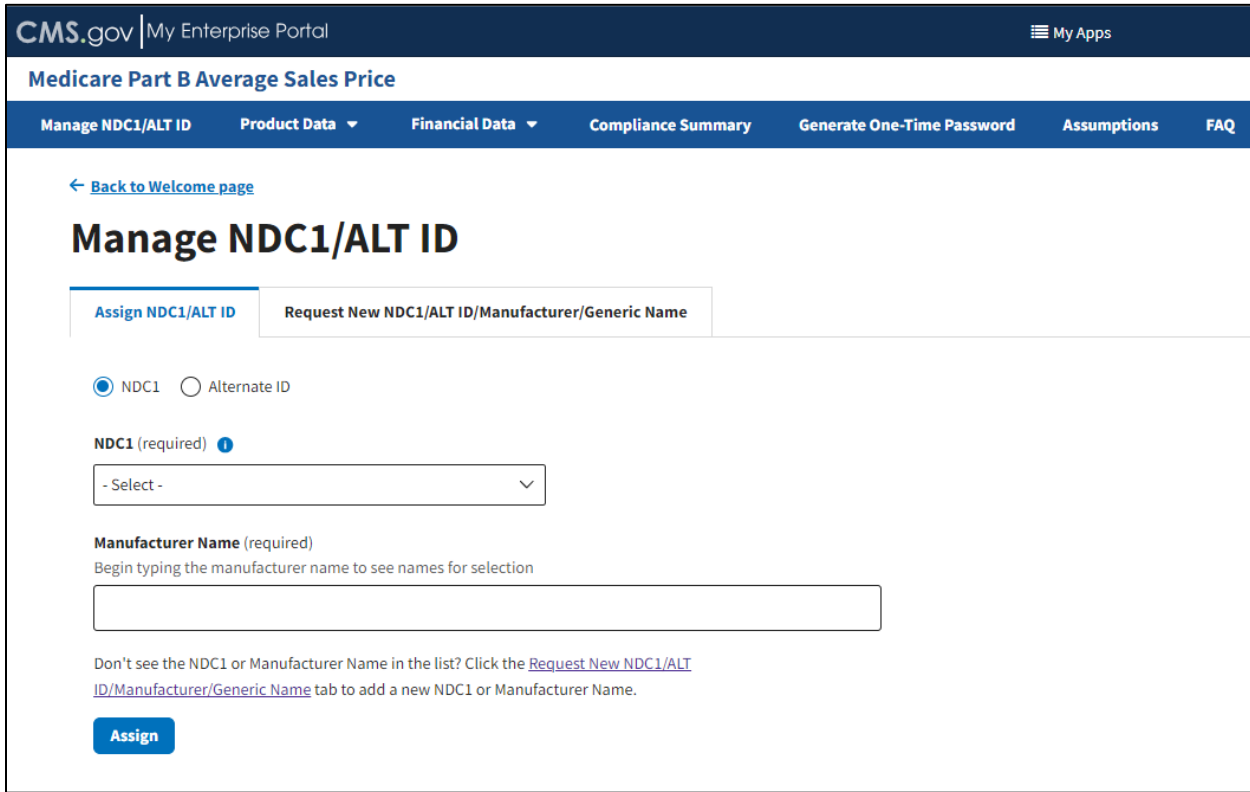
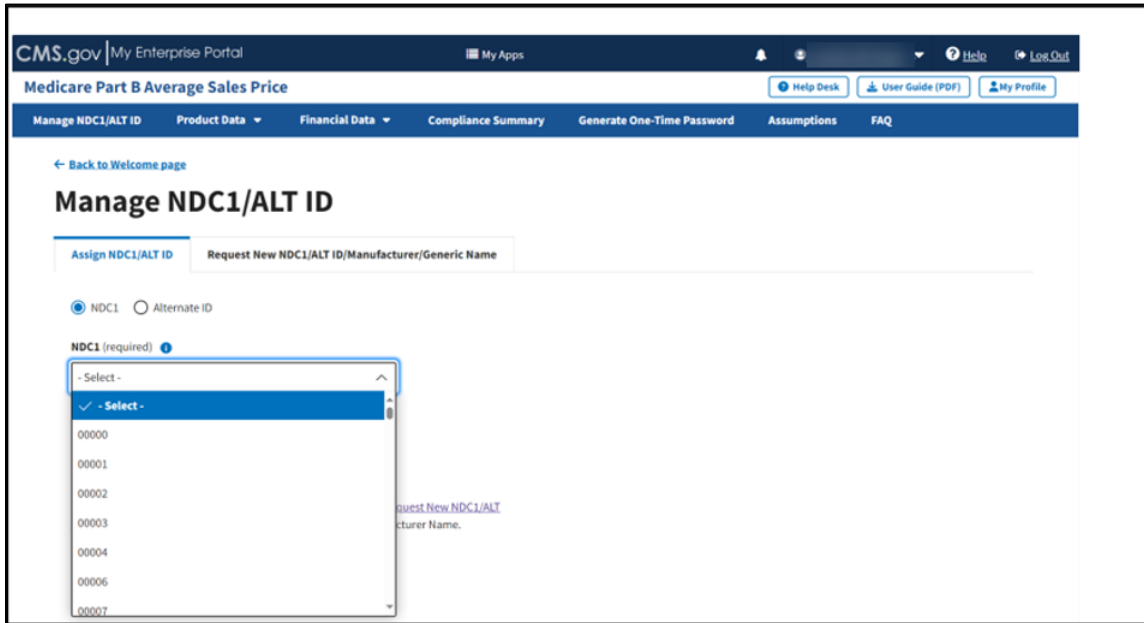

 A screenshot of the CMS.gov Manage NDC1/ALT ID page. The page has a dark blue header with the CMS logo and "My Enterprise Portal" on the left, and "My Apps" on the right. Below the header is a navigation bar with tabs: "Manage NDC1/ALT ID" (which is active and highlighted in blue), "Product Data", "Financial Data", "Compliance Summary", "Generate One-Time Password", "Assumptions", and "FAQ". Below the navigation bar is a back-link "← Back to Welcome page". The main content area has a title "Manage NDC1/ALT ID" and two tabs: "Assign NDC1/ALT ID" (selected) and "Request New NDC1/ALT ID/Manufacturer/Generic Name". Under the "Assign NDC1/ALT ID" tab, there are two radio buttons: "NDC1" (selected) and "Alternate ID". A required field "NDC1 (required)" with a blue info icon has a dropdown menu with the option "- Select -". A "Manufacturer Name (required)" field with a blue info icon has a text input field and placeholder text "Begin typing the manufacturer name to see names for selection". A note at the bottom left says "Don't see the NDC1 or Manufacturer Name in the list? Click the [Request New NDC1/ALT ID/Manufacturer/Generic Name](#) tab to add a new NDC1 or Manufacturer Name." A blue "Assign" button is at the bottom right.

Figure 9: Manage NDC1/ALT ID Page - Assign NDC1

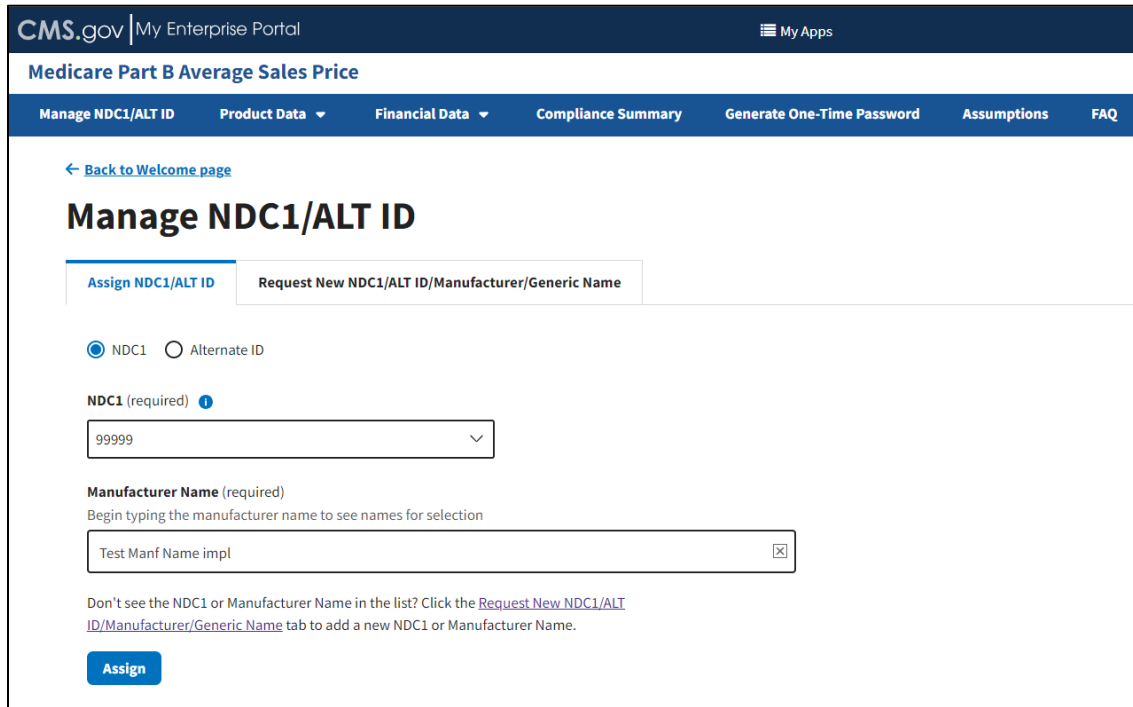
2. From the **Assign NDC1/ALT ID** tab, select the **NDC1** radio button to specify the product data you need to submit to the Module.
3. Under **NDC1 (required)**, click the **-Select-** drop-down menu to expand the list of submitted drugs and additional products in the Module to date; select the appropriate NDC1. Refer to *Figure 10*.



The screenshot shows the CMS.gov My Enterprise Portal interface. The top navigation bar includes 'My Apps', 'Help', 'Log Out', 'Help Desk', 'User Guide (PDF)', and 'My Profile'. The main menu has options for 'Manage NDC1/ALT ID', 'Product Data', 'Financial Data', 'Compliance Summary', 'Generate One-Time Password', 'Assumptions', and 'FAQ'. Below the menu, a link to 'Back to Welcome page' is present. The main content area is titled 'Manage NDC1/ALT ID'. It has two tabs: 'Assign NDC1/ALT ID' (selected) and 'Request New NDC1/ALT ID/Manufacturer/Generic Name'. A dropdown menu is open under the 'Assign NDC1/ALT ID' tab, showing a list of NDC1 codes from 00000 to 00007. The 'Select' option is highlighted.

Figure 10: Manage NDC1/ALT ID Page - Assign NDC1 Drop-down Menu

3. Under **Manufacturer Name (required)**, type and select the appropriate manufacturer. Refer to *Figure 11*.



The screenshot shows the CMS.gov My Enterprise Portal interface. The top navigation bar includes 'My Apps', 'Help', 'Log Out', 'Help Desk', 'User Guide (PDF)', and 'My Profile'. The main menu has options for 'Manage NDC1/ALT ID', 'Product Data', 'Financial Data', 'Compliance Summary', 'Generate One-Time Password', 'Assumptions', and 'FAQ'. Below the menu, a link to 'Back to Welcome page' is present. The main content area is titled 'Manage NDC1/ALT ID'. It has two tabs: 'Assign NDC1/ALT ID' (selected) and 'Request New NDC1/ALT ID/Manufacturer/Generic Name'. A dropdown menu is open under the 'Assign NDC1/ALT ID' tab, showing the value '99999'. Below the dropdown, a 'Manufacturer Name (required)' field contains the text 'Test Manf Name impl'. A note at the bottom of the page says: 'Don't see the NDC1 or Manufacturer Name in the list? Click the Request New NDC1/ALT ID/Manufacturer/Generic Name tab to add a new NDC1 or Manufacturer Name.' A 'Assign' button is located at the bottom of the page.

Figure 11: Manage NDC1/ALT ID Page - Enter NDC1 Manufacturer Name

4. Click **Assign** to submit your information.

A message displays confirming you have successfully added your selections. Refer to *Figure 12*.

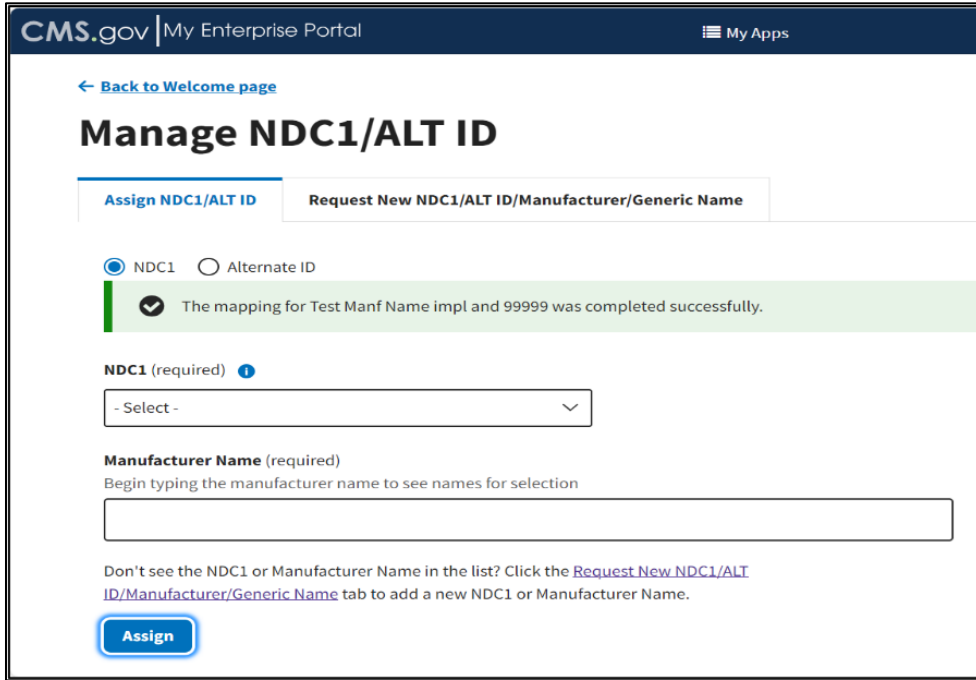
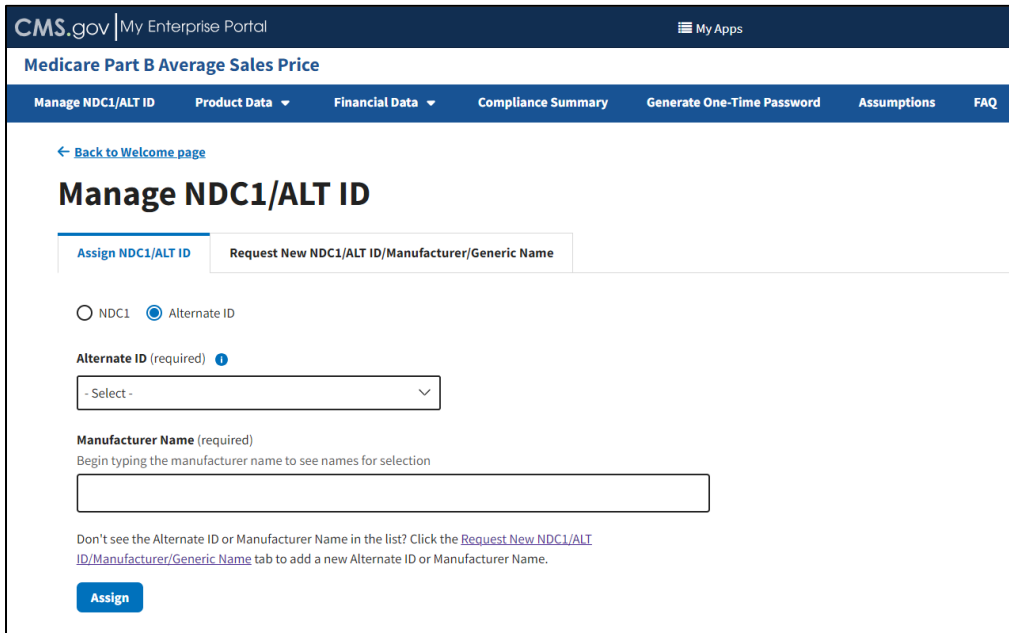

 The screenshot shows the CMS.gov 'My Enterprise Portal' interface. The title bar includes the CMS logo and 'My Enterprise Portal' text. A 'My Apps' button is in the top right. Below the title bar, a blue navigation bar has 'Back to Welcome page' and 'Manage NDC1/ALT ID' tabs. The 'Manage NDC1/ALT ID' tab is active. Underneath, there are two radio buttons: 'NDC1' (selected) and 'Alternate ID'. A green success message box contains the text: 'The mapping for Test Manf Name impl and 99999 was completed successfully.' Below this, there are two dropdown menus: 'NDC1 (required)' with the placeholder '- Select -' and 'Manufacturer Name (required)' with the placeholder 'Begin typing the manufacturer name to see names for selection'. A note at the bottom says: 'Don't see the NDC1 or Manufacturer Name in the list? Click the Request New NDC1/ALT ID/Manufacturer/Generic Name tab to add a new NDC1 or Manufacturer Name.' A blue 'Assign' button is at the bottom.

Figure 12: Manage NDC1/ALT ID - NDC1 Assigned Successfully

3.1.2 Assign by Alternate ID

Follow these steps to assign Alternate IDs:

1. From the Medicare Part B Average Sales Price homepage, click the **Manage NDC1/ALT ID** tab.
The **Manage NDC1/ALT ID** page opens and displays the **Assign NDC1/ALT ID** tab by default.
2. From the **Assign NDC1/ALT ID** tab, select the **Alternate ID** radio button.
Additional fields specific to assigning an Alternate ID display. Refer to *Figure 13*.



CMS.gov | My Enterprise Portal My Apps

Medicare Part B Average Sales Price

Manage NDC1/ALT ID Product Data Financial Data Compliance Summary Generate One-Time Password Assumptions FAQ

[← Back to Welcome page](#)

Manage NDC1/ALT ID

[Assign NDC1/ALT ID](#) [Request New NDC1/ALT ID/Manufacturer/Generic Name](#)

NDC1 Alternate ID

Alternate ID (required) ⓘ

- Select -

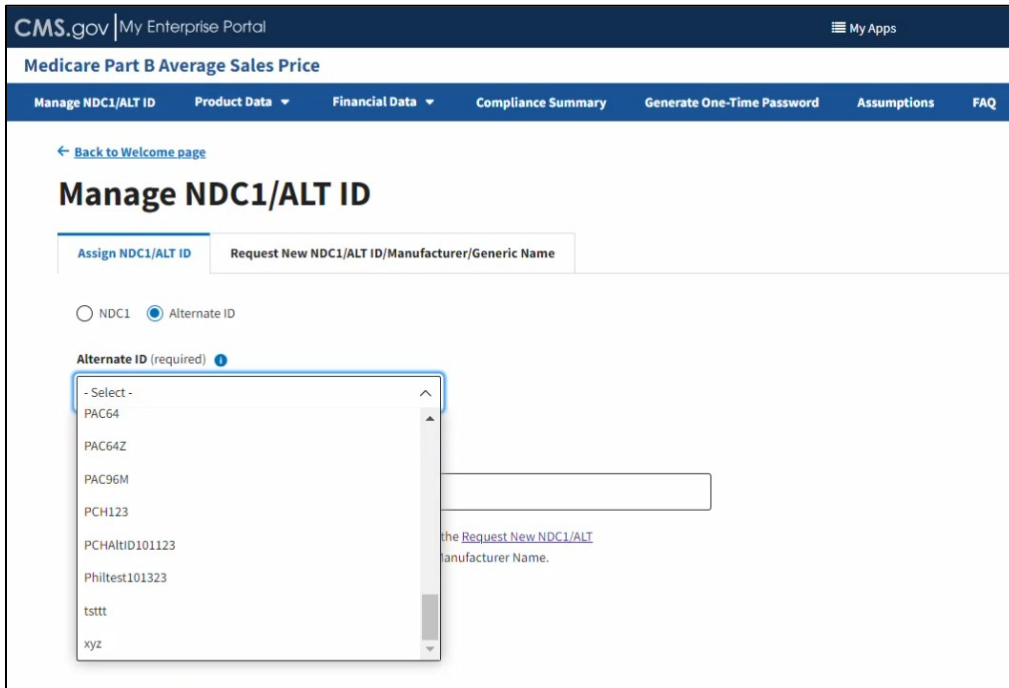
Manufacturer Name (required)
Begin typing the manufacturer name to see names for selection

Don't see the Alternate ID or Manufacturer Name in the list? Click the [Request New NDC1/ALT ID/Manufacturer/Generic Name](#) tab to add a new Alternate ID or Manufacturer Name.

Assign

Figure 13: Manage NDC1/ALT ID Page - Assign ALT ID

- Under **Alternate ID (required)**, click the **-Select-** drop-down menu to expand the list of submitted drugs and additional products in the Module to date; select the appropriate alternate ID. Refer to *Figure 14*.



CMS.gov | My Enterprise Portal My Apps

Medicare Part B Average Sales Price

Manage NDC1/ALT ID Product Data Financial Data Compliance Summary Generate One-Time Password Assumptions FAQ

[← Back to Welcome page](#)

Manage NDC1/ALT ID

[Assign NDC1/ALT ID](#) [Request New NDC1/ALT ID/Manufacturer/Generic Name](#)

NDC1 Alternate ID

Alternate ID (required) ⓘ

- Select -

PAC64

PAC64Z

PAC96M

PCH123

PCHAItID101123

Philtest101323

tsitt

xyz

Don't see the Alternate ID or Manufacturer Name in the list? Click the [Request New NDC1/ALT ID/Manufacturer/Generic Name](#) tab to add a new Alternate ID or Manufacturer Name.

Figure 14: Manage NDC1/ALT ID Page - Assign ALT ID Drop-down Menu

4. In the **Manufacturer Name (required)** auto-fill field, begin to type the manufacturer name; select the appropriate manufacturer from the list that generates. Refer to *Figure 15*.

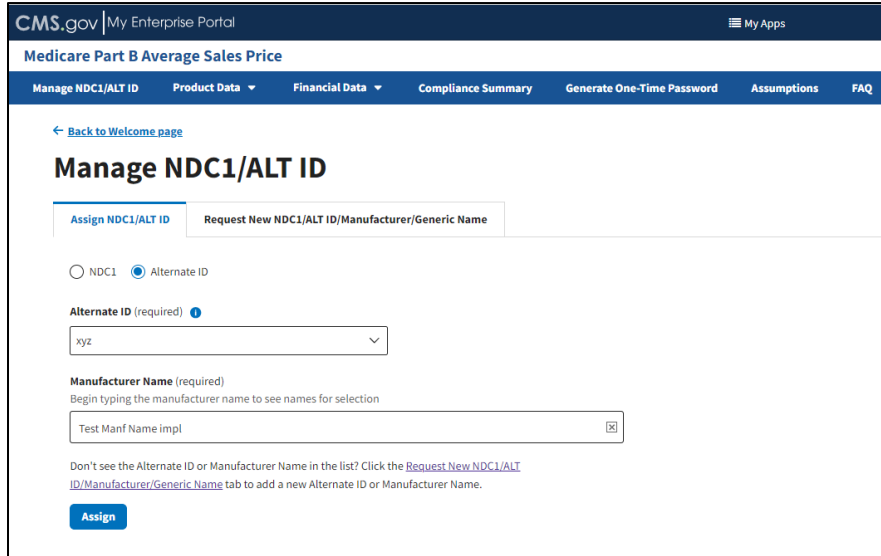

 A screenshot of the CMS.gov 'My Enterprise Portal' interface. The top navigation bar includes links for 'My Apps', 'Assign NDC1/ALT ID', 'Product Data', 'Financial Data', 'Compliance Summary', 'Generate One-Time Password', 'Assumptions', and 'FAQ'. Below the navigation is a breadcrumb trail with a link to 'Back to Welcome page'. The main content area is titled 'Manage NDC1/ALT ID'. It features two tabs: 'Assign NDC1/ALT ID' (selected) and 'Request New NDC1/ALT ID/Manufacturer/Generic Name' (disabled). The 'Assign NDC1/ALT ID' tab contains fields for 'Alternate ID (required)' (set to 'xyz') and 'Manufacturer Name (required)' (set to 'Test Manf Name impl'). A note at the bottom states: 'Don't see the Alternate ID or Manufacturer Name in the list? Click the Request New NDC1/ALT ID/Manufacturer/Generic Name tab to add a new Alternate ID or Manufacturer Name.' A blue 'Assign' button is at the bottom.

Figure 15: Manage NDC1/ALT ID Page - Enter ALT ID Manufacturer Name

5. Click **Assign** to submit your information.

A message displays confirming you have successfully added your selections. Refer to *Figure 16*.

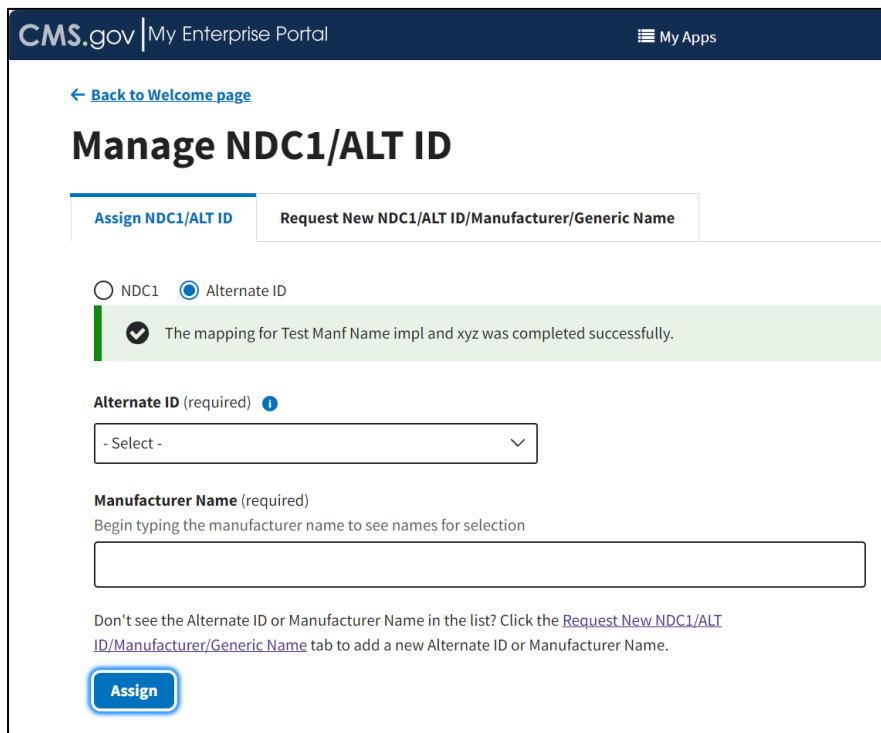

 A screenshot of the CMS.gov 'My Enterprise Portal' interface, showing the 'Manage NDC1/ALT ID' page after submission. The 'Assign NDC1/ALT ID' tab is selected. The 'Alternate ID (required)' field is empty. The 'Manufacturer Name (required)' field is empty. A green success message box at the top right of the page area says: 'The mapping for Test Manf Name impl and xyz was completed successfully.' A blue 'Assign' button is at the bottom.

Figure 16: Manage NDC1/ALT ID - ALT ID Assigned Successfully

3.1.3 Request New NDC1/ALT ID/Manufacturer/Generic Name

The following sections describe how to request a new NDC1, ALT ID, manufacturer, and generic name.

3.1.3.1 Request New NDC1

Follow these steps to request a new NDC1:

1. Navigate to the **Manage NDC1/ALT ID** page, which automatically opens on the **Assign NDC1/ALT ID** tab.
2. Click the **Request New NDC1/ALT ID/Manufacturer/Generic Name** tab.

The **Request New NDC1/ALT ID/Manufacturer/Generic Name** page opens, showing the status (**Pending**, **Approved**, or **Rejected**) for each submitted request. The Module organizes data by **Request Type**, **Requested Value** as well as **Request Date** and **Status (Pending, Approved, or Rejected)**. Refer to *Figure 17*.

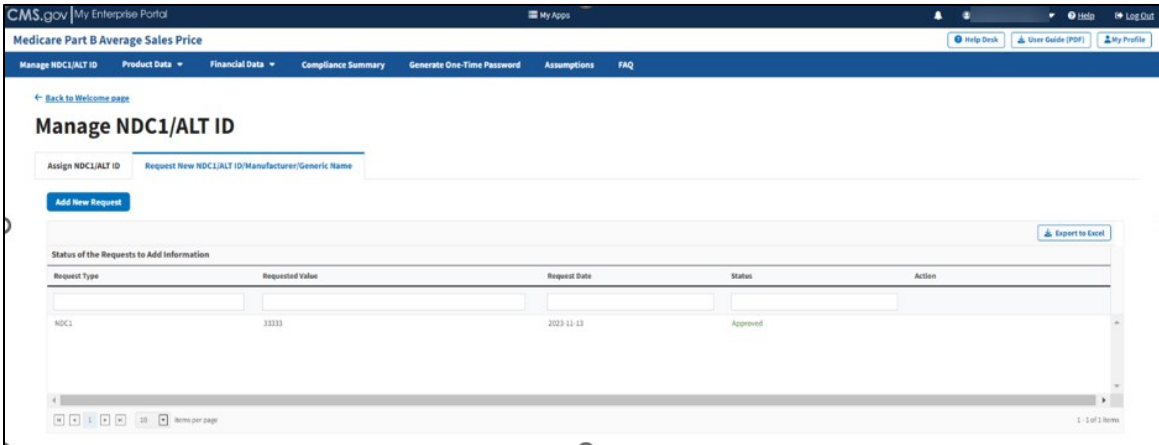


Figure 17: Request New NDC1/ALT ID/Manufacturer/Generic Name Page

3. Click the **Add New Request** button.

An **Add New NDC1/ALT ID/Manufacturer/Generic Name** window opens. Refer to *Figure 18*.

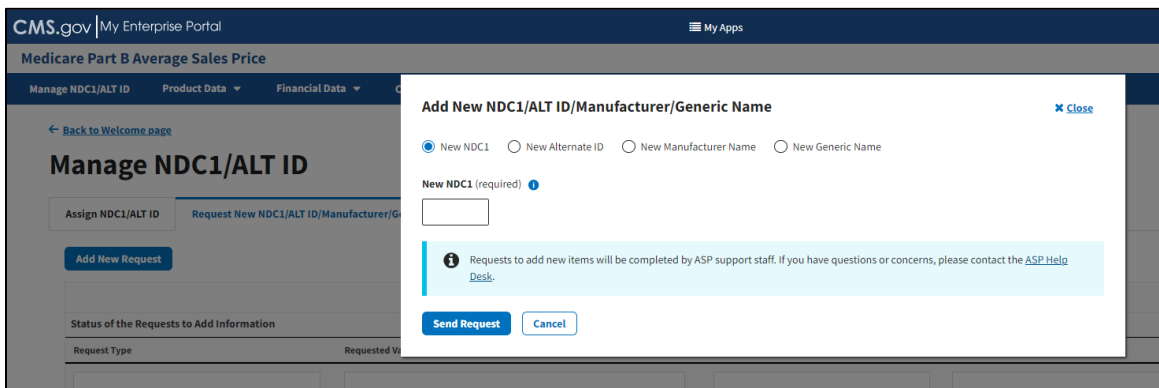


Figure 18: Request New NDC1/ALT ID/Manufacturer/Generic Name Page - Add New NDC1

4. Select the **New NDC1** radio button to specify the product data you need to submit to the Module. Note that the Module automatically defaults to select the **New NDC1**.
5. Under **New NDC1 (required)**, enter the appropriate NDC for the data product you are requesting to add to the Module. Refer to *Figure 19*.

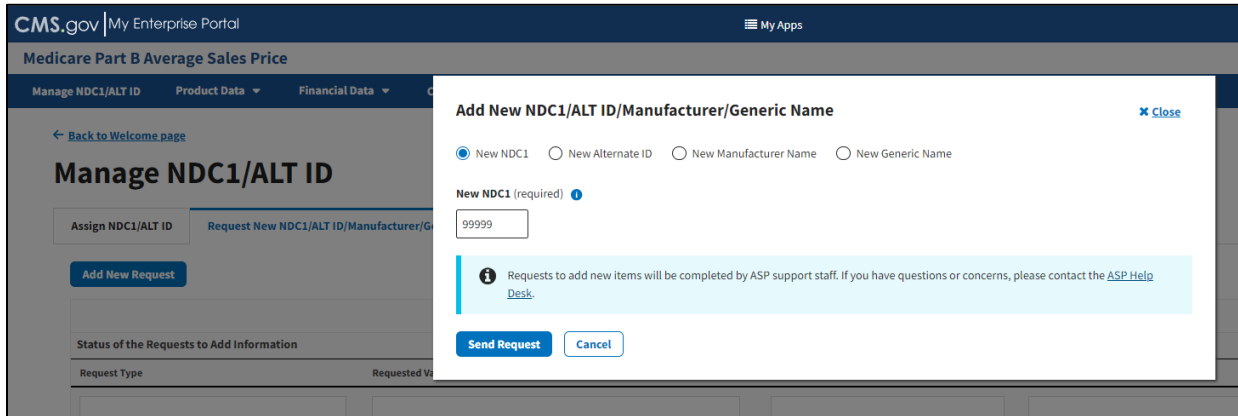


Figure 19: Request New NDC1 - Field Filled

6. Click **Send Request** to submit your information.

A message displays confirming you have successfully added your selections. Refer to *Figure 20*.

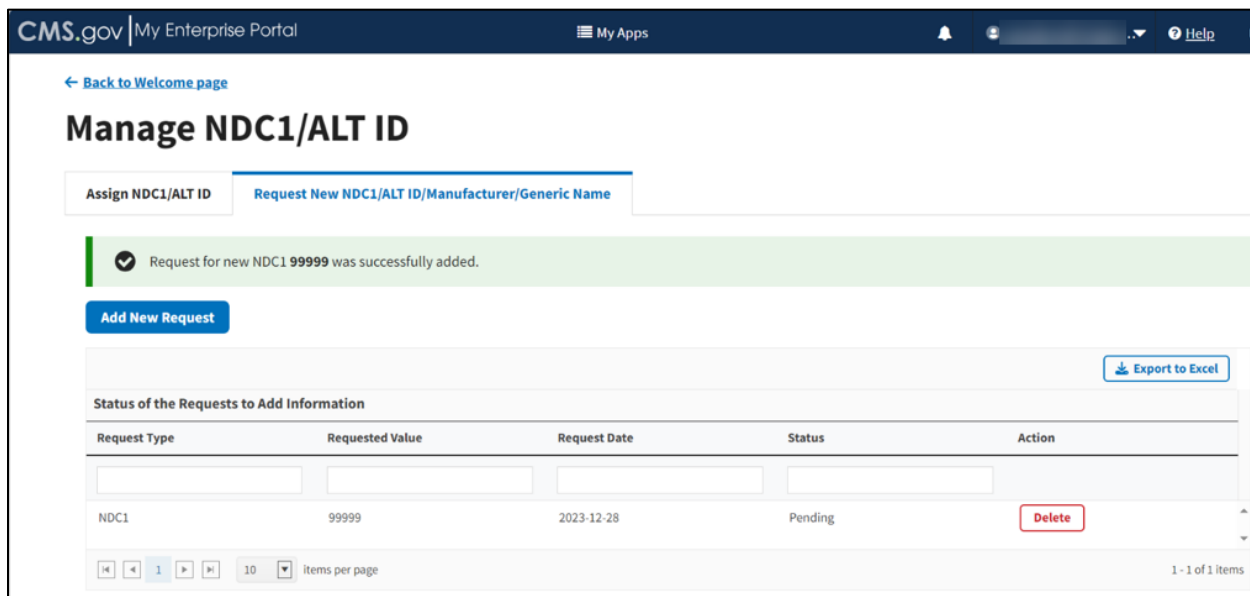


Figure 20: Request New NDC1 - NDC1 Successfully Added

3.1.3.2 Request New ALT ID

Follow these steps to request a new **Alternate ID**:

1. Click the **Add New Request** button. Refer to *Figure 20*.

An **Add New NDC1/ALT ID/Manufacturer/Generic Name** window opens. Note that the Module automatically defaults to the **New NDC1** tab.

2. Select the **New Alternate ID** radio button to specify the product data you need to submit to the Module. Refer to *Figure 21*.

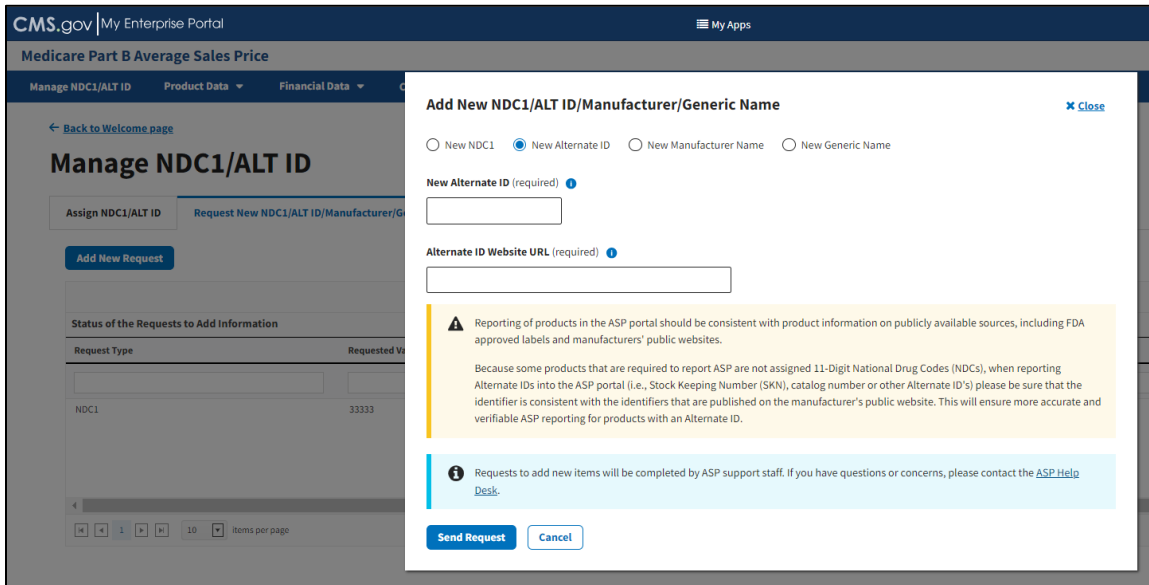

 A screenshot of the CMS ASP Portal. The main title is 'Manage NDC1/ALT ID'. A sub-dialog box is open titled 'Add New NDC1/ALT ID/Manufacturer/Generic Name'. It contains three radio buttons: 'New NDC1' (unchecked), 'New Alternate ID' (checked), 'New Manufacturer Name' (unchecked), and 'New Generic Name' (unchecked). Below these are two input fields: 'New Alternate ID (required)' and 'Alternate ID Website URL (required)'. A yellow warning box contains text about reporting consistency with FDA sources and website URLs. A blue info box contains text about support requests. At the bottom are 'Send Request' and 'Cancel' buttons.

Figure 21: Request New NDC1/ALT ID/Manufacturer/Generic Name Page - Add New ALT ID

3. Under **New Alternate ID (required)**, enter the appropriate alternate ID for the product you want to add to the Module.

Note: An Alternate ID is a manufacturer-selected product identifier that can be any combination of letters or numbers unique to the product (i.e., Stock Keeping Number (SKN) or product number). The **New Alternate ID** field allows up to a maximum of 23 characters and special characters (colon, dash, or period).

4. Under **Alternate ID Website URL (required)**, enter the hyperlink for the drug manufacturer's website. Refer to *Figure 22*.

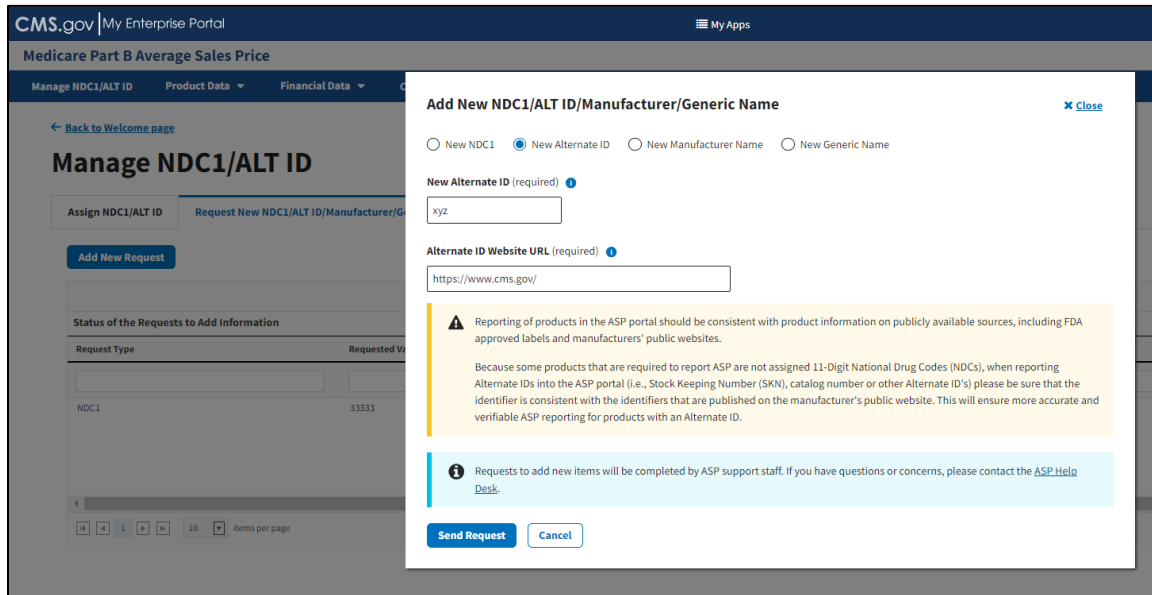

 A screenshot of the CMS.gov My Enterprise Portal. The main page shows a table of requests for managing NDC1/ALT IDs. An 'Add New Request' button is highlighted. A modal window titled 'Add New NDC1/ALT ID/Manufacturer/Generic Name' is open. In the modal, the 'New Alternate ID' field contains 'xyz' and the 'Alternate ID Website URL' field contains 'https://www.cms.gov/'. A note in the modal states: 'Reporting of products in the ASP portal should be consistent with product information on publicly available sources, including FDA approved labels and manufacturers' public websites.' Another note says: 'Requests to add new items will be completed by ASP support staff. If you have questions or concerns, please contact the [ASP Help Desk](#)'.

Figure 22: Request New Alternate ID - ALT ID Field Filled

- Click the **Send Request** button to submit your information.

A message displays confirming you have successfully added your selections. Refer to *Figure 23*.

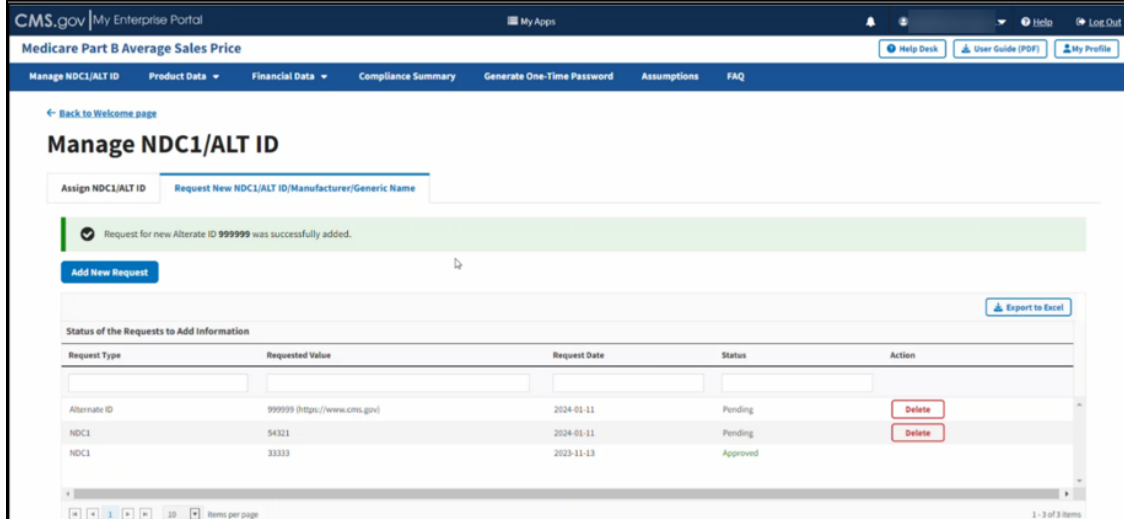

 A screenshot of the CMS.gov My Enterprise Portal. The main page shows a table of requests for managing NDC1/ALT IDs. A success message at the top of the page states: 'Request for new Alternate ID 999999 was successfully added.' Below this, a table shows the request history with one item successfully added. The table has columns: Request Type, Requested Value, Request Date, Status, and Action. The single item in the table is: 'Alternate ID' with value '999999 (https://www.cms.gov/)', 'Request Date' as '2024-01-11', 'Status' as 'Pending', and 'Action' with a 'Delete' button. There is also an 'Export to Excel' button in the table header.

Figure 23: Request New Alternate ID - ALT ID Successfully Added

3.1.3.3 Request New Manufacturer Name or New Generic Name

Follow these steps to request a **New Manufacturer** or **New Generic Name**:

- Click the **Add New Request** button. Refer to *Figure 23*.

An **Add New NDC1/ALT ID/Manufacturer/Generic Name** window opens. Note that the Module automatically defaults to the **New NDC1** tab.

2. Select either the **New Manufacturer Name** or the **New Generic Name** radio button to specify the product data you need to submit to the Module.

Additional fields display as the next page opens for either selection. Refer to *Figure 24* and *Figure 25*.

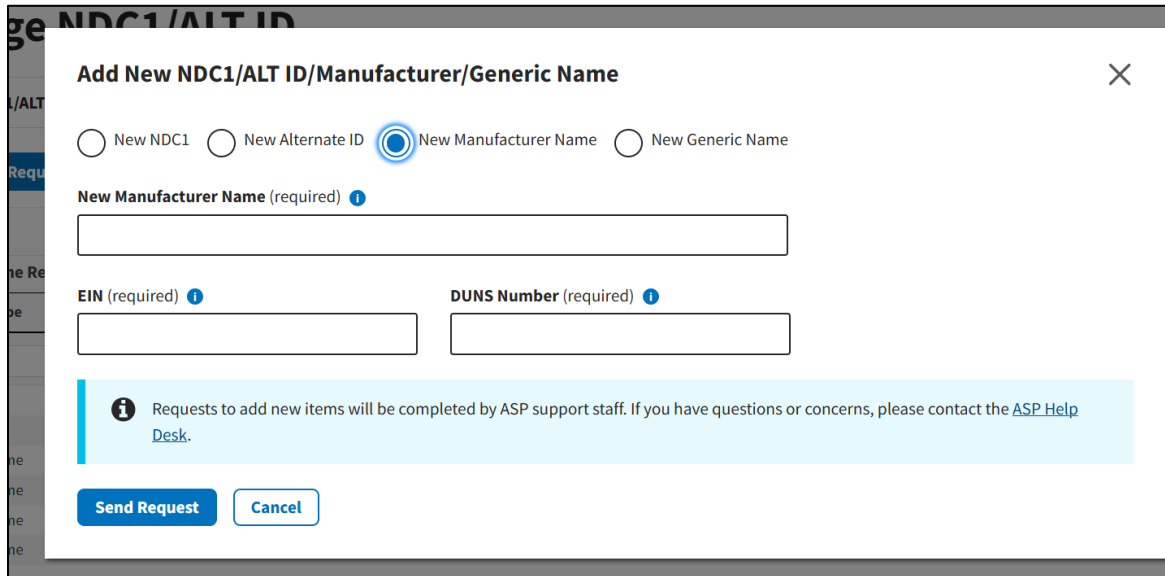

 A screenshot of a web-based application window titled "Add New NDC1/ALT ID/Manufacturer/Generic Name". The window has a light gray background with a dark gray header bar. At the top left, there is a vertical sidebar with the text "NDC1/ALT ID" and "Request". At the top right is a close button (an "X"). Below the title, there are four radio buttons: "New NDC1" (unselected), "New Alternate ID" (unselected), "New Manufacturer Name" (selected, indicated by a blue outline), and "New Generic Name" (unselected). Below the radio buttons are three input fields: "New Manufacturer Name (required)" with a blue info icon, "EIN (required)" with a blue info icon, and "DUNS Number (required)" with a blue info icon. Each input field has a corresponding empty text box. Below these fields is a light blue callout box containing the text: "Requests to add new items will be completed by ASP support staff. If you have questions or concerns, please contact the [ASP Help Desk](#)". At the bottom of the window are two buttons: "Send Request" (blue) and "Cancel" (white).

Figure 24: Request New Manufacturer Name

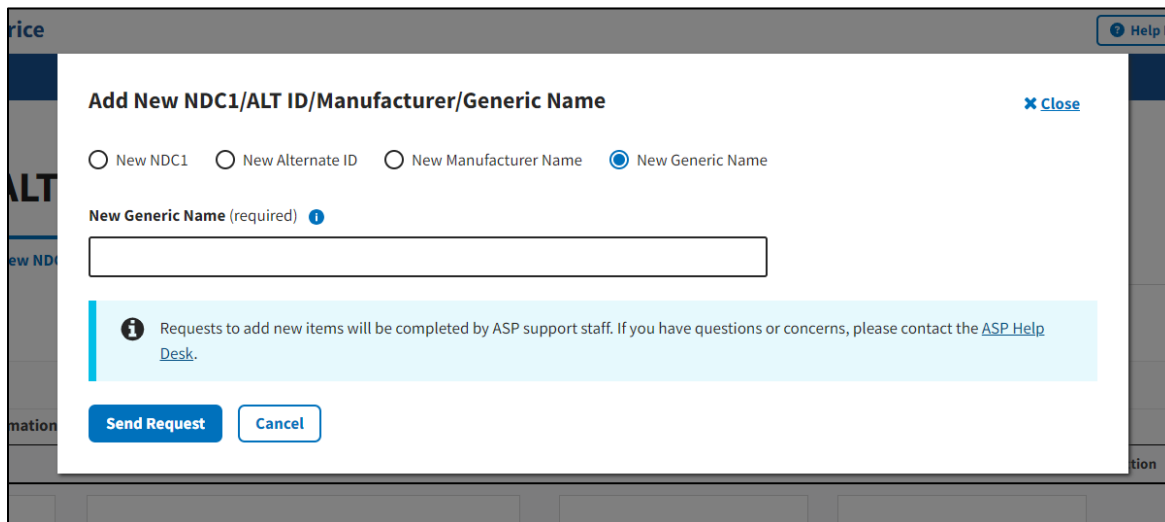

 A screenshot of a web-based application window titled "Add New NDC1/ALT ID/Manufacturer/Generic Name". The window has a light gray background with a dark gray header bar. At the top left, there is a vertical sidebar with the text "NDC1/ALT ID" and "Request". At the top right is a close button (an "X"). Below the title, there are four radio buttons: "New NDC1" (unselected), "New Alternate ID" (unselected), "New Manufacturer Name" (unselected), and "New Generic Name" (selected, indicated by a blue outline). Below the radio buttons is one input field: "New Generic Name (required)" with a blue info icon, which has an empty text box. Below this field is a light blue callout box containing the text: "Requests to add new items will be completed by ASP support staff. If you have questions or concerns, please contact the [ASP Help Desk](#)". At the bottom of the window are two buttons: "Send Request" (blue) and "Cancel" (white).

Figure 25: Request New Generic Name

3. Under **New Manufacturer Name (required)** or **New Generic Name (required)**, enter the appropriate information for the data product you want to add to the Module. Refer to *Figure 26* and *Figure 27*.

Note: For **New Manufacturer Name** requests, users will need to submit business identification information including the manufacturer's Employer Identification Number (EIN) and Data Universal Numbering System (DUNS) number.

ASP Help Desk.' Two buttons at the bottom are 'Send Request' and 'Cancel'." data-bbox="159 117 832 367"/>

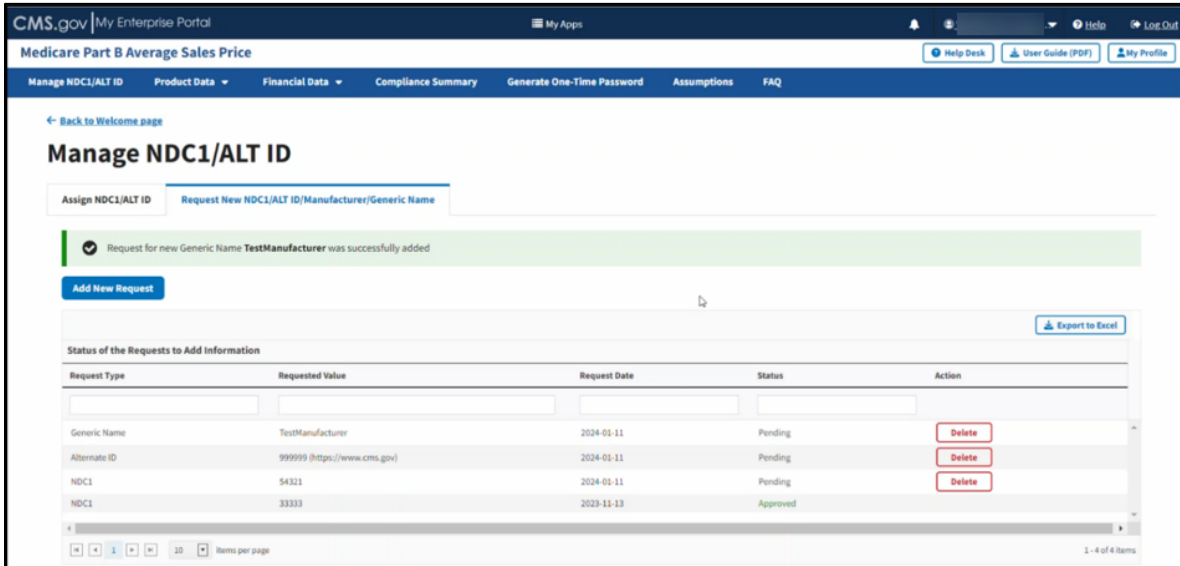
Figure 26: Request New Manufacturer Name - Field Populated

ASP Help Desk.' Two buttons at the bottom are 'Send Request' and 'Cancel'." data-bbox="154 403 837 639"/>

Figure 27: Request New Generic Name - Field Populated

4. Click **Send Request** to submit your information for either selection.

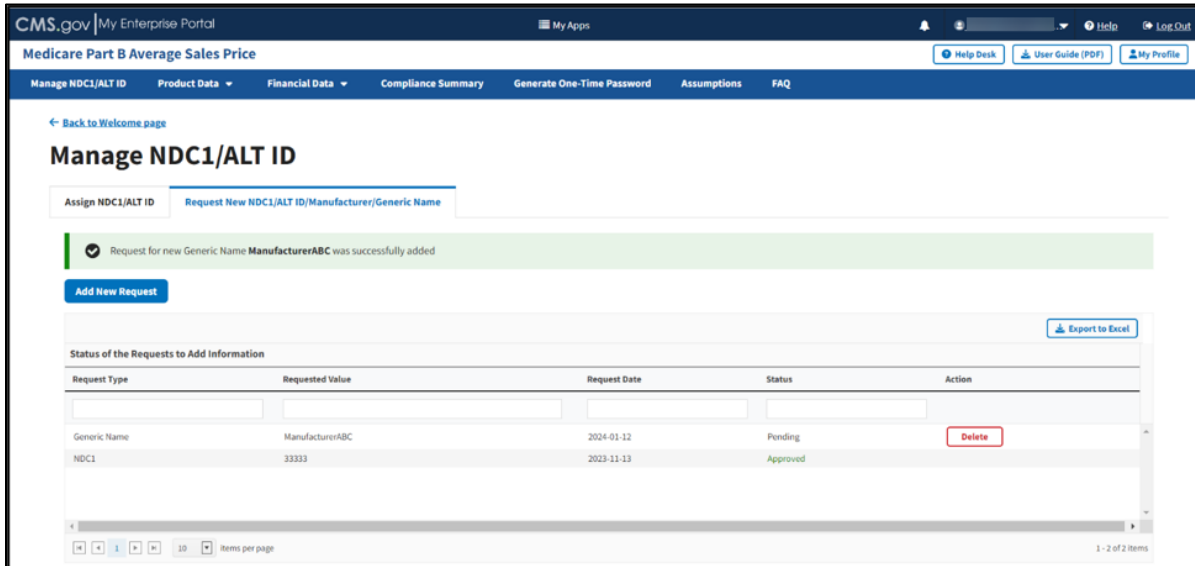
A message displays confirming you have successfully added your selections. Refer to *Figure 28* and *Figure 29*.



Request for new Generic Name **TestManufacturer** was successfully added

Request Type	Requested Value	Request Date	Status	Action
Generic Name	TestManufacturer	2024-01-11	Pending	Delete
Alternate ID	999999 (https://www.cms.gov)	2024-01-11	Pending	Delete
NDC1	54321	2024-01-11	Pending	Delete
NDC1	33333	2023-11-13	Approved	

Figure 28: Request New Manufacturer Name - Successfully Added



Request for new Generic Name **ManufacturerABC** was successfully added

Request Type	Requested Value	Request Date	Status	Action
Generic Name	ManufacturerABC	2024-01-12	Pending	Delete
NDC1	33333	2023-11-13	Approved	

Figure 29: Request New Generic Name - Successfully Added

Note: ASP support staff complete requests to add new items. Contact asphelpdesk@dcca.com for further questions or concerns about the process.

3.2 Product Data

Drug manufacturers must submit quarterly drug pricing data using a file transfer process, or through online data entry in the ASP module. Drug data consists of product and financial data. Click the **Product Data** tab on the Medicare Part B Average Sales Price homepage to view the drop-down menu for the **Add/Update Product Data**, **Upload Product Data**, and **View Drugs** tabs. Refer to *Figure 30*.

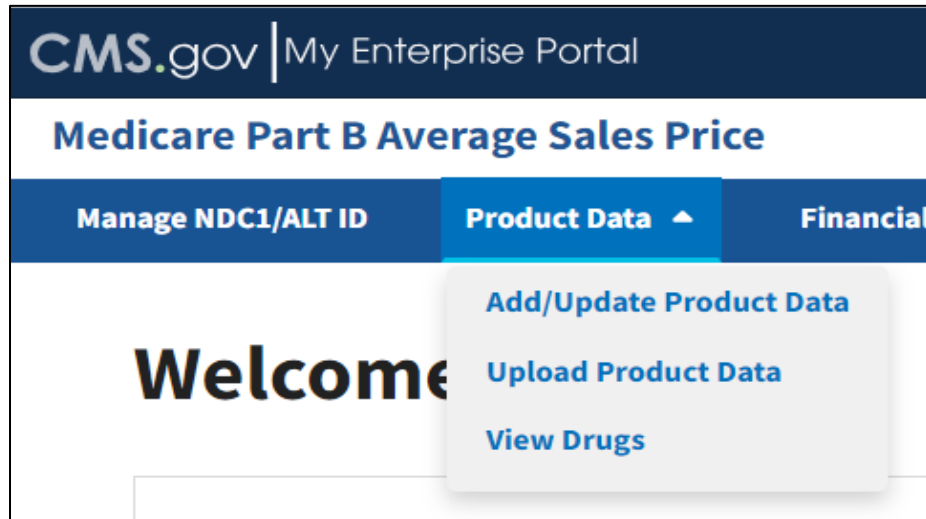


Figure 30: Product Data - Main Drop-down Menu

The following sections describe how to add/update, upload product data, and view drugs.

3.2.1 Add/Update Product Data

Follow these steps to add and/or update product data:

1. From the Medicare Part B Average Sales Price homepage, click the **Product Data** tab; then select the **Add/Update Product Data** tab.

The **Add/Update Product Data** page opens with default selections. Refer to *Figure 31*.

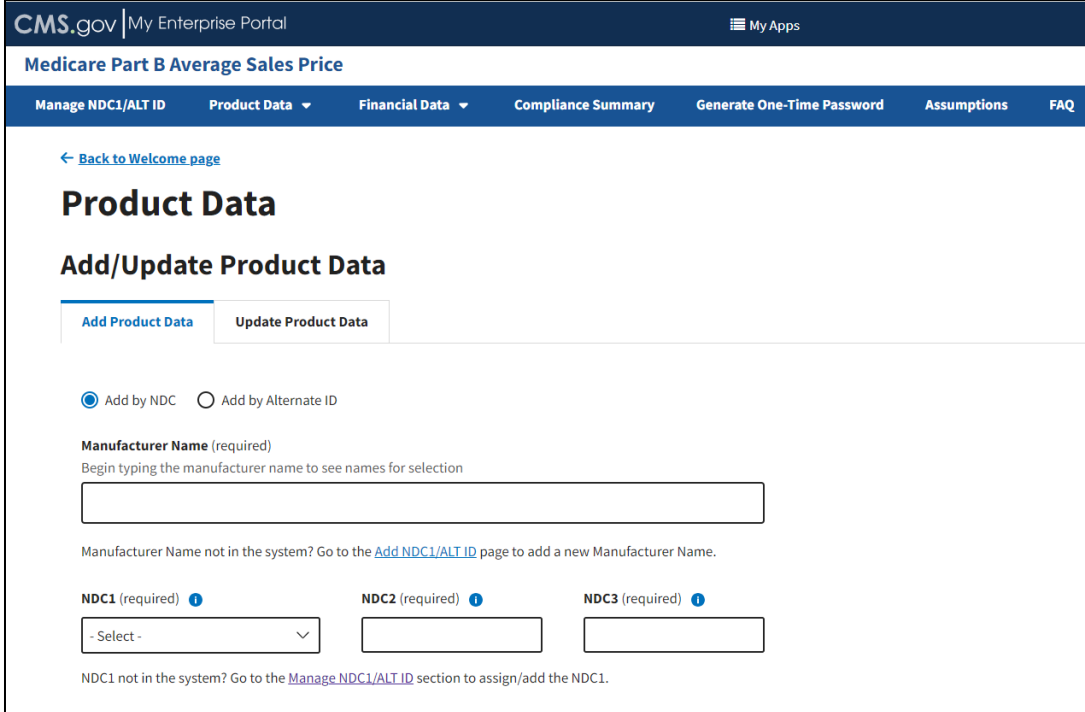


Figure 31: Add/Update Product Data

Note: It is imperative that the spelling matches each time you enter product data for the same drug manufacturer. The spelling must also match when entering data under the **Upload Product Data** tab.

3.2.1.1 Add Product Data by NDC

Follow these steps to add product data by NDC:

1. From the **Add Product Data** page, select the **Add by NDC** radio button if it is not already selected when the page opens.
2. In the **Manufacturer Name (required)** field, begin to type and then select the appropriate manufacturer.
3. Under **NDC1 (required)**, click the **-Select-** drop-down menu to expand the list of submitted drugs and additional products in the Module to date; select your required NDC1* code.
4. Enter your 4-digit number in the **NDC2* (required)** field.
5. Enter your 2-digit number in the **NDC3* (required)** field.

As you complete the **NDC3* (required)** field, the **Add Product Data** page expands to display multiple drop-down menus and empty fields.

6. Enter or select the required information as follows:
 - a. Select the **Drug has brand name** checkbox if the product you are submitting has a brand name. (If so, an empty field displays to submit the brand name; type information here as needed.)
 - b. Click the **Generic Name (required)** drop-down menu; select the generic name you need to enter for your product.

Note: Return to the **Manage NDC1/ALT ID** page if you cannot find the appropriate generic name in the system. Refer to *Section 3.1 - Manage NDC1/ALT ID* for guidance.

- c. Enter the volume per item in the **Volume Per Item (required)** field.
- d. Click the **Unit for Volume Per Item (required)** drop-down; select the appropriate option for your product.
- e. Enter the appropriate number in the **Number of Items per NDC (required)** field.
- f. Click the **Package Type (required)** drop-down; select the appropriate package type.
- g. Enter the strength in the **Strength (required)** field.
- h. Click the **Unit for Strength (required)** drop-down; select the appropriate unit.
- i. Enter the FDA application number in the **FDA Application Number (required)** field.
- j. Enter the FDA application supplement number in the **FDA Application Supplement Number** field, if applicable.

Note: Click the **Add Additional FDA Application Numbers** button if applicable for the drug, and repeat steps i and j.

- k. Enter the FDA approval date in the **FDA Approval Date (required)** field.
- l. Click the **FDA Approval Type (required)** drop-down; select the appropriate approval type.

- m. Enter the first marketing date in the **First Marketing Date (required)** field.
- n. Enter the date of first sale in the **Date of First Sale (required)** field.
- o. Enter the WAC in the **Wholesale Acquisition Cost (required)** field.

Note: The **Wholesale Acquisition Cost (required)** field is required and displays when the **First Marketing Date** occurs after the current reporting period.

Note: The date of first sale cannot occur before the FDA approval date and must occur prior to the current reporting period start date.

7. Confirm your selections. Refer to *Figure 32*.

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Product Data

Add/Update Product Data

[Add Product Data](#) [Update Product Data](#)

Add by NDC Add by Alternate ID

Manufacturer Name (required) *
Begin typing the manufacturer name to see names for selection

Manufacturer Name not in the system? Go to the [Add NDC1/ALT ID](#) page to add a new Manufacturer Name.

NDC1 (required) * **NDC2** (required) * **NDC3** (required) *

Drug has a brand name

Generic Name (required) *
Begin typing the generic name to see names for selection

Generic Name not in the system? Go to the [Add NDC1/ALT ID](#) page to add a new generic name.

Volume per Item (required) * **Unit for Volume per Item** (required) *

Number of Items per NDC (required) * **Package Type** (required) *

Strength (required) * **Unit for Strength** (required) *

FDA Application (Primary)

FDA Application Number (required) * <input type="text" value="000001"/>	FDA Application Supplement Number * <input type="text" value="0001"/>
--	--

[Add Additional FDA Application Numbers](#)

FDA Approval Date (required) *
MM/DD/YYYY

FDA Approval Type (required) *

First Marketing Date (required) *
MM/DD/YYYY

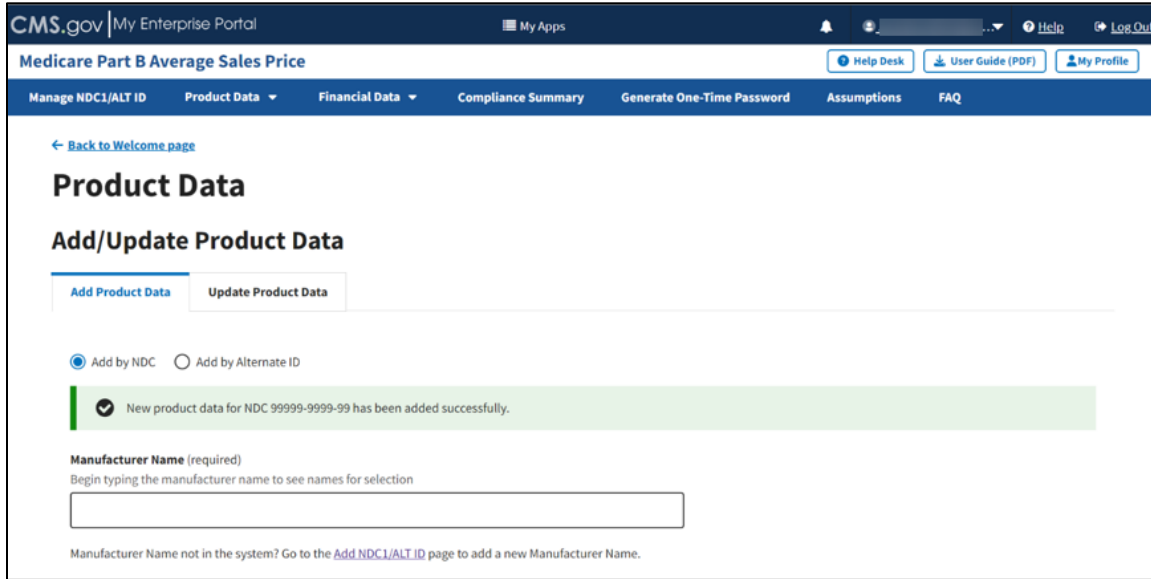
Date of First Sale for this NDC (required) *
MM/DD/YYYY

[Add Product Data](#)

Figure 32: Add/Update Product Data Fields Populated

8. Click **Add Product Data** to submit your information.

A message displays confirming you have successfully added your selections. Refer to *Figure 33*.



The screenshot shows the CMS.gov Product Data page. At the top, there is a navigation bar with links for 'My Apps', 'Help Desk', 'User Guide (PDF)', and 'My Profile'. Below the navigation bar, the page title is 'Medicare Part B Average Sales Price' and the sub-section is 'Product Data'. There are two buttons: 'Add Product Data' (highlighted in blue) and 'Update Product Data'. Below these buttons, there are two radio buttons: 'Add by NDC' (selected) and 'Add by Alternate ID'. A green success message box contains the text: 'New product data for NDC 99999-9999-99 has been added successfully.' Below the message box, there is a search field for 'Manufacturer Name (required)' with the placeholder 'Begin typing the manufacturer name to see names for selection'. At the bottom of the page, there is a note: 'Manufacturer Name not in the system? Go to the [Add NDC1/ALT ID](#) page to add a new Manufacturer Name.'

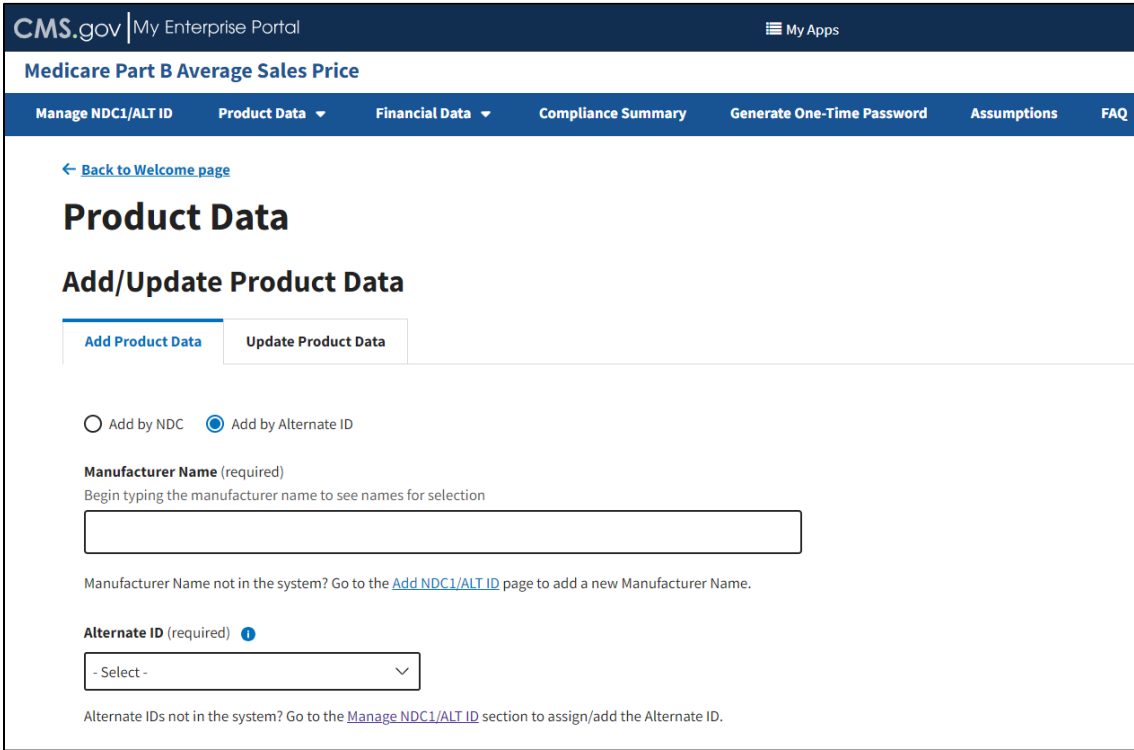
Figure 33: Add/Update Product Data Successfully Added

Note: It is imperative that the spelling matches each time you enter product data for the same drug manufacturer. The spelling must also match when entering data under the **Upload Product Data** tab.

3.2.1.2 Add Product Data by Alternate ID

Follow these steps to add product data by alternate ID:

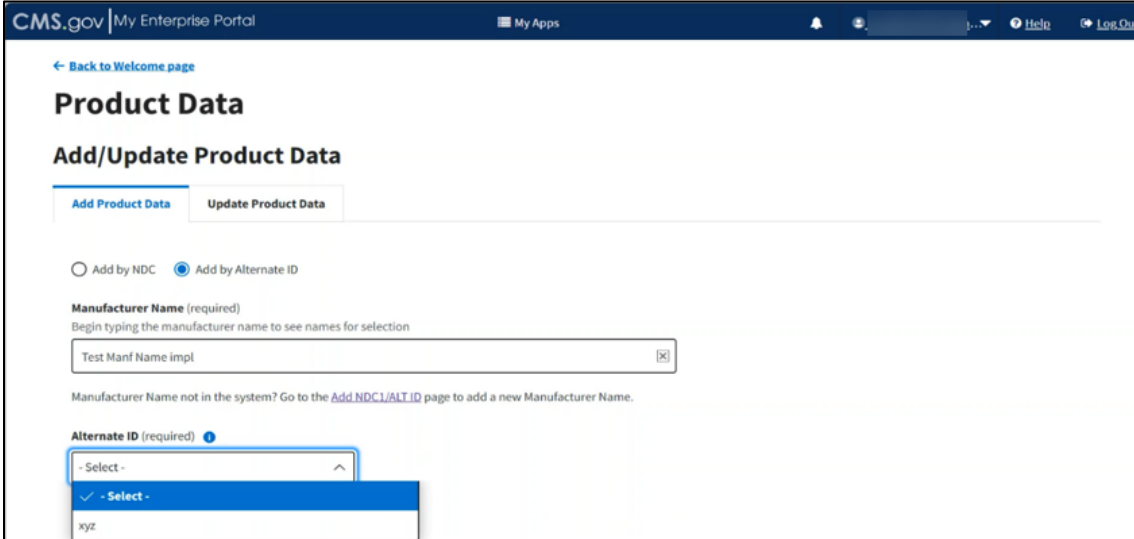
1. From the **Add/Update Product Data** page, select the **Add by Alternate ID** radio button. The **Add Product Data** page expands to display additional empty fields. Refer to *Figure 34*.



The screenshot shows the CMS.gov Product Data page. At the top, there are navigation links: Manage NDC1/ALT ID, Product Data, Financial Data, Compliance Summary, Generate One-Time Password, Assumptions, and FAQ. Below these are links to Back to Welcome page and Product Data. A sub-navigation bar for 'Add/Update Product Data' has 'Add Product Data' (selected) and 'Update Product Data' buttons. Under 'Add Product Data', there are two radio buttons: 'Add by NDC' (unchecked) and 'Add by Alternate ID' (checked). A 'Manufacturer Name (required)' field is present with a placeholder 'Begin typing the manufacturer name to see names for selection'. Below it is a note: 'Manufacturer Name not in the system? Go to the [Add NDC1/ALT ID](#) page to add a new Manufacturer Name.' Under 'Alternate ID (required)', there is a dropdown menu with options '- Select -', 'xyz', and 'Test Manf Name impl' (which is highlighted with a blue selection bar). A note below the dropdown says: 'Alternate IDs not in the system? Go to the [Manage NDC1/ALT ID](#) section to assign/add the Alternate ID.'

Figure 34: Add Product Data by Alternate ID

2. Under **Manufacturer Name (required)**, begin to type and then select the appropriate manufacturer.
3. Under **Alternate ID (required)**, click the **-Select-** drop-down to expand the list. Select the required alternate ID code. Refer to *Figure 35*.



This screenshot is identical to Figure 34, but the 'Test Manf Name impl' value is now selected in the 'Alternate ID (required)' dropdown menu, as indicated by the blue selection bar around the option.

Figure 35: Add Product Data by Alternate ID - Fields Populated

As you complete the **Alternate ID (required)** field, the **Add Product Data** page expands to show multiple drop-down menus and empty fields.

4. Enter or select the required information as follows:
 - a. Enter the uniform resource locator (URL) to the manufacturer website in the **Manufacturer's Website URL (required)** field for verification purposes.
 - b. Select the **Drug has a brand name** checkbox if the product you are submitting has a brand name. (If so, an empty field displays in which to enter the brand name; type information here as needed.)
 - c. Click the **Generic Name (required)** drop-down; select the generic name you need to enter for your product.

Note: Return to the **Manage NDC1/ALT ID** page if you cannot find the appropriate generic name in the system. Refer to *Section 3.1 - Manage NDC1/ALT ID* for guidance.

- d. Enter the volume per item in the **Volume Per Item (required)** field.
 - e. Click the **Unit for Volume Per Item (required)** drop-down; select the appropriate option for your product.

Note: For skin substitute products such as powders, sheets or discs, enter "One" for Volume and "Each" for Unit for Volume.

- f. Enter the appropriate number in the **Number of Items per Alternate ID (required)** field.
 - g. Click the **Package Type (required)** drop-down; select the appropriate package type. Package Type is not applicable to skin substitute sheets.
 - h. Enter the strength in the **Strength (required)** field.

Note: For skin substitute products, strength is determined by calculating the area of the product.

- i. Click the **Unit for Strength (required)** drop-down; select the appropriate unit.
 - j. Enter the FDA registration number in the **FDA Registration Number (required)** field.
 - k. Enter the FDA approval date in the **FDA Approval Date (required)** field.
 - l. Enter the FDA approval type in the **FDA Approval Type (required)** field.
 - m. Enter the first marketing date in the **First Marketing Date (required)** field.
 - n. Enter the date of first sale in the **Date of First Sale for this ALT ID (required)** field.

Note: The date of first sale cannot occur before the FDA approval date and must occur prior to the current reporting period start date.

5. Confirm your selections; click **Add Product Data** to submit your information. Refer to *Figure 36*.

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Product Data

Add/Update Product Data

[Add Product Data](#) [Update Product Data](#)

Add by NDC Add by Alternate ID

Manufacturer Name (required) i
Begin typing the manufacturer name to see names for selection

Test Manf Name impl

Manufacturer Name not in the system? Go to the [Add NDC1/ALT ID](#) page to add a new Manufacturer Name.

Alternate ID (required) i

xy2

Alternate ID Website URL (required) i

TestData.com

Drug has a brand name

Generic Name (required) i
Begin typing the generic name to see names for selection

GENERIC

Generic Name not in the system? Go to the [Add NDC1/ALT ID](#) page to add a new generic name.

Volume per Item (required) i	Unit for Volume per Item (required) i
1	Capsule

Number of Items per Alternate ID (required) i	Package Type (required) i
30	SINGLE SOURCE

Strength (required) i	Unit for Strength (required) i
10	%

FDA Registration Number (required) i	FDA Approval Date (required) i
000009	MM/DD/YYYY 12/01/2022

FDA Approval Type (required) i	First Marketing Date (required) i
OTHER	MM/DD/YYYY 01/01/2023

Date of First Sale for this ALT ID (required) i
MM/DD/YYYY 02/01/2023

[Add Product Data](#)

Figure 36: Add Product Data by Alternate ID - Additional Fields

A message displays confirming you have successfully added your product data. Refer to *Figure 37*.

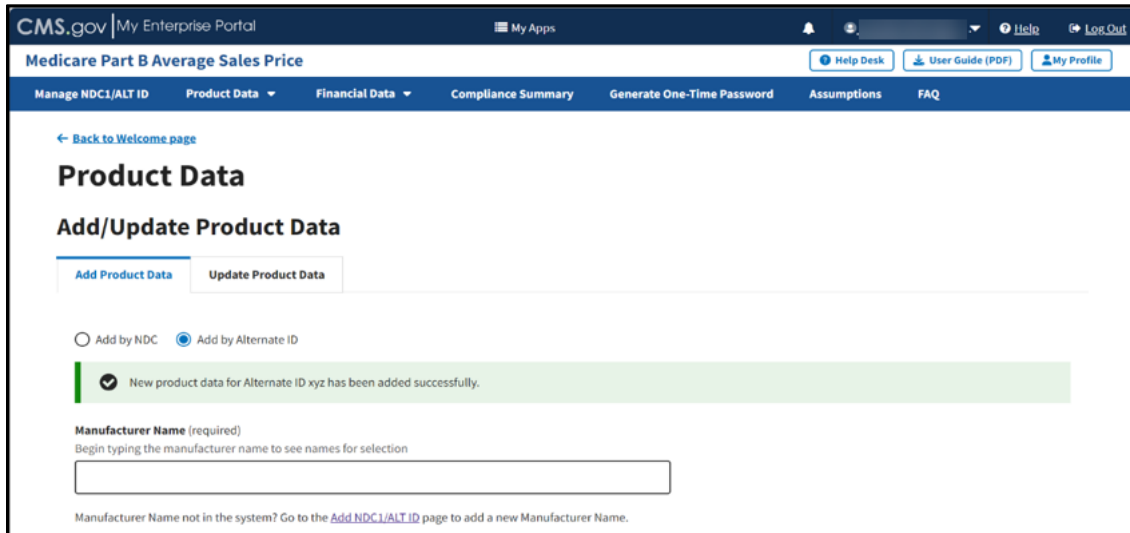

 A screenshot of the CMS.gov Product Data page. The page title is 'Product Data' and the sub-section is 'Add/Update Product Data'. There are two tabs: 'Add Product Data' and 'Update Product Data', with 'Update Product Data' being the active tab. Below the tabs, there are two radio buttons: 'Add by NDC' (unchecked) and 'Add by Alternate ID' (checked). A green success message box contains the text: 'New product data for Alternate ID xyz has been added successfully.' Below the message box, there is a field labeled 'Manufacturer Name (required)' with a placeholder 'Begin typing the manufacturer name to see names for selection' and a text input field. At the bottom of the page, there is a note: 'Manufacturer Name not in the system? Go to the [Add NDC1/ALT ID](#) page to add a new Manufacturer Name.'.

Figure 37: Product Data by Alternate ID Added Successfully

Note: It is imperative that the spelling matches each time you enter product data for the same drug manufacturer. The spelling must also match when entering data under the **Upload Product Data** tab.

3.2.1.3 Update Product Data by NDC

Follow these steps to update product data by NDC:

1. From the **Add/Update Product Data** page, select the **Update Product Data** tab; then, select the **Update by NDC** radio button if it is not already selected when the page opens. Refer to *Figure 38*.

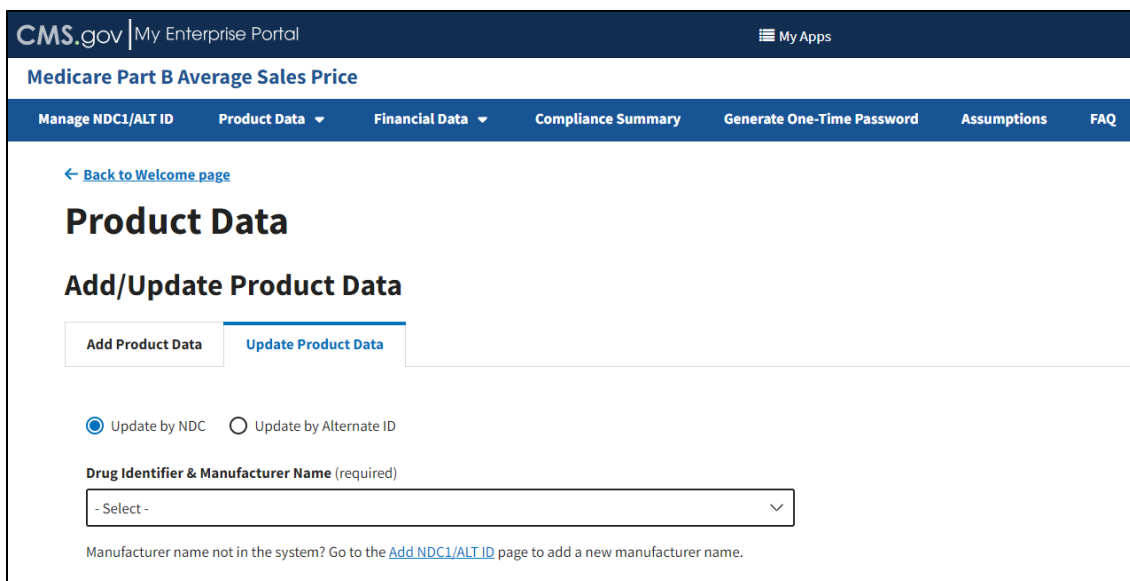

 A screenshot of the CMS.gov Product Data page, similar to Figure 37 but with the 'Update Product Data' tab selected. The 'Update by NDC' radio button is checked. Below the tabs, there is a field labeled 'Drug Identifier & Manufacturer Name (required)' with a dropdown menu showing the option '- Select -'. At the bottom of the page, there is a note: 'Manufacturer name not in the system? Go to the [Add NDC1/ALT ID](#) page to add a new manufacturer name.'.

Figure 38: Update Product Data - Drug Identifier & Manufacturer Name

2. In the **Drug Identifier & Manufacturer Name (required)** drop-down menu, click - **Select-** to expand the list of submitted drugs and additional products in the Module to date; select the appropriate drug identifier.

The page automatically loads the product data for that specific drug. Refer to *Figure 39*.

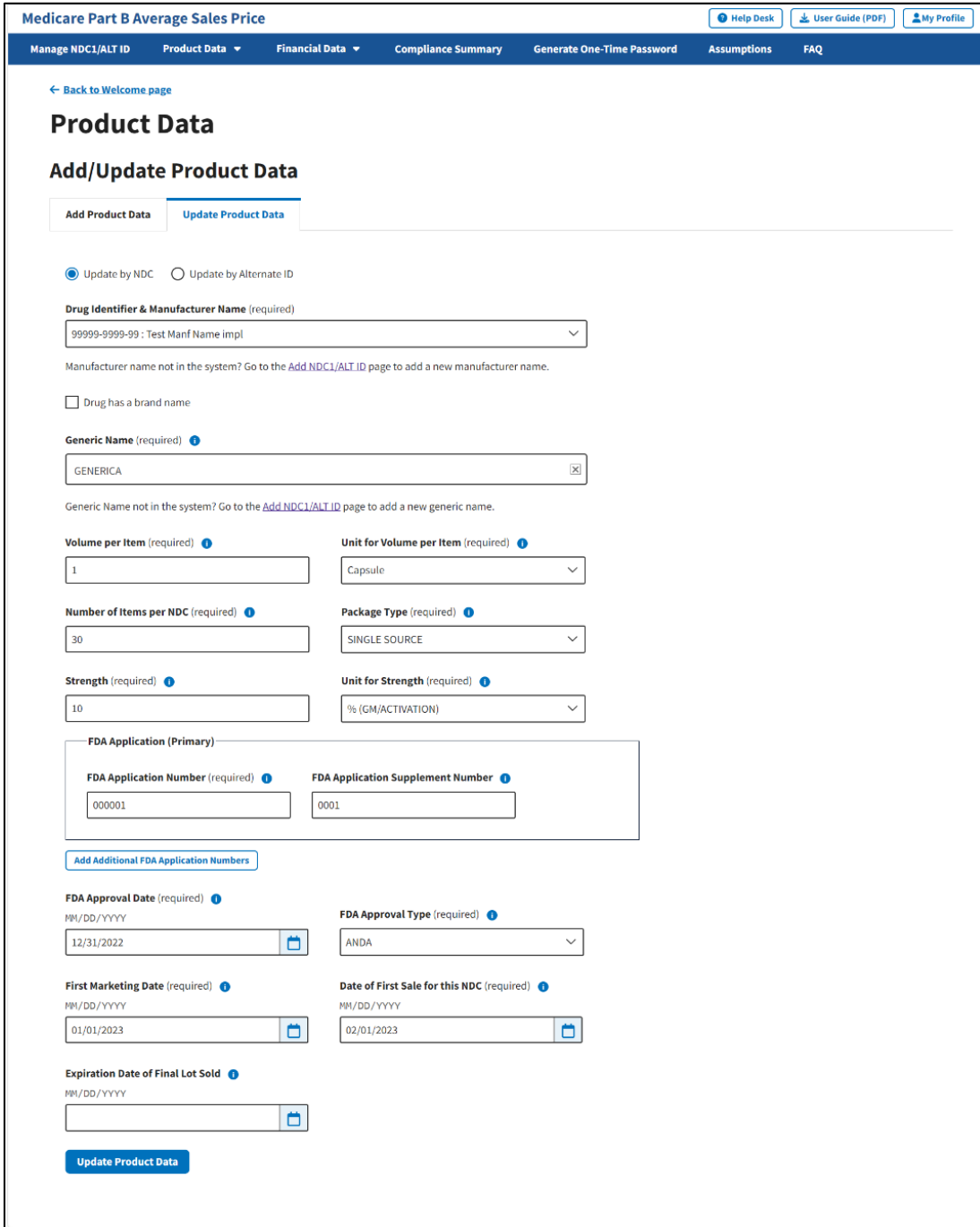

 The screenshot shows the 'Product Data' section of the CMS ASP Module Submitter. At the top, there are tabs for 'Add Product Data' and 'Update Product Data', with 'Update Product Data' being the active tab. Below these tabs, there are two radio buttons: 'Update by NDC' (selected) and 'Update by Alternate ID'. A dropdown menu labeled 'Drug Identifier & Manufacturer Name (required)' contains the entry '99999-9999-99 : Test Manf Name impl'. A note below the dropdown says 'Manufacturer name not in the system? Go to the [Add NDC1/ALT ID](#) page to add a new manufacturer name.' A checkbox 'Drug has a brand name' is present but unchecked. The 'Generic Name (required)' field contains 'GENERIC' with a delete icon. A note below it says 'Generic Name not in the system? Go to the [Add NDC1/ALT ID](#) page to add a new generic name.' Below this, there are fields for 'Volume per Item (required)' (value '1') and 'Unit for Volume per Item (required)' (value 'Capsule'). There are also fields for 'Number of Items per NDC (required)' (value '30') and 'Package Type (required)' (value 'SINGLE SOURCE'). For 'Strength (required)', the value is '10' and the 'Unit for Strength (required)' is '% (GM/ACTIVATION)'. A section for 'FDA Application (Primary)' contains fields for 'FDA Application Number (required)' (value '000001') and 'FDA Application Supplement Number (required)' (value '0001'). A link 'Add Additional FDA Application Numbers' is available. Below this, there are fields for 'FDA Approval Date (required)' (value '12/31/2022') and 'FDA Approval Type (required)' (value 'ANDA'). There are also fields for 'First Marketing Date (required)' (value '01/01/2023') and 'Date of First Sale for this NDC (required)' (value '02/01/2023'). A field for 'Expiration Date of Final Lot Sold (required)' (value 'MM/DD/YYYY') is present but empty. At the bottom, there is a blue 'Update Product Data' button.

Figure 39: Update Product Data by NDC

3. Review all your information in the appropriate boxes previously submitted in *Section 3.2 - Product Data*.
4. Confirm your selections; click **Update Product Data** to submit any changes in your drug product data.

A message displays confirming you have successfully updated your product data. Refer to *Figure 40*.

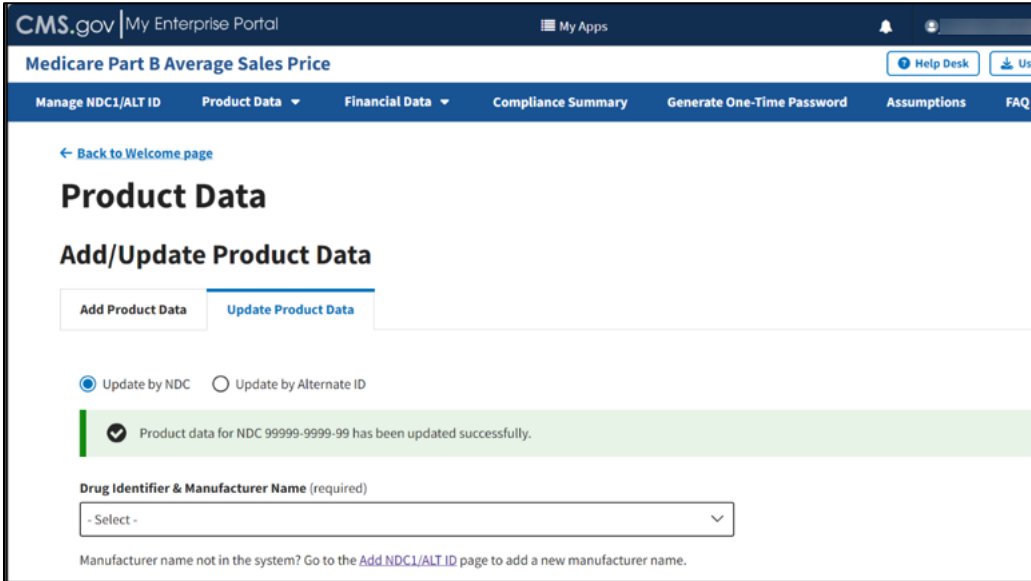

 A screenshot of the CMS.gov Product Data page. The page title is 'Medicare Part B Average Sales Price'. The 'Update Product Data' tab is selected. Below the tabs, there are two radio buttons: 'Update by NDC' (selected) and 'Update by Alternate ID'. A green success message box contains the text 'Product data for NDC 99999-9999-99 has been updated successfully.' Below the message box is a dropdown menu labeled 'Drug Identifier & Manufacturer Name (required)' with the placeholder '- Select -'. At the bottom of the page, a note says 'Manufacturer name not in the system? Go to the [Add NDC1/ALT ID](#) page to add a new manufacturer name.'.

Figure 40: Update Product Data by NDC - Data Updated Successfully

3.2.1.4 Update Product Data by Alternate ID

Follow these steps to update product data by Alternate ID:

1. From the **Add/Update Product Data** page, select the **Update Product Data** tab; then, select the **Update by Alternate ID** radio button. Refer to *Figure 41*.

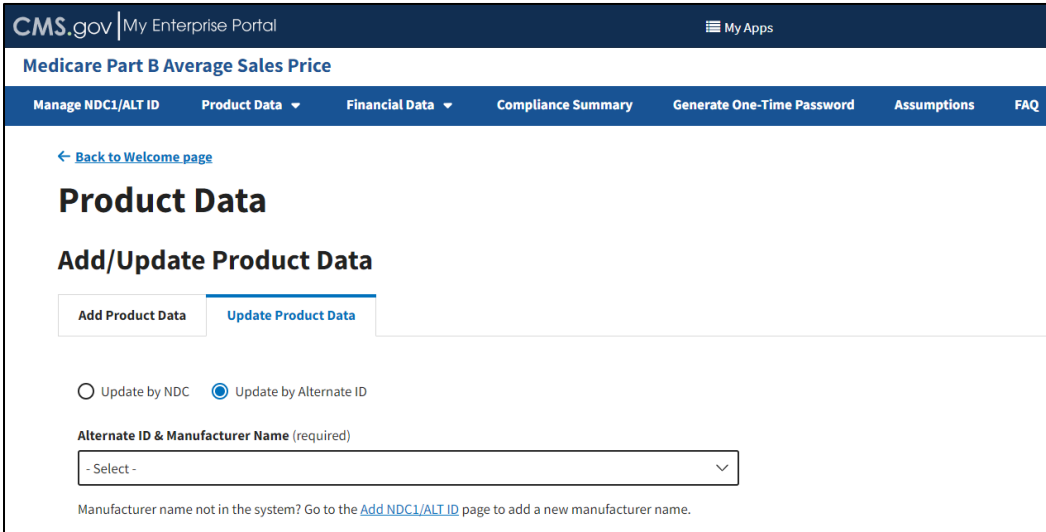

 A screenshot of the CMS.gov Product Data page. The page title is 'Medicare Part B Average Sales Price'. The 'Update Product Data' tab is selected. Below the tabs, there are two radio buttons: 'Update by NDC' (unchecked) and 'Update by Alternate ID' (selected). A dropdown menu labeled 'Alternate ID & Manufacturer Name (required)' with the placeholder '- Select -' is visible. At the bottom of the page, a note says 'Manufacturer name not in the system? Go to the [Add NDC1/ALT ID](#) page to add a new manufacturer name.'.

Figure 41: Update Product Data by Alternate ID

- Under the **Alternate ID & Manufacturer Name (required)** drop-down; click the **-Select-** drop-down to expand the list; select the appropriate information.

The page automatically loads the product data for that specific drug. Refer to *Figure 42*.

Note: Additional fields display on the next page. Ensure that you complete all required fields, and that all added financial information is accurate.

Medicare Part B Average Sales Price

Manage NDC1/ALT ID Product Data Financial Data Compliance Summary Generate One-Time Password Assumptions FAQ Help Desk User

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Product Data

Add/Update Product Data

[Add Product Data](#) [Update Product Data](#)

Update by NDC Update by Alternate ID

Alternate ID & Manufacturer Name (required)
 xyz : Test Manf Name impl

Manufacturer name not in the system? Go to the [Add NDC1/ALT ID](#) page to add a new manufacturer name.

Alternate ID Website URL (required) [i](#)
 http://cms.gov

Drug has a brand name

Generic Name (required) [i](#)
 GENERICA

Generic Name not in the system? Go to the [Add NDC1/ALT ID](#) page to add a new generic name.

Volume per Item (required) i	Unit for Volume per Item (required) i
1	Capsule

Number of Items per Alternate ID (required) i	Package Type (required) i
30	SINGLE SOURCE

Strength (required) i	Unit for Strength (required) i
10	%

FDA Registration Number (required) i	FDA Approval Date (required) i
000009	MM/DD/YYYY 12/01/2022

FDA Approval Type (required) i	First Marketing Date (required) i
OTHER	MM/DD/YYYY 01/01/2023

Date of First Sale for this ALT ID (required) i	Expiration Date of Final Lot Sold i
MM/DD/YYYY 02/01/2023	MM/DD/YYYY

[Update Product Data](#)

Figure 42: Update Product Data by Alternate ID - Drug Identifier Drop-down Menu

3. Review all your information in the appropriate boxes previously submitted in *Section 3.2 - Product Data*.
4. Confirm your selections; click **Update Product Data** to submit any changes in your drug product data.

A message displays confirming you have successfully updated your product data. Refer to *Figure 43*.

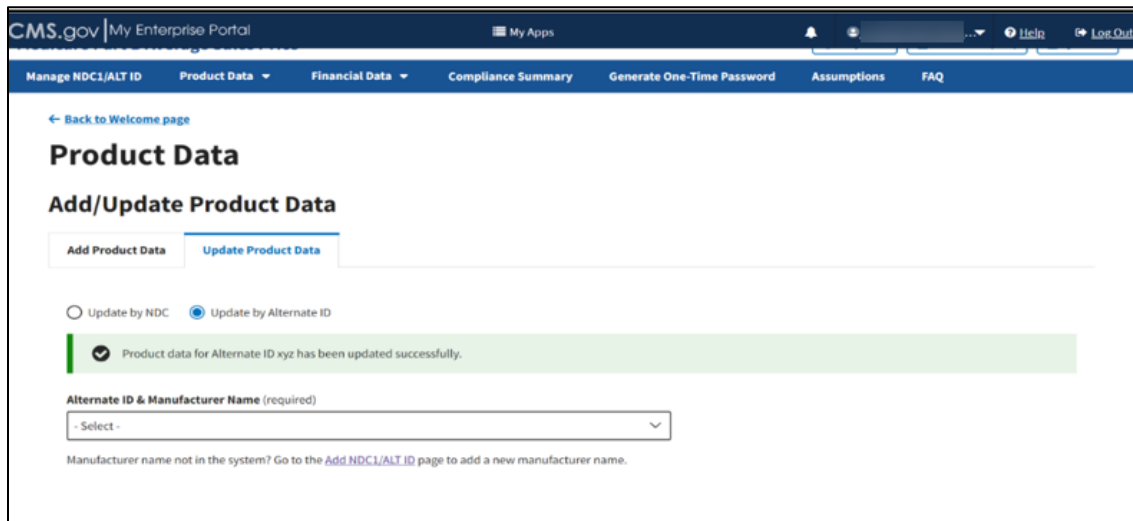

 A screenshot of the CMS.gov Product Data page. The page has a dark blue header with the CMS logo and a navigation bar with links like 'Manage NDC1/ALT ID', 'Product Data', 'Financial Data', 'Compliance Summary', 'Generate One-Time Password', 'Assumptions', and 'FAQ'. Below the header, there's a 'Product Data' section with a sub-section 'Add/Update Product Data'. Underneath that, there are two buttons: 'Add Product Data' and 'Update Product Data', with 'Update Product Data' being the active one. Below the buttons, there are two radio buttons: 'Update by NDC' (unchecked) and 'Update by Alternate ID' (checked). A green success message box contains the text 'Product data for Alternate ID xyz has been updated successfully.' At the bottom of the page, there's a dropdown menu labeled '- Select -' and a note: 'Manufacturer name not in the system? Go to the Add NDC1/ALT ID page to add a new manufacturer name.'.

Figure 43: Update Product Data by Alternate ID - Updated Successfully

3.2.2 Upload Product Data

Follow these steps to upload product data:

1. From the Medicare Part B Average Sales Price homepage, click the **Product Data** tab; then select the **Upload Product Data** tab.

The **Upload Product Data** page opens, listing the financial quarter and year for the upcoming reporting period. Refer to *Figure 44*.

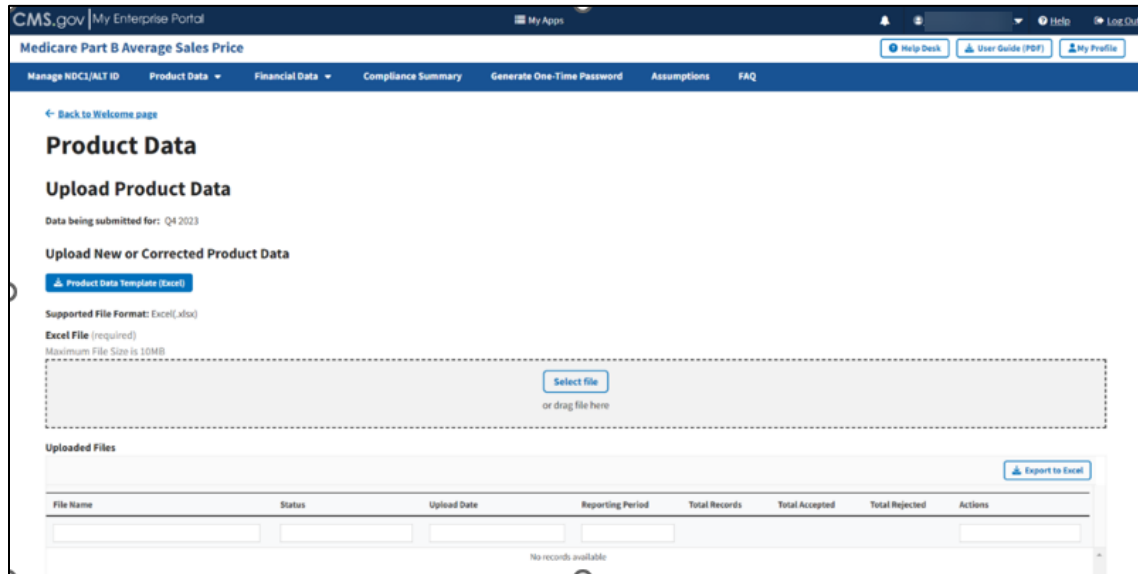


Figure 44: Upload Product Data - New or Corrected

Note: Click the **Product Data Template (Excel)** box to download a copy of the product data template.

- Upon preparing your **.xlsx file (required)** and verifying your information for accuracy, click **Select File**; then select the Excel file in the dialog box. You may also drag the file into the **Select File** box. Refer to *Figure 45*.

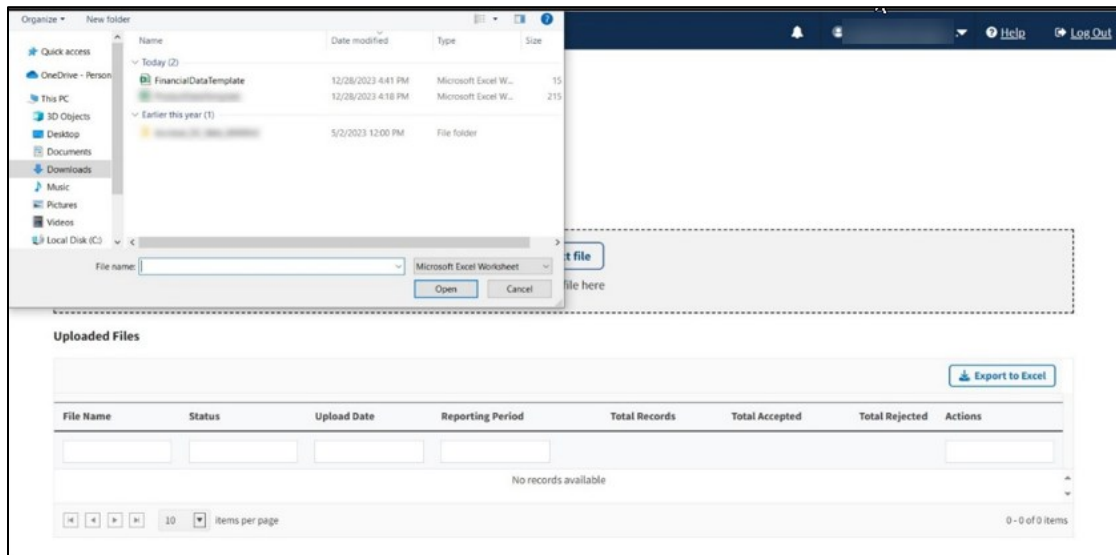
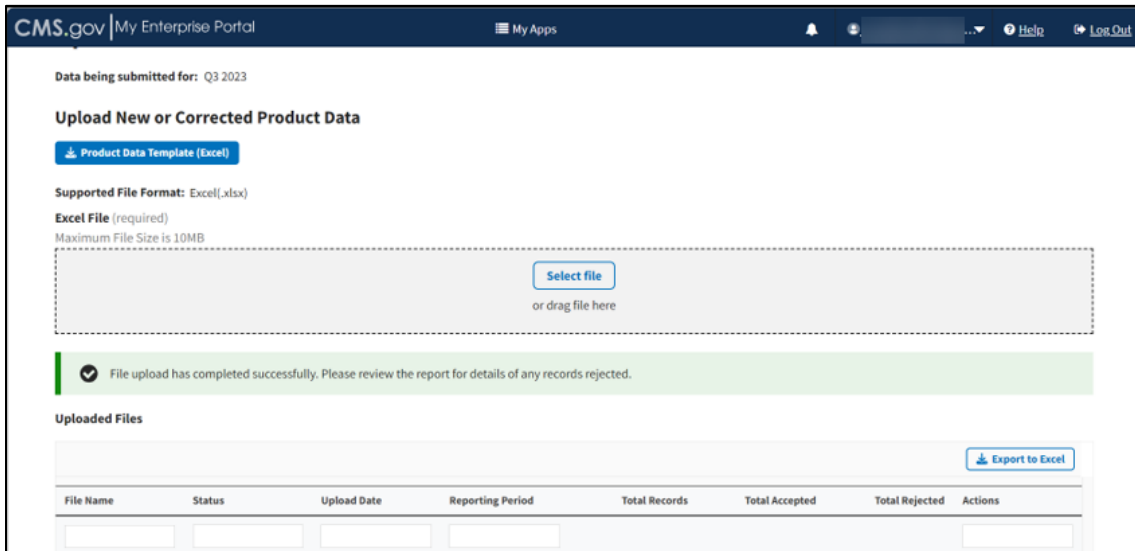


Figure 45: Upload Product Data - Uploading Files from Desktop

A download bar displays as your file uploads. A message displays confirming you have successfully uploaded your **.xlsx file**. Refer to *Figure 46*.

Note: If the Module cannot process your file, an error message displays, and a **New Report** generates under **Uploaded Files**.



Data being submitted for: Q3 2023

Upload New or Corrected Product Data

[Product Data Template \(Excel\)](#)

Supported File Format: Excel(.xlsx)

Excel File (required)
Maximum File Size is 10MB

Select file
or drag file here

File upload has completed successfully. Please review the report for details of any records rejected.

Uploaded Files

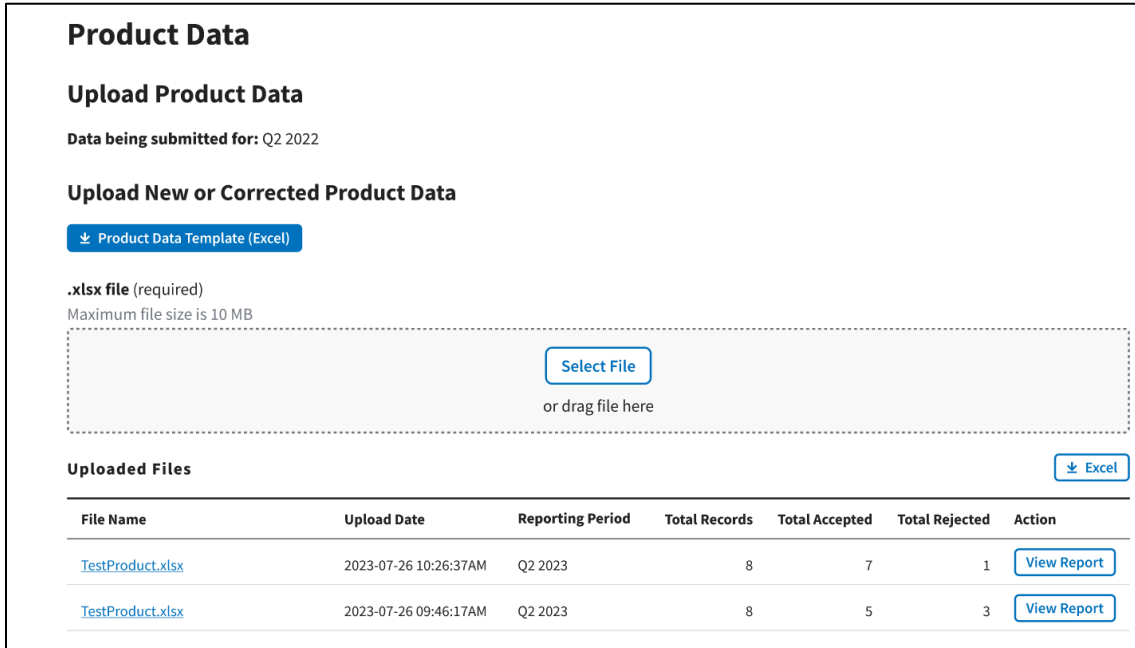
[Export to Excel](#)

File Name	Status	Upload Date	Reporting Period	Total Records	Total Accepted	Total Rejected	Actions
TestProduct.xlsx		2023-07-26 10:26:37AM	Q3 2023	8	7	1	View Report

Figure 46: Upload Product Data - New File Successfully Uploaded

3. Refresh your browser to allow the system to update and display your new file.

The **Uploaded Files** section displays files you uploaded recently as well as previous files still in the Module. Refer to *Figure 47*.



Product Data

Upload Product Data

Data being submitted for: Q2 2022

Upload New or Corrected Product Data

[Product Data Template \(Excel\)](#)

.xlsx file (required)
Maximum file size is 10 MB

Select File
or drag file here

Uploaded Files

[Excel](#)

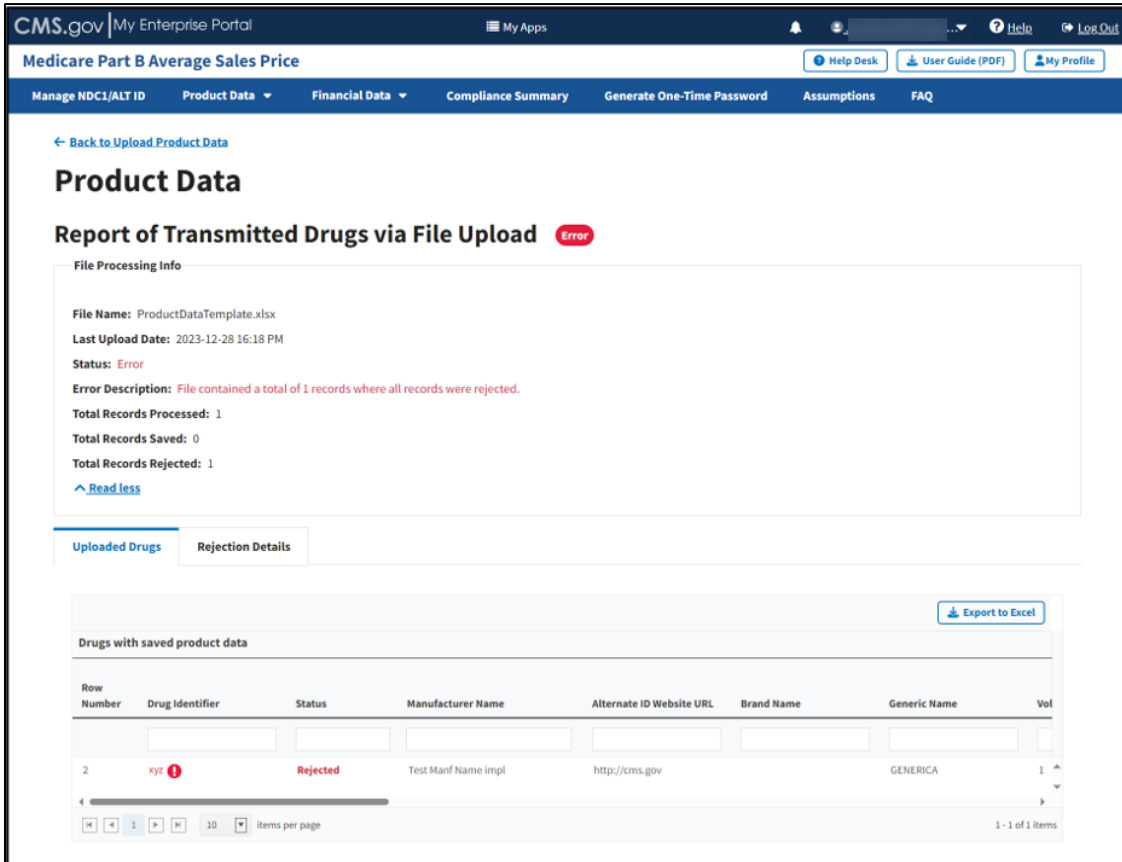
File Name	Upload Date	Reporting Period	Total Records	Total Accepted	Total Rejected	Action
TestProduct.xlsx	2023-07-26 10:26:37AM	Q2 2023	8	7	1	View Report
TestProduct.xlsx	2023-07-26 09:46:17AM	Q2 2023	8	5	3	View Report

Figure 47: Upload Product Data - Uploaded Files

Each uploaded file displays the **File Name**, **Upload Date**, **Reporting Period**, **Total Records**, **Total Accepted**, **Total Rejected**, and **Action** categories submitted to the Module.

4. Click **View Report** under **Action** in the **Uploaded Files** section to view the full report for a submitted file.

The report opens on the next page. Refer to *Figure 48*.


 A screenshot of the CMS.gov Product Data report. The top navigation bar includes links for "My Apps", "Help Desk", "User Guide (PDF)", and "My Profile". The main title is "Medicare Part B Average Sales Price" with a sub-section "Product Data". Below this, a heading "Report of Transmitted Drugs via File Upload" is displayed, with a red "Error" icon. A "File Processing Info" section shows the following details:

- File Name: ProductDataTemplate.xlsx
- Last Upload Date: 2023-12-28 16:18 PM
- Status: Error
- Error Description: File contained a total of 1 records where all records were rejected.
- Total Records Processed: 1
- Total Records Saved: 0
- Total Records Rejected: 1

 Below this, there are two tabs: "Uploaded Drugs" (selected) and "Rejection Details". The "Uploaded Drugs" tab shows a table with one row of data:

Row Number	Drug Identifier	Status	Manufacturer Name	Alternate ID Website URL	Brand Name	Generic Name	Vol
2	xyz ①	Rejected	Test Manf Name impl	http://cms.gov	GENERICA	1	

 The "Rejection Details" tab is not visible in the screenshot.

Figure 48: Upload Product Data - Full Report of Transmitted Drugs via File Upload

5. Click the **Read More** tab under the **Report of Transmitted Drugs via File Upload** to view all **File Processing Information** related to this report.

The report lists all uploaded drugs with saved product data in the ASP system. The Module organizes the full list by row number and includes each drug identifier, status, and all previously submitted information from the **Add Product Data** sections.

Note: The Module highlights errors in red. Hover over the red text to display information about the specific error.

6. Click the **Rejection Details** tab.

A listing of drugs with rejected product data displays. Refer to *Figure 49*.

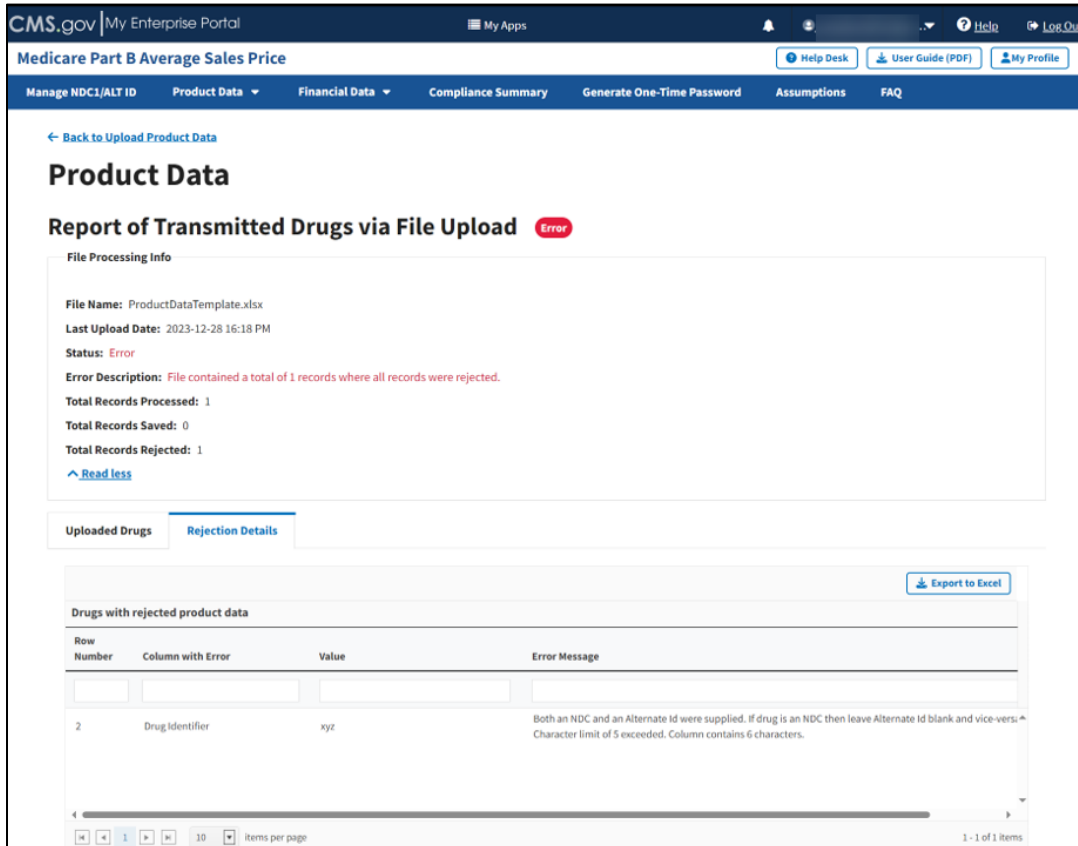

 The screenshot shows a web interface for the CMS ASP Module. The top navigation bar includes links for 'My Apps', 'Help Desk', 'User Guide (PDF)', and 'Log Out'. The main menu has options like 'Manage NDC1/ALT ID', 'Product Data', 'Financial Data', 'Compliance Summary', 'Generate One-Time Password', 'Assumptions', and 'FAQ'. Below the menu, a link '← Back to Upload Product Data' is visible. The main content area is titled 'Product Data' and contains a sub-section 'Report of Transmitted Drugs via File Upload' with an 'Error' status indicator. A 'File Processing Info' box displays the file name 'ProductDataTemplate.xlsx', the last upload date '2023-12-28 16:18 PM', and a status of 'Error'. An error description states: 'File contained a total of 1 records where all records were rejected.' It also shows 'Total Records Processed: 1', 'Total Records Saved: 0', and 'Total Records Rejected: 1'. A link '▲ Read less' is present. Below this, there are two tabs: 'Uploaded Drugs' (selected) and 'Rejection Details'. The 'Rejection Details' tab is currently inactive. A table titled 'Drugs with rejected product data' is shown, with columns for 'Row Number', 'Column with Error', 'Value', and 'Error Message'. One row is listed: Row 2, Column with Error 'Drug Identifier', Value 'xyz', and Error Message 'Both an NDC and an Alternate Id were supplied. If drug is an NDC then leave Alternate Id blank and vice-versa. Character limit of 5 exceeded. Column contains 6 characters.'. At the bottom of the table, there are navigation buttons for page numbers and items per page, and a note '1 - 1 of 1 items'. An 'Export to Excel' button is located at the top right of the table.

Figure 49: Upload Product Data - Reported Rejection Details

The Module lists all errors found in submitted data by **Row Number**, **Column with Error**, and **Error Message** under **Drugs with Rejected Product Data**.

7. Return to the **Add/Update Product Data** section of the Module to request any changes to your product data.

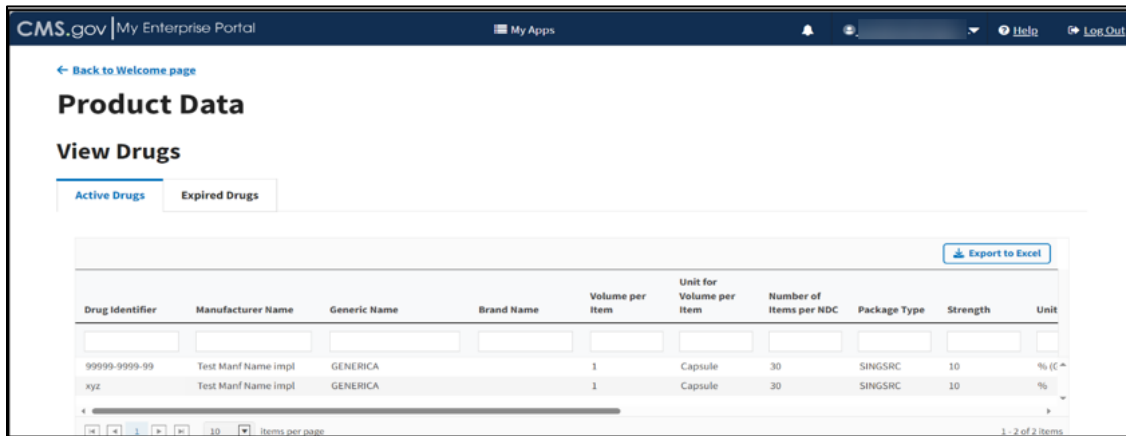
3.2.3 View Drugs

Drug manufacturers can use the ASP module to view drug data submitted during the current reporting period. However, manufacturers cannot update or edit drug data using this feature. From the Medicare Part B Average Sales Price homepage, click the **Product Data** tab; then select the **View Drugs** tab to view the **View Drugs** page.

The following sections describe how to view active and expired drugs.

3.2.3.1 View Active Drugs

From the **View Drugs** page, the **Active Drugs** tab displays by default. Refer to *Figure 50*.


 A screenshot of a web browser showing the CMS.gov My Enterprise Portal. The title bar says "CMS.gov | My Enterprise Portal". The main content area is titled "Product Data" and "View Drugs". There are two tabs: "Active Drugs" (which is selected) and "Expired Drugs". Below the tabs is a table with the following data:

Drug Identifier	Manufacturer Name	Generic Name	Brand Name	Volume per Item	Unit for Volume per Item	Number of Items per NDC	Package Type	Strength	Unit
99999-9999-99	Test Manf Name impl	GENERIC		1	Capsule	30	SINGSRC	10	% (C)
XYZ	Test Manf Name impl	GENERIC		1	Capsule	30	SINGSRC	10	%

 At the bottom of the table, there are navigation arrows and a "1 - 2 of 3 items" message. An "Export to Excel" button is located in the top right corner of the table area.

Figure 50: Product Data - View Active Drugs

Follow these steps to view submitted drug data for **Active Drugs** from the **View Drugs** page:

1. Scroll through the list of active drugs to view submitted data and status. The Module organizes all active drugs by **Drug Identifier**, **Manufacturer Name**, **Generic Name**, **Brand Name**, **Volume per Item**, **Unit for Volume per Item**, **Number of Items per NDC**, **Package Type**, and **Strength** categories, and previously submitted information from the **Add Product Data** sections.
2. Click the arrows on the bottom left to scroll through all submitted drugs by page. View, filter, and sort active drugs by clicking on the category name.

Note: Click the **Export to Excel** button to download all products under the **Compliance Summary**.

3.2.3.2 View Expired Drugs

Follow these steps to view submitted drug data for **Expired Drugs**:

1. From the **View Drugs** page, select the **Expired Drugs** tab. The **Expired Drugs** page opens. Refer to *Figure 51*.

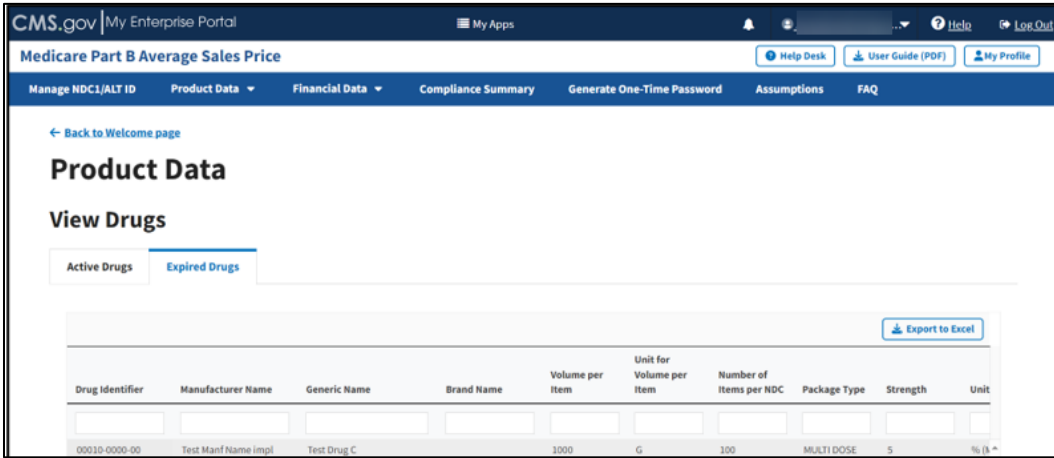

 A screenshot of the CMS.gov Product Data - View Expired Drugs page. The page has a dark blue header with the CMS logo and a "My Enterprise Portal" link. Below the header is a navigation bar with links for "Manage NDC1/ALT ID", "Product Data", "Financial Data", "Compliance Summary", "Generate One-Time Password", "Assumptions", and "FAQ". The main content area is titled "Product Data" and "View Drugs". It shows a table with columns for "Drug Identifier", "Manufacturer Name", "Generic Name", "Brand Name", "Volume per Item", "Unit for Volume per Item", "Number of Items per NDC", "Package Type", "Strength", and "Unit". A row of data is shown: "00010-0000-00", "Test Manf Name impl", "Test Drug C", "1000", "G", "100", "MULTIDOSE", "5", "% (L)".

Figure 51: Product Data - View Expired Drugs

2. Scroll through the list of expired drugs to view submitted data and status.

The Module organizes expired drugs by **Drug Identifier**, **Manufacturer Name**, **Generic Name**, **Brand Name**, **Volume per Item**, **Unit for Volume per Item**, **Number of Items per NDC**, **Package Type**, and **Strength** categories, and previously submitted information from the **Add Product Data** sections.

3. Click the arrows on the bottom left to scroll through all submitted drugs by page. View, filter, and sort active drugs by clicking on the category name.
4. Click the **Export to Excel** button to download all expired drug products.

3.3 Financial Data

Click the **Financial Data** tab on the Medicare Part B Average Sales Price homepage to view the drop-down menu tabs, **Add/Update Financial Data for Current Quarter**, **Upload Financial Data for Current Quarter**, **Restate Financial Data or Add for Prior Quarters**, and **Upload Financial Data for Prior Quarters**. Refer to *Figure 52*.

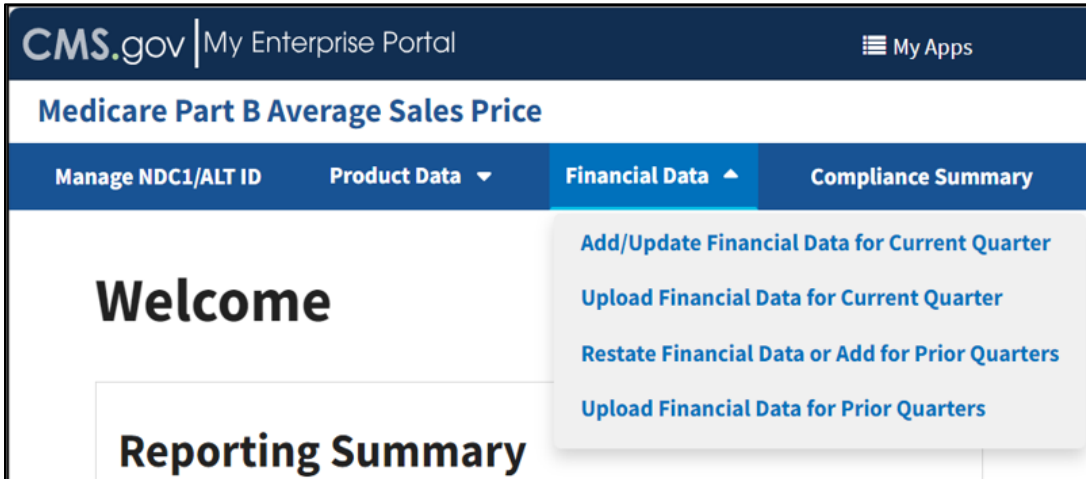

 A screenshot of the CMS.gov Financial Data - Main Drop-down page. The page has a dark blue header with the CMS logo and a "My Enterprise Portal" link. Below the header is a navigation bar with links for "Manage NDC1/ALT ID", "Product Data", "Financial Data", and "Compliance Summary". The main content area is titled "Welcome" and "Reporting Summary". On the right side, there is a list of options: "Add/Update Financial Data for Current Quarter", "Upload Financial Data for Current Quarter", "Restate Financial Data or Add for Prior Quarters", and "Upload Financial Data for Prior Quarters".

Figure 52: Financial Data - Main Drop-down

The following sections describe how to add/update and upload financial data.

3.3.1 Add/Update Financial Data

To add or update financial data, click the **Add/Update Financial Data for Current Quarter** tab.

Note: If you are a manufacturer of certain drugs that contain variable amounts of product, such as radiopharmaceuticals and blood clotting factors, your data should be reported to CMS at the HCPCS level rather than the NDC level. CMS maintains and publishes a list of these drugs on a quarterly basis on the [ASP Reporting page](#) under the **Reporting Resources** section. If you are a manufacturer of a drug that contains variable amounts of product, please check the **“ASP Report in Units Other than NDC”** document prior to submitting your financial data for the quarter. Should you have any questions, please contact sec303aspdata@cms.hhs.gov.

The **Add/Update Financial Data** page opens with default selections. Refer to *Figure 53*.

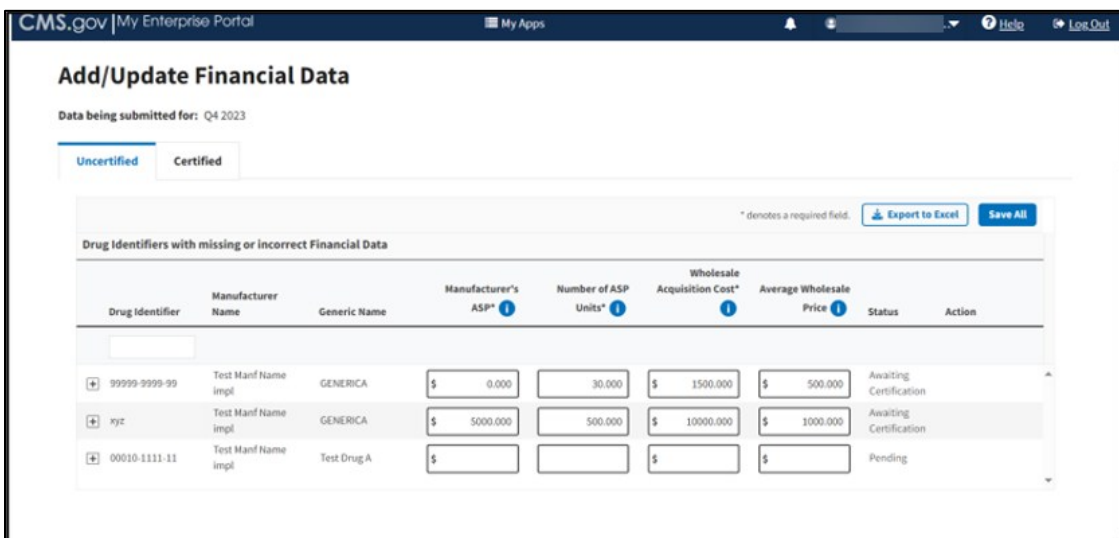


Figure 53: Add/Update Financial Data

Note: The Module collects data submissions for the upcoming financial quarter. As an example, figures in this section feature data submitted for Q1 2023.

The next section describes steps to indicate if your 505(b)(2) products have any therapeutic equivalent changes. If this is not applicable to you, you may skip ahead to the next section, *Section 3.3.1.2 - Add/Update Financial Data for Uncertified Drugs*.

3.3.1.1 Add Therapeutic Equivalent Changes for 505(b)(2) Drugs

If you are associated with any 505(b)(2) products, the system displays a prompt to review the list of products and indicate if any of those products have therapeutic equivalent changes. Refer to *Figure 54*.

Note: If you are associated with any 505(b)(2) products, you must complete these steps before proceeding with adding or editing data. If you are not associated with 505(b)(2) products or are a new user, skip ahead to *Section 3.3.1.2 - Add/Update Financial Data for Uncertified Drugs*.

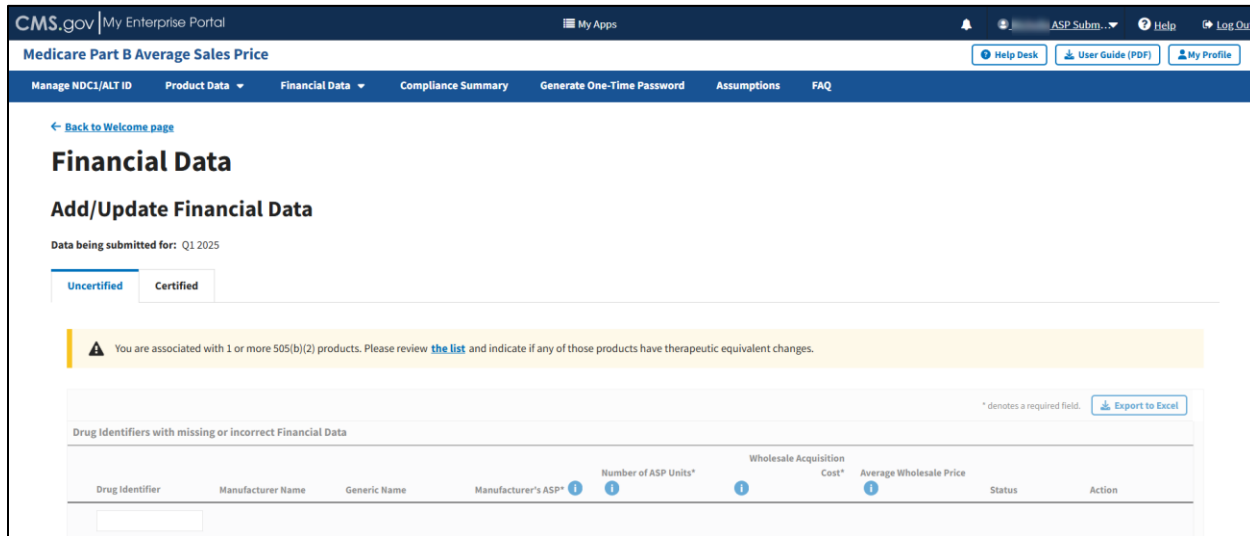

 This screenshot shows the CMS ASP Module Submitter User Guide interface. The top navigation bar includes links for "My Apps", "Help Desk", "User Guide (PDF)", and "Log Out". The main content area is titled "Financial Data" and "Add/Update Financial Data". It displays a table for "Drug Identifiers with missing or incorrect Financial Data". The table columns are: Drug Identifier, Manufacturer Name, Generic Name, Manufacturer's ASP*, Number of ASP Units*, Wholesale Acquisition Cost*, Average Wholesale Price, Status, and Action. A note at the top of the table area states: "⚠ You are associated with 1 or more 505(b)(2) products. Please review [the list](#) and indicate if any of those products have therapeutic equivalent changes." An "Export to Excel" button is located in the top right corner of the table area. The status bar at the bottom of the table indicates that the "Cost*" field is required.

Figure 54: Add/Update Financial Data 505(b)(2)

Follow these steps to review your list of products and indicate any therapeutic equivalent changes:

Click [the list](#) hyperlink in the prompt. Refer to *Figure 54*. The list of products displays.

5. Select **Yes** or **No** in the drop-down menu for each product to indicate whether your product has a therapeutic equivalent. Refer to *Figure 56* and *Figure 56*.

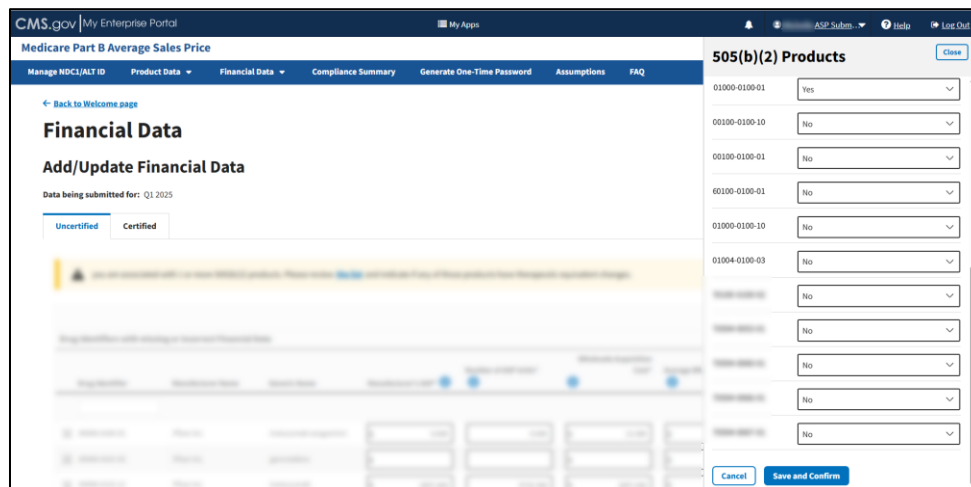
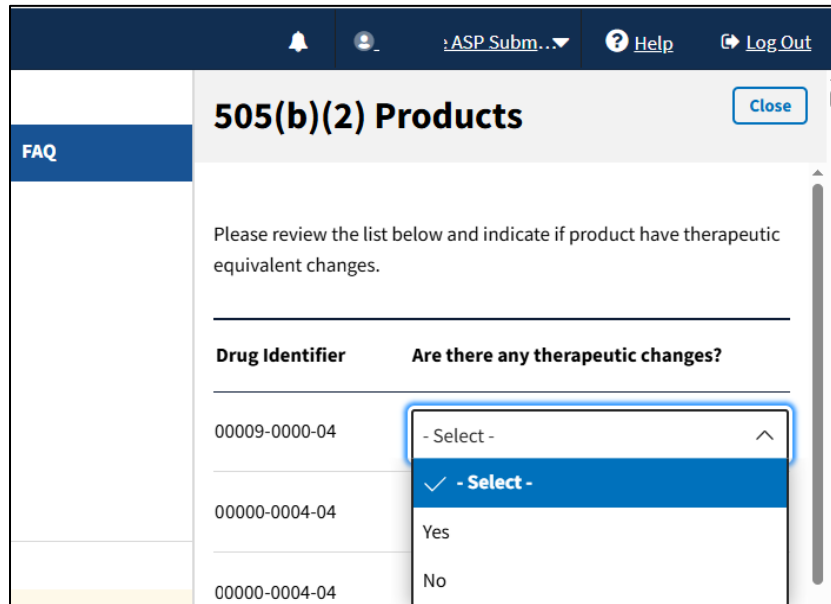

 This screenshot shows the "505(b)(2) Products" list. A modal window is open on the right side of the screen, titled "505(b)(2) Products". It contains a table with two columns: "Product ID" and "Therapeutic Equivalent". The "Product ID" column lists various identifiers such as 01000-0100-01, 00100-0100-10, 00100-0100-01, 60100-0100-01, 01000-0100-10, 01004-0100-03, and 01004-0100-04. The "Therapeutic Equivalent" column contains dropdown menus with the options "Yes" and "No". The "Save and Confirm" button is located at the bottom right of the modal.

Figure 55: Add/Update Financial Data 505(b)(2) Products List



Drug Identifier	Are there any therapeutic changes?
00009-0000-04	<div style="border: 1px solid #ccc; padding: 5px; width: 150px;"> - Select - ✓ - Select - Yes No </div>
00000-0004-04	
00000-0004-04	

Figure 56: Add/Update Financial Data 505(b)(2) Products List

6. Once you have reviewed the list and made your selections, click **Save and Confirm**. A confirmation message displays asking if you have reviewed all of your products.
7. If you have finished reviewing your products, click **Confirm**. Refer to *Figure 57*.

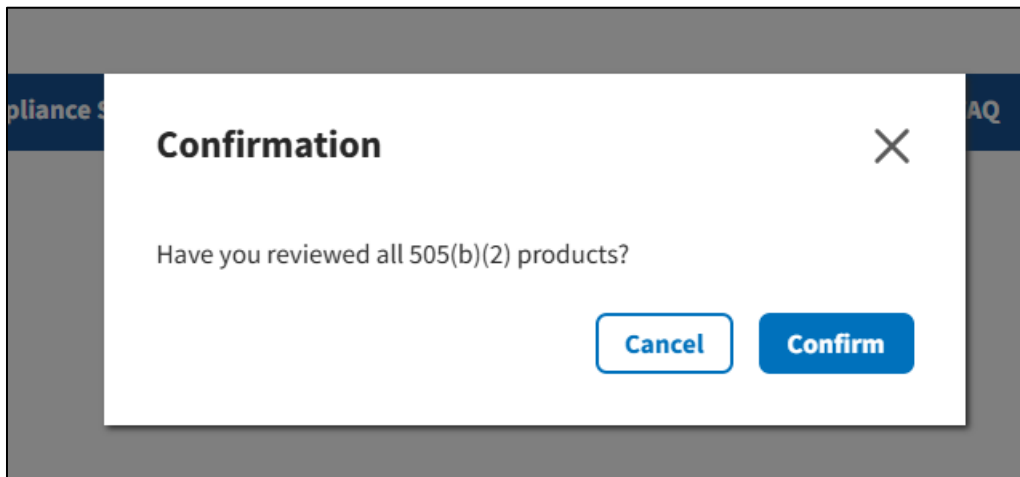


Figure 57: 505(b)(2) Confirmation

A message displays confirming you have successfully updated your therapeutic changes. Refer *Figure 58*.

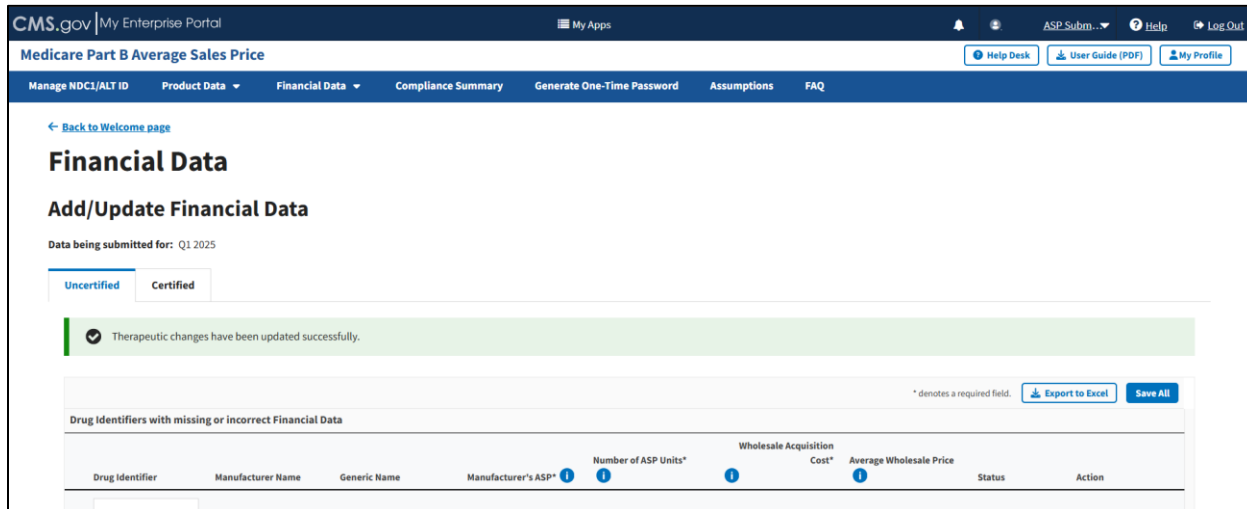

 A screenshot of the CMS ASP Module Submitter User Guide. The top navigation bar includes links for "My Apps", "Help Desk", "User Guide (PDF)", and "My Profile". The main content area is titled "Financial Data" and "Add/Update Financial Data". A message box indicates "Therapeutic changes have been updated successfully." Below this, a table lists "Drug Identifiers with missing or incorrect Financial Data". The columns are: Drug Identifier, Manufacturer Name, Generic Name, Manufacturer's ASP*, Number of ASP Units*, Wholesale Acquisition Cost*, Average Wholesale Price, Status, and Action. The "Manufacturer's ASP*", "Number of ASP Units*", and "Wholesale Acquisition Cost*" fields are marked with an asterisk (*) to denote required fields. Buttons for "Export to Excel" and "Save All" are located at the bottom right of the table.

Figure 58: Add/Update Financial Data 505(b)(2) Successfully Updated

3.3.1.2 Add/Update Financial Data for Uncertified Drugs

Follow these steps to add/update financial data for uncertified drugs:

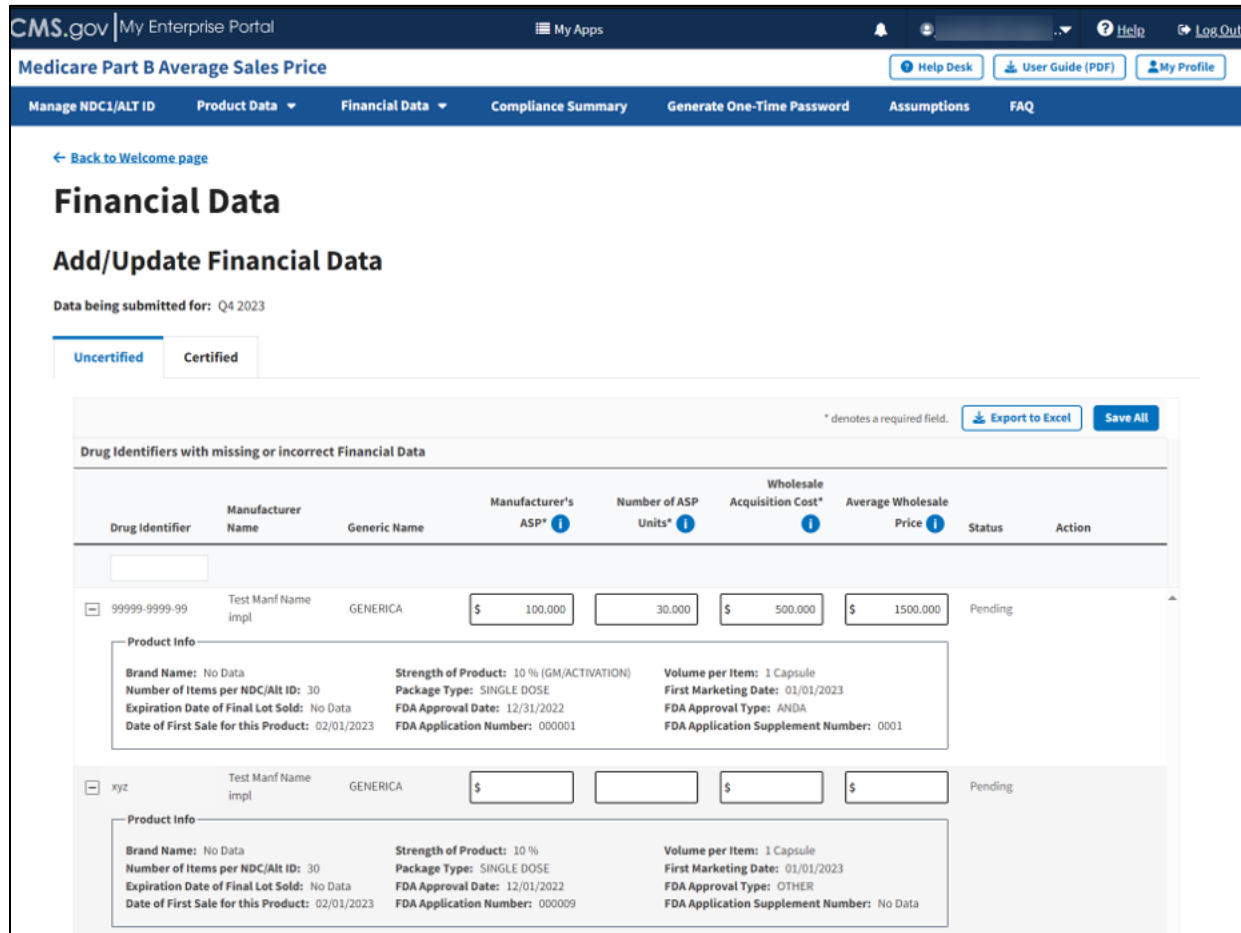
1. From the **Add/Update Financial Data** page, select the **Uncertified Drugs** tab if it is not already selected.

Note: The Module denotes the **Manufacturer's ASP**, **Number of ASP Units**, and **Wholesale Acquisition Cost** fields with an (*) to indicate that each field is required.

2. Enter or edit any missing or inaccurate financial data for your submitted drug products.
3. As you add or update information onto the page, click the **Save All** button to save your changes in the Module.

Note: As an alternative to entering data directly into the Module, under **Drug Identifiers with Missing or Incorrect Financial Data**, you can click the Excel box on the right side to convert all information on this page into an Excel file. You can upload the Excel file after making your updates. Refer to *Section 3.3.3 - Upload Financial Data*.

4. Scroll through the list of submitted drugs and products on the page. Filter through all the information by clicking on the category name.
5. Click the plus symbol on each row of the table to expand each product's information and view additional categories, including **Brand Name**, **FDA Approval** and all other information previously submitted or acknowledged in the **Product Data** section. Refer to *Figure 59*.



Financial Data

Add/Update Financial Data

Data being submitted for: Q4 2023

Uncertified **Certified**

* denotes a required field. [Export to Excel](#) [Save All](#)

Drug Identifiers with missing or incorrect Financial Data

Drug Identifier	Manufacturer Name	Generic Name	Manufacturer's ASP*	Number of ASP Units*	Wholesale Acquisition Cost*	Average Wholesale Price*	Status	Action
99999-9999-99	Test Manf Name impl	GENERIC	\$ 100.000	30.000	\$ 500.000	\$ 1500.000	Pending	
xyz	Test Manf Name impl	GENERIC	\$	\$	\$		Pending	

Product Info

Row 1 (Drug Identifier: 99999-9999-99)

- Brand Name: No Data
- Number of Items per NDC/Alt ID: 30
- Expiration Date of Final Lot Sold: No Data
- Date of First Sale for this Product: 02/01/2023
- Strength of Product: 10 % (GM/ACTIVATION)
- Package Type: SINGLE DOSE
- FDA Approval Date: 12/31/2022
- FDA Application Number: 000001
- Volume per Item: 1 Capsule
- First Marketing Date: 01/01/2023
- FDA Approval Type: ANDA
- FDA Application Supplement Number: 0001

Row 2 (Drug Identifier: xyz)

- Brand Name: No Data
- Number of Items per NDC/Alt ID: 30
- Expiration Date of Final Lot Sold: No Data
- Date of First Sale for this Product: 02/01/2023
- Strength of Product: 10 %
- Package Type: SINGLE DOSE
- FDA Approval Date: 12/01/2022
- FDA Application Number: 000009
- Volume per Item: 1 Capsule
- First Marketing Date: 01/01/2023
- FDA Approval Type: OTHER
- FDA Application Supplement Number: No Data

Figure 59: Add/Update Financial Data - Drug Identifiers With Missing or Incorrect Data

6. Enter and review your information to ensure the highest level of accuracy in data reporting.
7. Click the **Save All** button to submit your information to the Module.

A message displays confirming you have successfully updated your financial data. Refer to *Figure 60*.

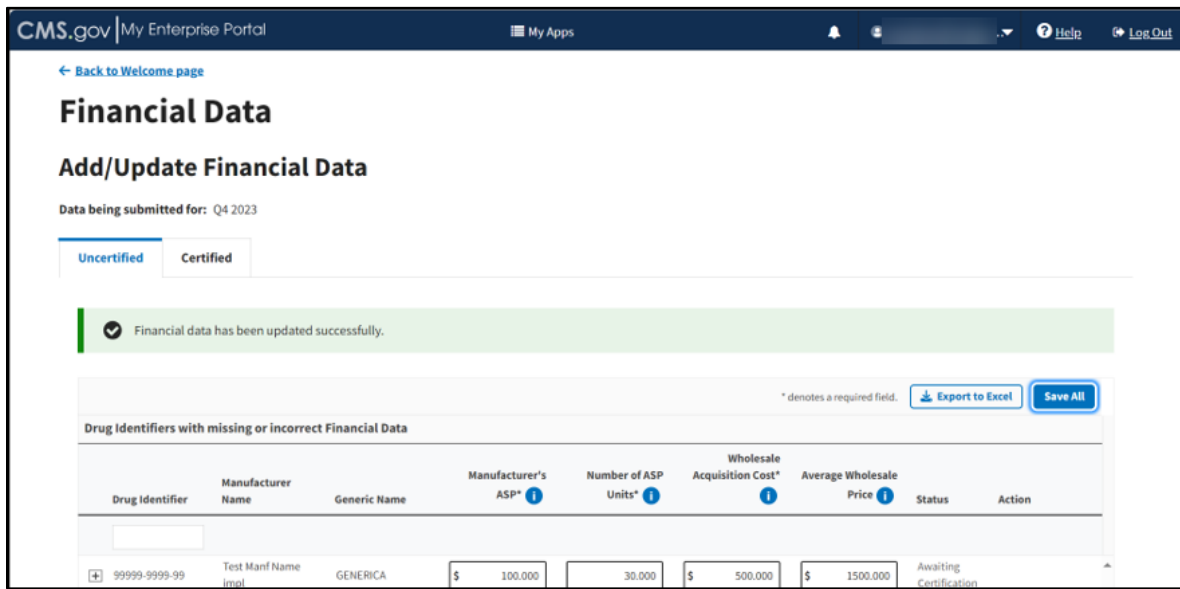


Figure 60: Add/Update Financial Data Successfully Added

Note: When there is an error in the submitted data or a missing field, the page highlights each box in yellow to flag an error.

Each row with errors displays a **View Alerts** button. Refer to *Figure 61*.

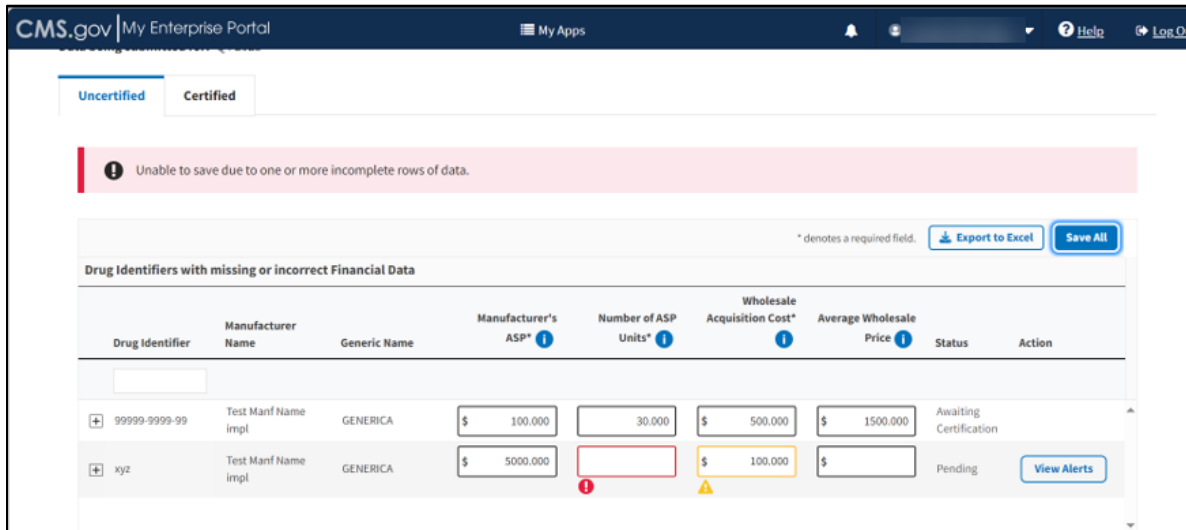


Figure 61: Add/Update Financial Data - Error Menu

- Click the **View Alerts** button for more information regarding the data reporting errors in your submitted financial data.

A side panel opens and displays a listing with descriptions of various errors and warnings. Refer to *Figure 62*.

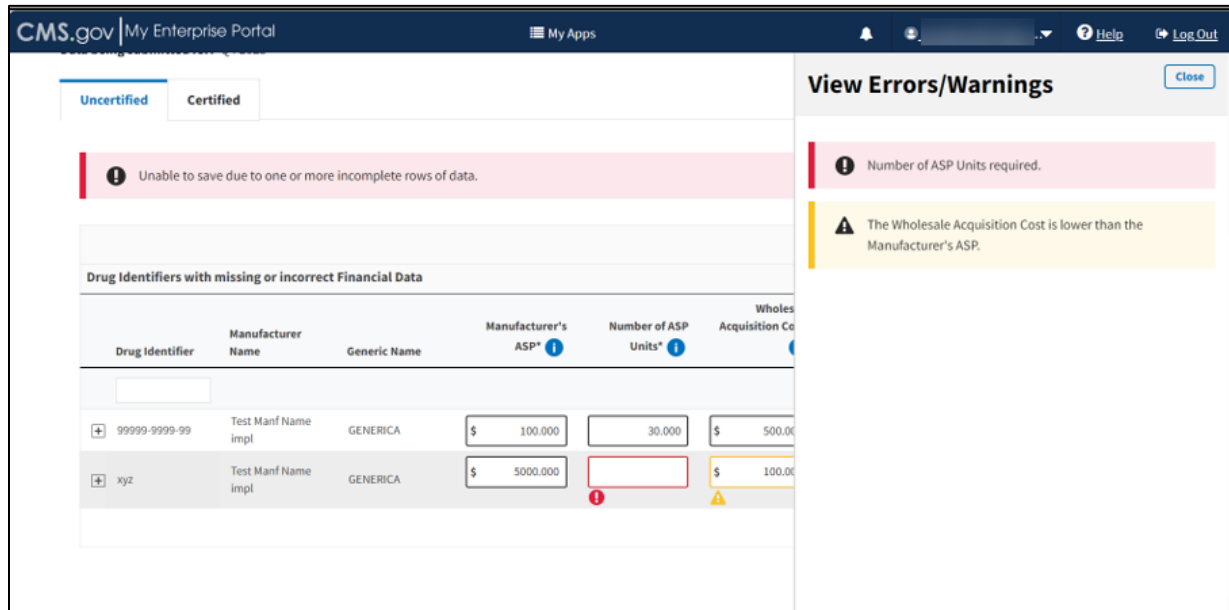


Figure 62: Add/Update Financial Data - View Errors/Warnings Page

- Click **Save Changes** once you address any errors and confirm your product data is accurate.

A message displays confirming that you have successfully added your data. Refer to *Figure 63*.

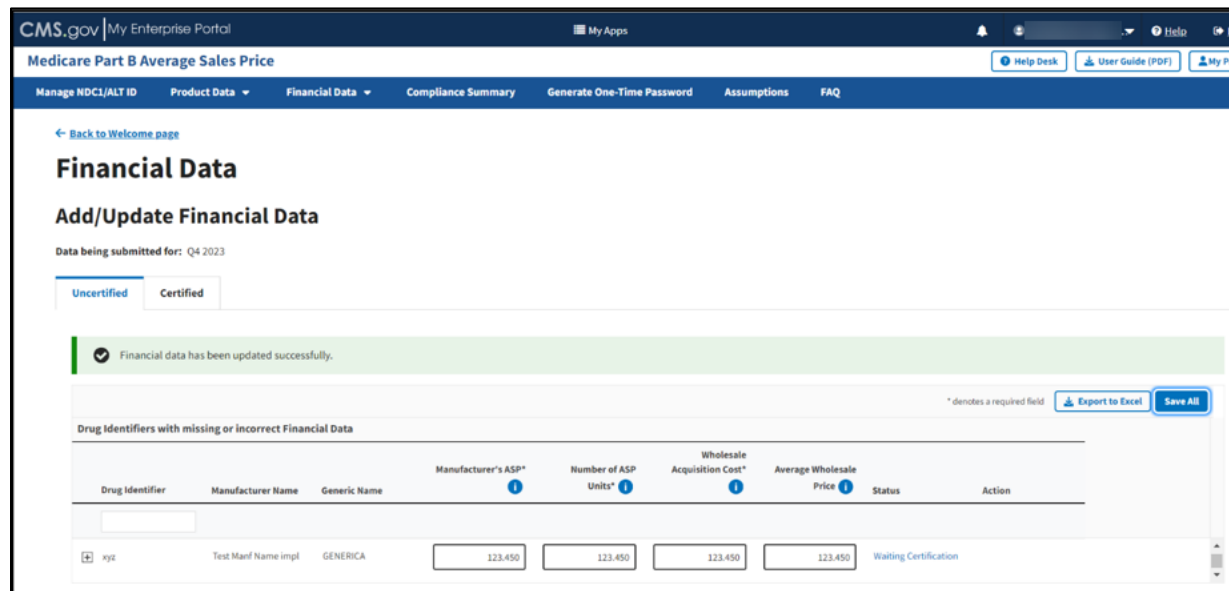


Figure 63: Add/Update Financial Data - Successfully Updated

3.3.2 Add/Update Financial Data - Certified Drugs

The Module collects data submissions for the upcoming financial quarter. Follow these steps to view submitted data for certified drugs:

- From the **Add/Update Financial Data** page, select the **Certified Drugs** tab.

The **Certified Drugs** page opens. Refer to *Figure 64*.

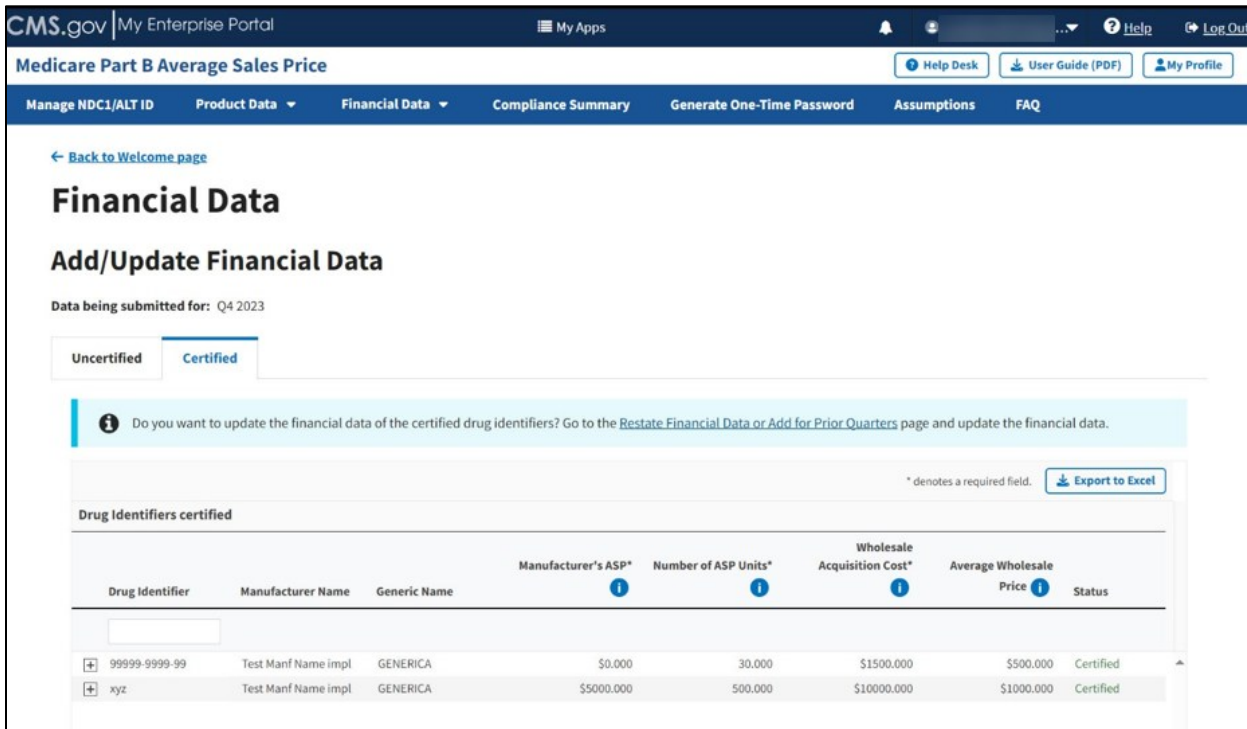

 The screenshot shows the CMS ASP Module interface. At the top, there is a navigation bar with links for 'My Apps', 'Help Desk', 'User Guide (PDF)', and 'My Profile'. Below the navigation bar, the page title is 'Medicare Part B Average Sales Price' and the sub-section is 'Financial Data'. The main content area is titled 'Financial Data' and 'Add/Update Financial Data'. A note at the top says 'Data being submitted for: Q4 2023'. Below this, there are two tabs: 'Uncertified' and 'Certified', with 'Certified' being selected. A callout box with an information icon says: 'Do you want to update the financial data of the certified drug identifiers? Go to the [Restate Financial Data or Add for Prior Quarters](#) page and update the financial data.' A note at the bottom right of the table area says '* denotes a required field.' An 'Export to Excel' button is also present. The main table is titled 'Drug Identifiers certified' and has columns for 'Drug Identifier', 'Manufacturer Name', 'Generic Name', 'Manufacturer's ASP*', 'Number of ASP Units*', 'Wholesale Acquisition Cost*', 'Average Wholesale Price', and 'Status'. Two rows of data are shown: one for '99999-9999-99' and another for 'xyz'. Both rows show 'Test Manf Name impl' as the manufacturer name and 'GENERICA' as the generic name. The 'Manufacturer's ASP*' column shows '\$0.000' and '\$5000.000' respectively. The 'Number of ASP Units*' column shows '30.000' and '500.000'. The 'Wholesale Acquisition Cost*' column shows '\$1500.000' and '\$10000.000'. The 'Average Wholesale Price' column shows '\$500.000' and '\$1000.000'. The 'Status' column shows 'Certified' for both rows.

Figure 64: Add/Update Financial Data - Certified Drugs

Note: To update financial data for Certified drug identifiers, refer to the steps in *Section 3.4.1- Add/Update Restate Financial Data*.

- Under **Drug Identifiers certified**, click the **Export to Excel** button to convert all information on this page into an Excel file.

Note: The Module denotes the **Manufacturer's ASP**, **Number of ASP Units**, and **Wholesale Acquisition Cost** fields with an (*) to indicate that each field is required.

- Scroll through the list of certified drugs and products on the page. Filter through all the information by clicking on your preferred category name.
- Click the arrows on the bottom left to scroll through all submitted drugs by page.
- Click on the plus symbol on each row of the table to expand each product's information and view additional categories, including **Brand Name**, **FDA Approval** and all other information previously submitted or acknowledged in the **Product Data** section. Refer to *Figure 65*.

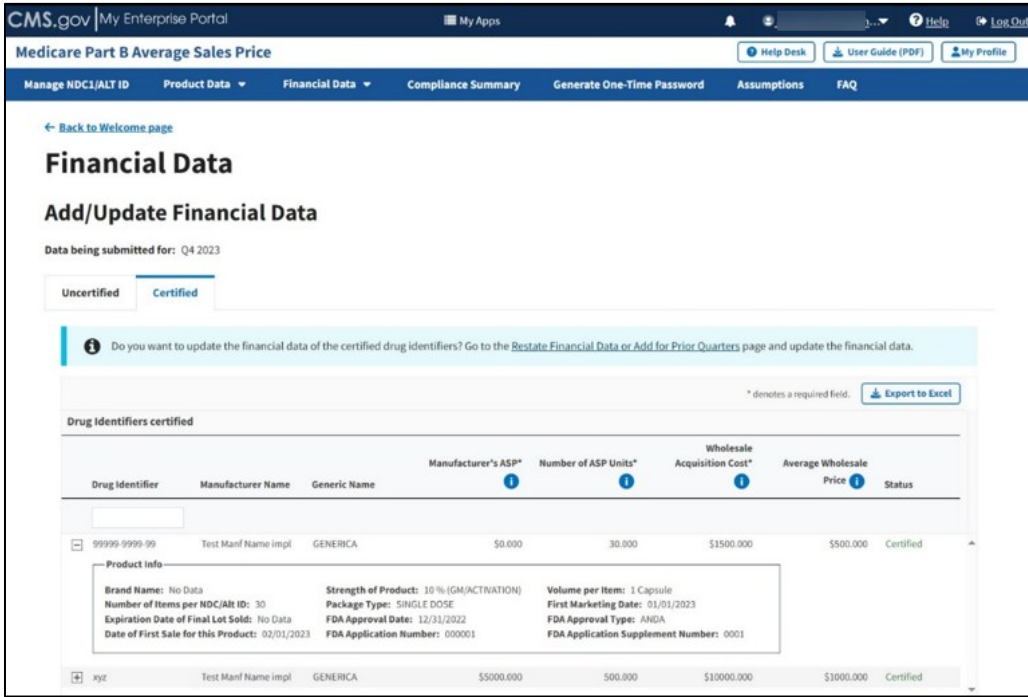

 This screenshot shows the 'Financial Data' section of the Medicare Part B Average Sales Price application. The top navigation bar includes links for 'Manage NDC1/ALT ID', 'Product Data', 'Financial Data', 'Compliance Summary', 'Generate One-Time Password', 'Assumptions', and 'FAQ'. The 'Financial Data' tab is selected. A sub-menu bar at the top of the page includes 'Back to Welcome page', 'Financial Data', 'Add/Update Financial Data', 'More Information', 'Upload Financial Data for Current Quarter', 'Upload New or Corrected Financial Data', and 'View Reports'. The main content area is titled 'Financial Data' and 'Add/Update Financial Data'. It shows a table for 'Drug Identifiers certified' with columns for Drug Identifier, Manufacturer Name, Generic Name, Manufacturer's ASP*, Number of ASP Units*, Wholesale Acquisition Cost*, Average Wholesale Price, and Status. A note at the top of the table area says, 'Do you want to update the financial data of the certified drug identifiers? Go to the [Restate Financial Data or Add for Prior Quarters](#) page and update the financial data.' An 'Export to Excel' button is located in the top right corner of the table area. Below the table, a 'Product Info' box displays detailed product information: Brand Name: No Data, Strength of Product: 10 % (GM/ACT/NATION), Volume per Item: 1 Capsule; Number of Items per NDC/Alt ID: 30, Package Type: SINGLE DOSE, First Marketing Date: 01/01/2023; Expiration Date of Final Lot Sold: No Data, FDA Approval Date: 12/31/2022, FDA Approval Type: ANDA; Date of First Sale for this Product: 02/01/2023, FDA Application Number: 000001, FDA Application Supplement Number: 00001.

Figure 65: Add/Update Financial Data - Certified Drugs More Information

3.3.3 Upload Financial Data

Follow these steps to upload financial data:

1. From the Medicare Part B Average Sales Price homepage, click the **Financial Data** tab; then select the **Upload Financial Data for Current Quarter** tab.
2. The **Upload New or Corrected Financial Data** page opens, listing the financial quarter and year for the upcoming reporting period. Refer to *Figure 66*.

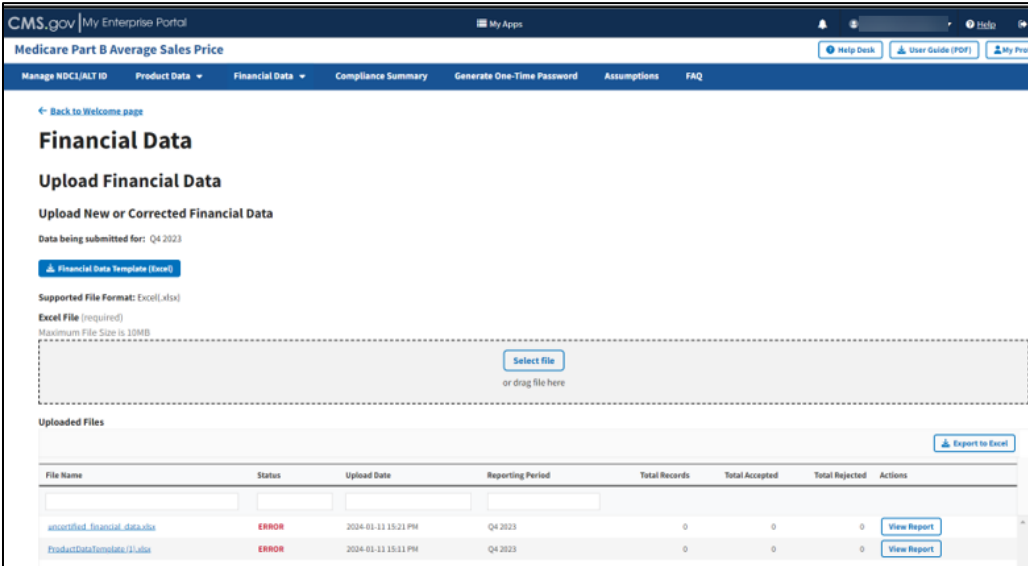

 This screenshot shows the 'Upload Financial Data' section of the Medicare Part B Average Sales Price application. The top navigation bar includes links for 'Manage NDC1/ALT ID', 'Product Data', 'Financial Data', 'Compliance Summary', 'Generate One-Time Password', 'Assumptions', and 'FAQ'. The 'Financial Data' tab is selected. A sub-menu bar at the top of the page includes 'Back to Welcome page', 'Financial Data', 'Add/Update Financial Data', 'More Information', 'Upload Financial Data for Current Quarter', 'Upload New or Corrected Financial Data', and 'View Reports'. The main content area is titled 'Upload Financial Data' and 'Upload New or Corrected Financial Data'. It shows a table for 'Uploaded Files' with columns for File Name, Status, Upload Date, Reporting Period, Total Records, Total Accepted, Total Rejected, and Actions. A note at the top of the table area says, 'Data being submitted for: Q4 2023'. A 'Financial Data Template (Excel)' button is located in the top right corner of the table area. Below the table, a 'Select file' button and a 'or drag file here' text input field are shown. The table data shows two files: 'uncertified_financial_data.xlsx' and 'ProductDataTemplate11.xlsx', both with an 'ERROR' status, uploaded on 2024-01-11 at 15:21 PM, and reporting for Q4 2023. Each file has a 'View Report' button in the 'Actions' column.

Figure 66: Upload Product Data - New or Corrected

Note: Under Data being submitted for: (current quarter), click **Financial Data Template (Excel)** to download a financial data template.

3. If you are associated with any 505(b)(2) products, you may be prompted to indicate if your products have any therapeutic equivalent changes before you can proceed. Refer to *Section 3.3.1.1 - Add Therapeutic Equivalent Changes for 505(b)(2) Drugs* for more information.
4. Upon preparing your **.xlsx file (required)** and verifying your information for accuracy, click **Select File** to browse your desktop and upload the file to the Module. You may also drag the file into the **Select File** box. Refer to *Figure 67*.

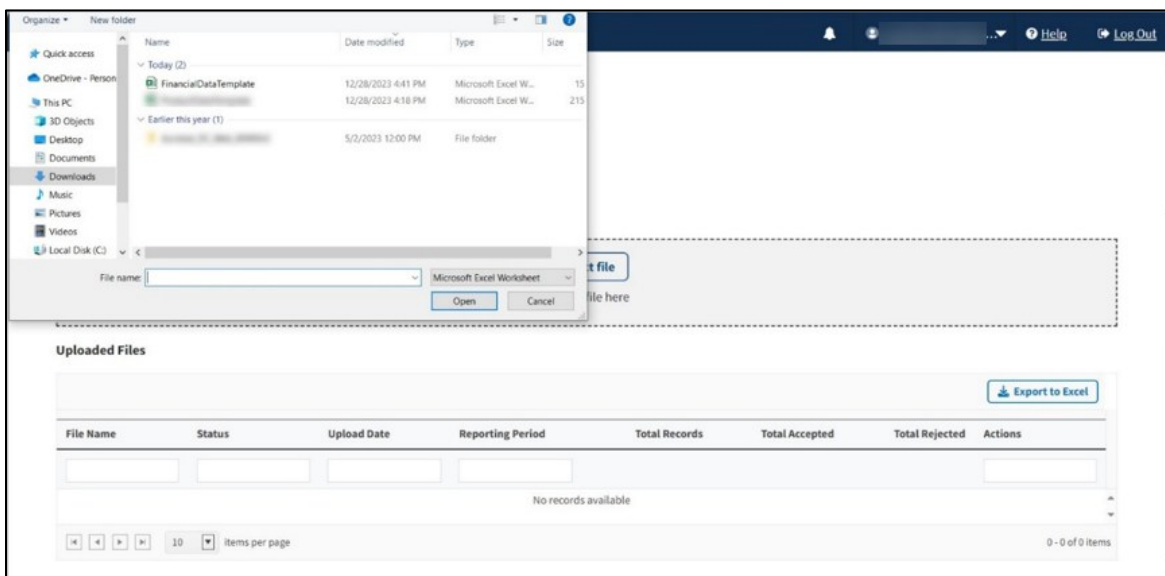
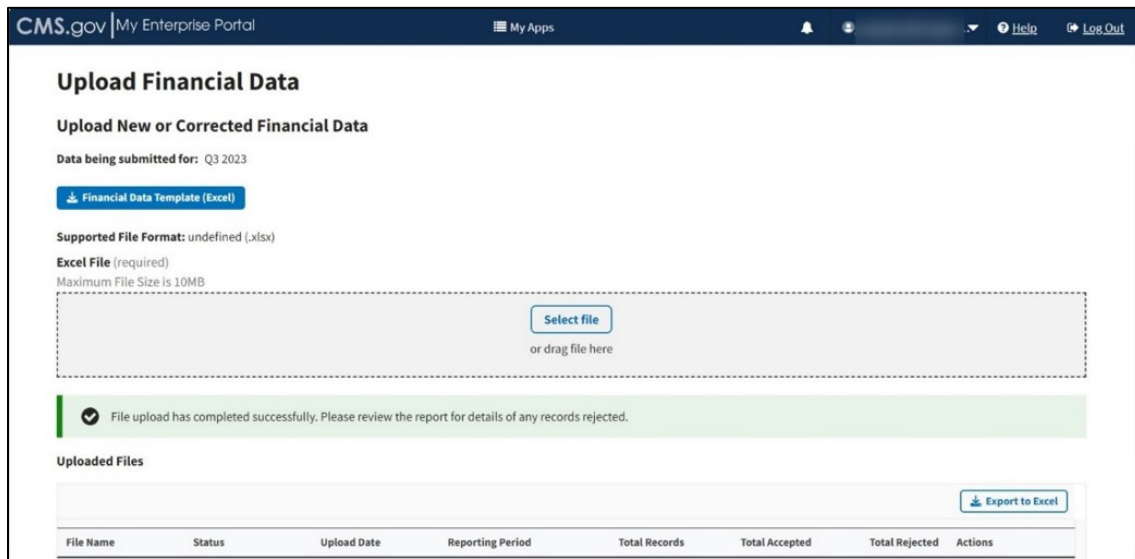


Figure 67: Upload Financial Data - Uploading Files From Desktop

A download bar displays as your file uploads. A message displays confirming you have successfully uploaded your **.xlsx file**. Refer to *Figure 68*.

Note: If the Module cannot process your file, an error message displays, and a **New Report** generates under **Uploaded Files**.



Upload Financial Data

Upload New or Corrected Financial Data

Data being submitted for: Q3 2023

[Financial Data Template \(Excel\)](#)

Supported File Format: undefined (.xlsx)

Excel File (required)
Maximum File Size is 10MB

Select file
or drag file here

File upload has completed successfully. Please review the report for details of any records rejected.

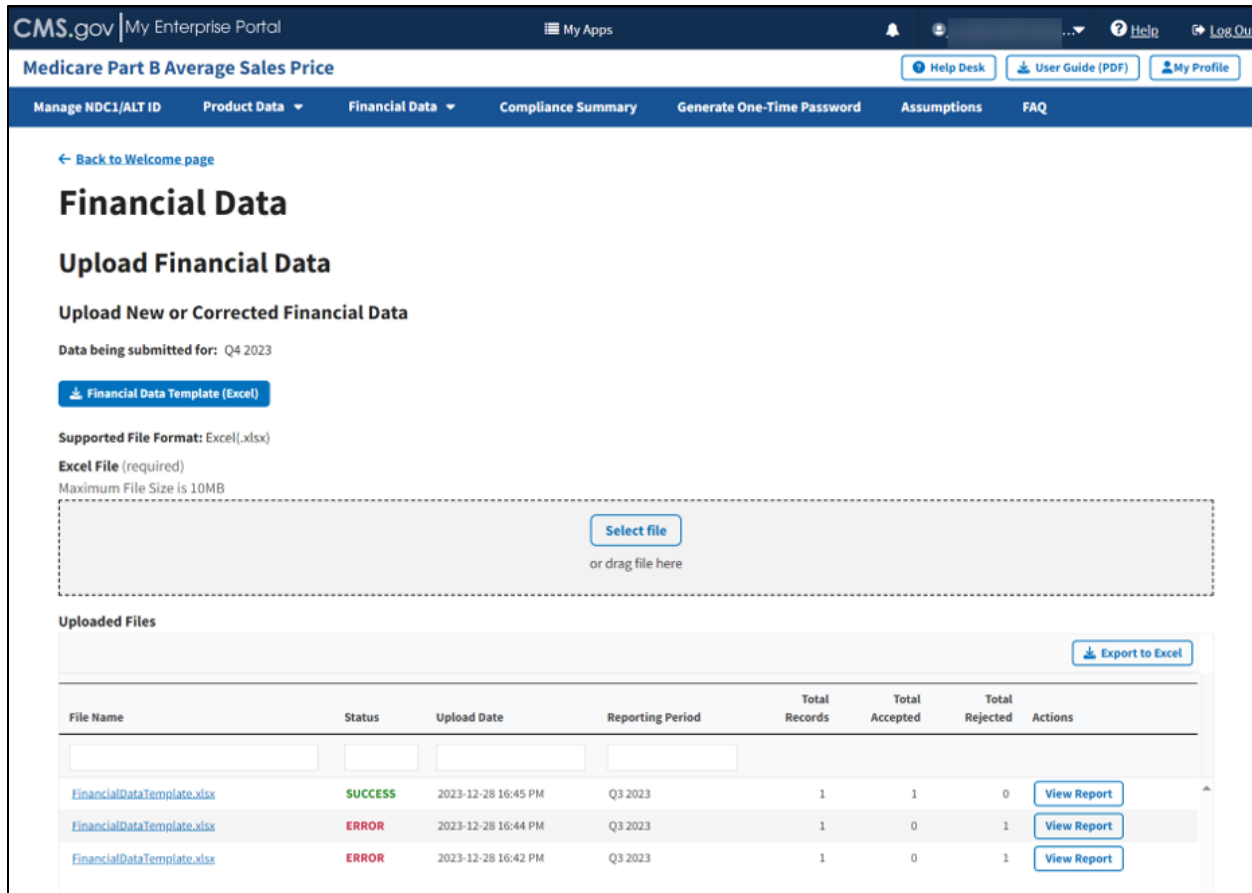
Uploaded Files

File Name	Status	Upload Date	Reporting Period	Total Records	Total Accepted	Total Rejected	Actions
FinancialDataTemplate.xlsx	SUCCESS	2023-12-28 16:45 PM	Q3 2023	1	1	0	View Report
FinancialDataTemplate.xlsx	ERROR	2023-12-28 16:44 PM	Q3 2023	1	0	1	View Report
FinancialDataTemplate.xlsx	ERROR	2023-12-28 16:42 PM	Q3 2023	1	0	1	View Report

[Export to Excel](#)

Figure 68: Upload Financial Data Page - New File Successfully Uploaded

5. The **Uploaded Files** section displays files you uploaded recently as well as previous files still in the Module. Refer to *Figure 69*.



Financial Data

Upload Financial Data

Upload New or Corrected Financial Data

Data being submitted for: Q4 2023

[Financial Data Template \(Excel\)](#)

Supported File Format: Excel(.xlsx)

Excel File (required)
Maximum File Size is 10MB

Select file
or drag file here

Uploaded Files

File Name	Status	Upload Date	Reporting Period	Total Records	Total Accepted	Total Rejected	Actions
FinancialDataTemplate.xlsx	ERROR	2023-12-28 16:45 PM	Q3 2023	1	0	1	View Report
FinancialDataTemplate.xlsx	ERROR	2023-12-28 16:44 PM	Q3 2023	1	0	1	View Report
FinancialDataTemplate.xlsx	ERROR	2023-12-28 16:42 PM	Q3 2023	1	0	1	View Report

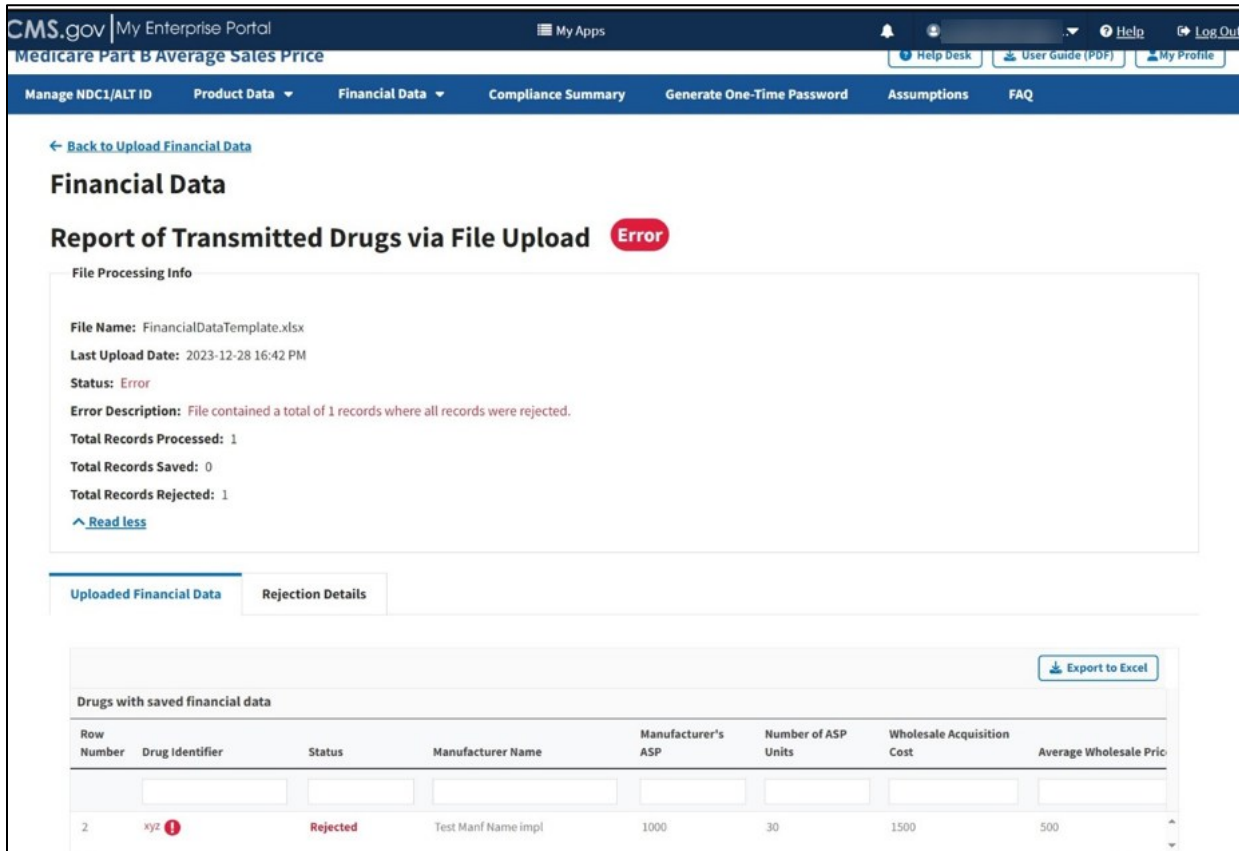
[Export to Excel](#)

Figure 69: Upload Financial Data - Uploaded Files

Each uploaded file displays the **File Name**, **Status**, **Upload Date**, **Reporting Period**, **Total Records**, **Total Accepted**, **Total Rejected**, and **Actions** categories submitted to the Module.

6. Click **View Report** under **Actions** in the **Uploaded Files** section to view the full report for a submitted file.

The report opens on the next page. Refer to *Figure 70*.



The screenshot shows the CMS ASP Module interface. At the top, there is a navigation bar with links for "My Apps", "Help Desk", "User Guide (PDF)", and "My Profile". Below the navigation bar, the main content area is titled "Financial Data" and "Report of Transmitted Drugs via File Upload". A red "Error" button is visible next to the title. The "File Processing Info" section contains the following details:

- File Name:** FinancialDataTemplate.xlsx
- Last Upload Date:** 2023-12-28 16:42 PM
- Status:** Error
- Error Description:** File contained a total of 1 records where all records were rejected.
- Total Records Processed:** 1
- Total Records Saved:** 0
- Total Records Rejected:** 1

Below this section are two tabs: "Uploaded Financial Data" (selected) and "Rejection Details". The "Rejection Details" tab is currently inactive. The "Uploaded Financial Data" tab displays a table titled "Drugs with saved financial data". The table has the following columns: Row Number, Drug Identifier, Status, Manufacturer Name, Manufacturer's ASP, Number of ASP Units, Wholesale Acquisition Cost, and Average Wholesale Price. One row is visible, showing a drug with Row Number 2, Drug Identifier "xyz", Status "Rejected", Manufacturer Name "Test Manf Name impl", Manufacturer's ASP 1000, Number of ASP Units 30, Wholesale Acquisition Cost 1500, and Average Wholesale Price 500. An "Export to Excel" button is located at the top right of the table.

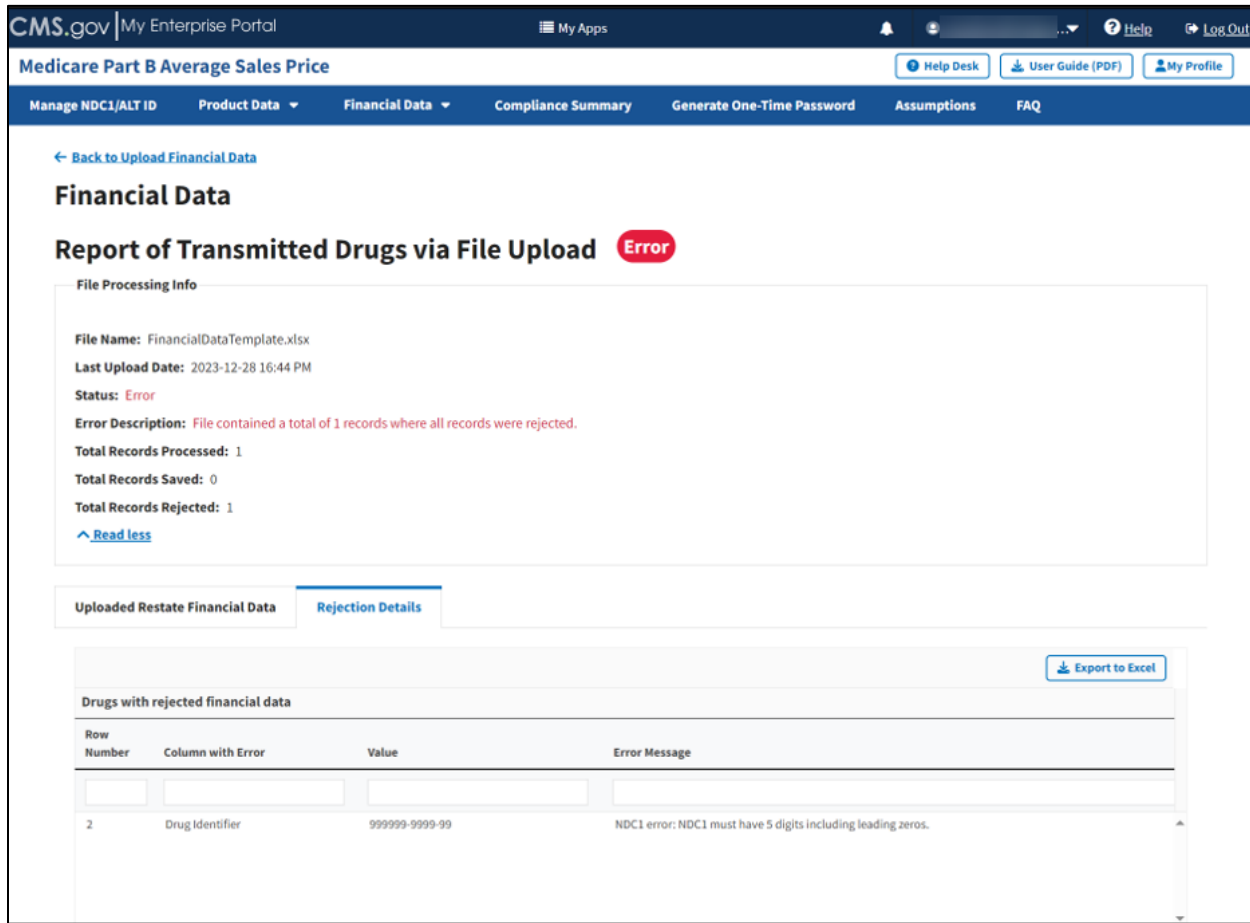
Figure 70: Upload Financial Data - Report of Transmitted Drugs via File Upload

The report lists all drugs with saved product data in the ASP system. The Module organizes the full list by row number and includes each drug identifier, status, and all previously submitted information from the **Add Product Data** sections.

Note: The Module highlights errors in red. Hover over the red text to display information about the specific error.

7. Click the **Rejection Details** tab.

A listing of drug identifiers with rejected financial data displays. Refer to *Figure 71*.


 A screenshot of the CMS ASP Module's 'Financial Data' section. At the top, a red 'Error' button is visible next to the title 'Report of Transmitted Drugs via File Upload'. Below this, the 'File Processing Info' section displays the following details:

- File Name: FinancialDataTemplate.xlsx
- Last Upload Date: 2023-12-28 16:44 PM
- Status: Error
- Error Description: File contained a total of 1 records where all records were rejected.
- Total Records Processed: 1
- Total Records Saved: 0
- Total Records Rejected: 1

 A link to 'Read less' is also present. Below this, there are two tabs: 'Uploaded Restate Financial Data' (selected) and 'Rejection Details'. The 'Rejection Details' tab shows a table titled 'Drugs with rejected financial data' with one row of data:

Row Number	Column with Error	Value	Error Message
2	Drug Identifier	999999-9999-99	NDC1 error: NDC1 must have 5 digits including leading zeros.

 An 'Export to Excel' button is located at the top right of this table.

Figure 71: Upload Financial Data - Reported Rejection Details

The Module lists all errors found in submitted data by **Row Number**, **Column with Error**, **Value** and **Error Message** under **Drugs with rejected financial data**.

8. Return to the **Add/Update Financial Data for Current Quarter** section of the Module to request any changes to your product data.

3.4 Restating Financial Data

Manufacturers of drugs and biologicals payable under Medicare Part B have an obligation to report accurate ASP data to CMS, including addressing data miscalculations and other errors in previously submitted data. Upon identifying an error, manufacturers must submit corrected data through the ASP Module. Additionally, CMS may identify an error and contact the manufacturer to request corrected data for prior quarters.

CMS evaluates resubmitted data and decides whether to issue a restatement of the payment limit. Criteria evaluated includes, but is not limited to, timing of the corrected data, changes to the payment limit, and/or administrative burden.

The following sections describe how to add/update or upload restate financial data using the online data entry process.

3.4.1 Add/Update Restate Financial Data

Follow these steps to add/update restate financial data:

1. Click the **Financial Data** tab; select **Restate Financial Data or Add for Prior Quarters**. Refer to *Figure 72*.

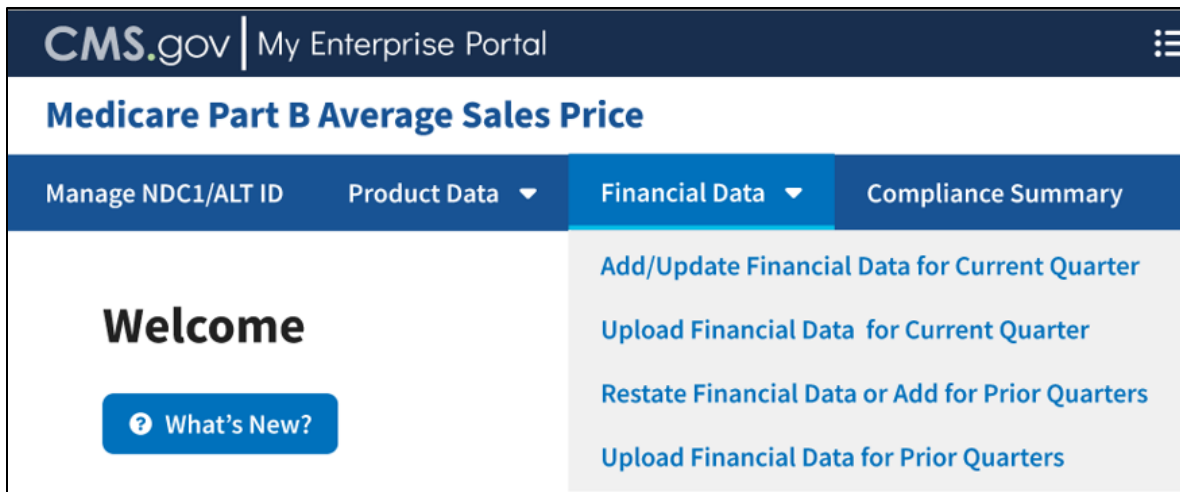


Figure 72: Financial Data - Main Dropdown

The **Restate Financial Data or Add for Prior Quarters** page opens, listing the financial quarter and year for the upcoming reporting period. Refer to *Figure 73*.

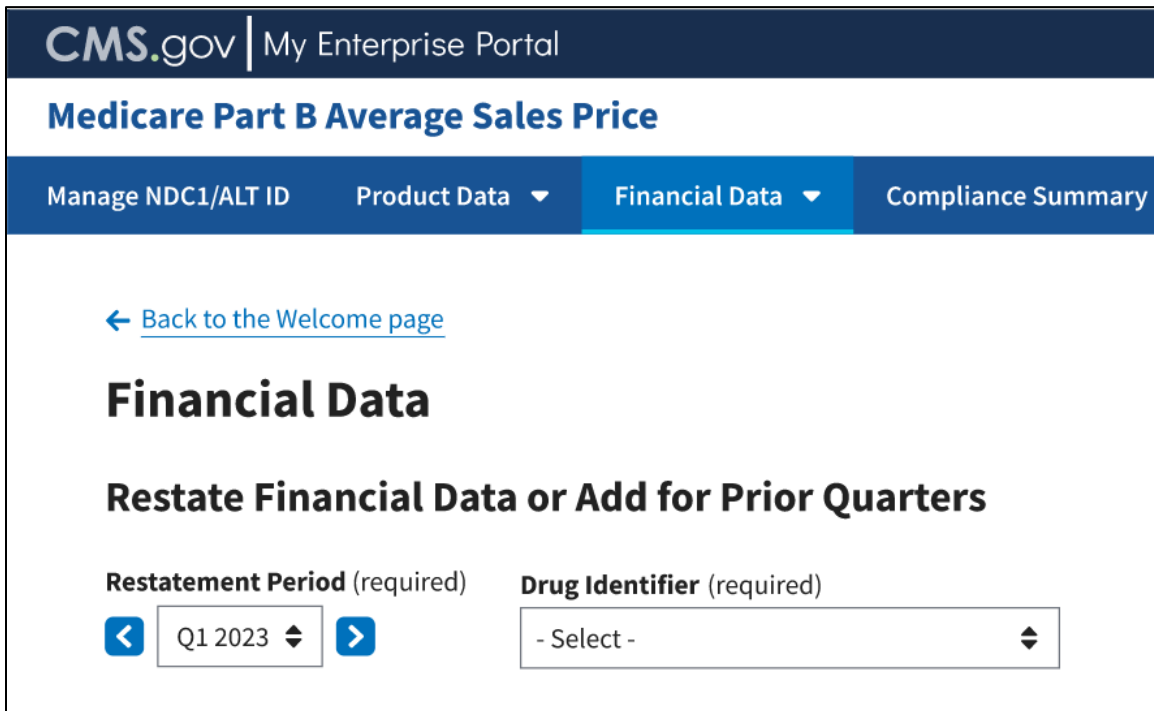


Figure 73: Financial Data - Add/Update Restate Financial Data

Note: Click the **Restatement Period (required)** drop-down in the top left to scroll through previous quarters. Click the blue arrows to navigate to a previous quarter starting with the most recent or next quarter.

2. Click the **-Select-** box under **Drug Identifier (required)** to expand the list of submitted drugs in the Module. Refer to *Figure 74*.

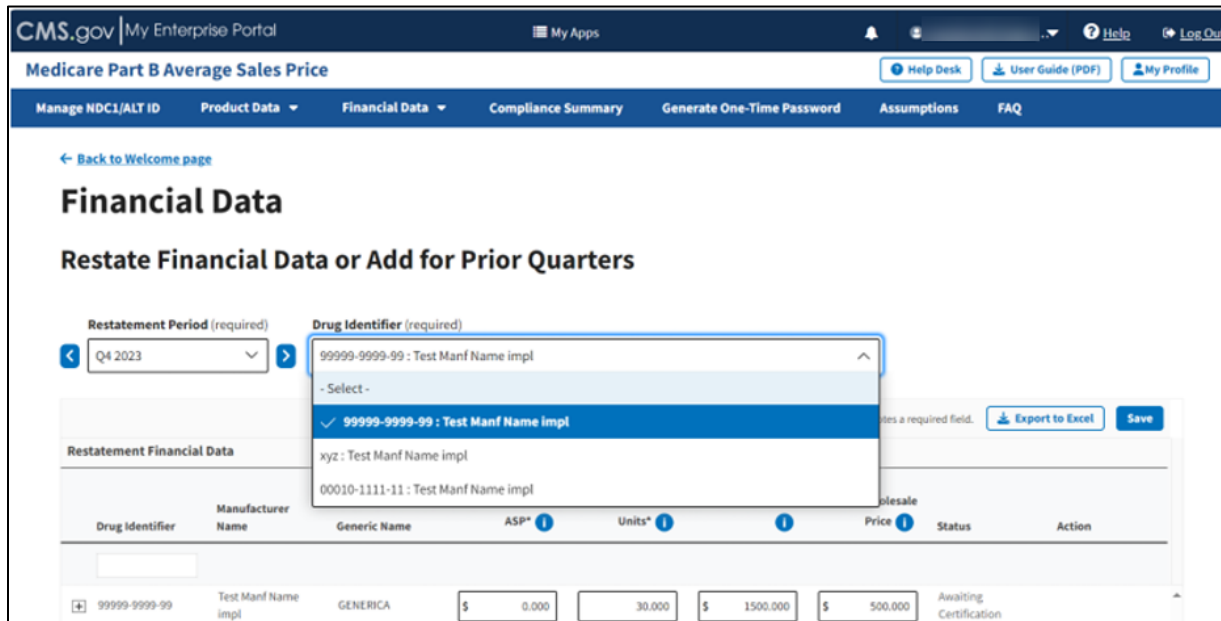


Figure 74: Add/Update Restate Financial Data - Drug Identifier Drop-down

Select the **Drug Identifier** you need to close the drop-down. Once you click a product, the **Review Restatement List** expands to show the selected restatement. Refer to *Figure 75*.

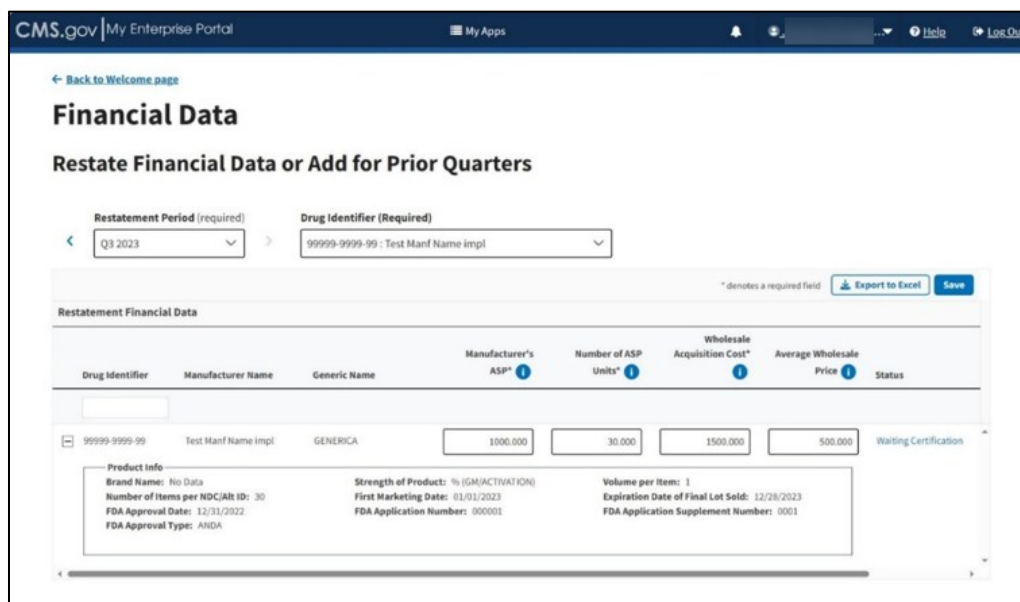
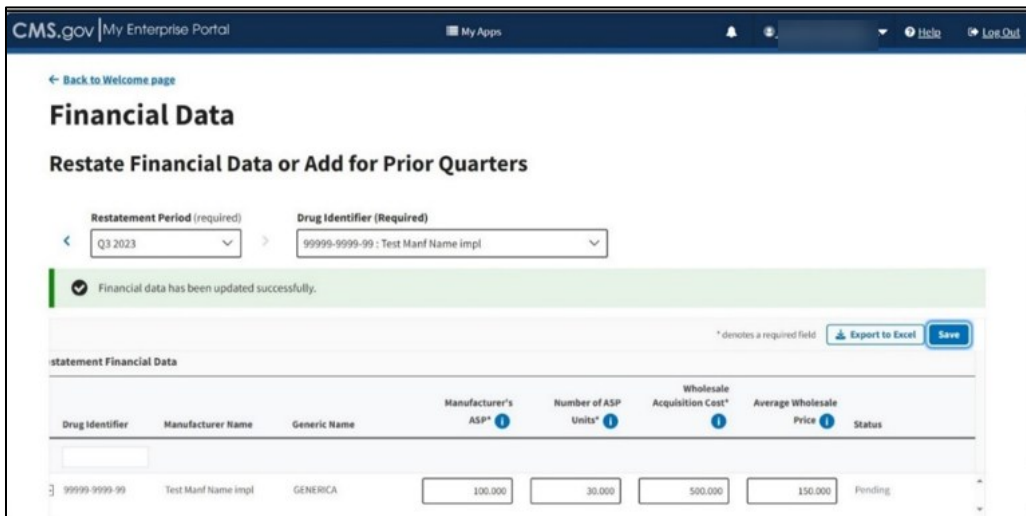


Figure 75: Add/Update Restate Page - Review Restatement List

3. Review and make any corrections necessary for the drug to the **Manufacturer's ASP**, **Number of ASP Units**, **Wholesale Acquisition Cost (all required)** and **Average Wholesale Price** fields.
4. Click the **plus** symbol on each row of the table to expand each product's information and view additional categories previously submitted or acknowledged in the **Product Data** section.
5. Click the **Save** button to submit your data.

A message displays confirming you have successfully updated your **Restate Financial Data**. Refer to *Figure 76*.



Drug Identifier	Manufacturer Name	Generic Name	Manufacturer's ASP*	Number of ASP Units*	Wholesale Acquisition Cost*	Average Wholesale Price*	Status
99999-9999-99	Test Manf Name Impl	GENERIC	100.000	30.000	500.000	150.000	Pending

Figure 76: Add/Update Restate Page - Restate Data Successfully Saved

6. Contact your Certifier to recertify the corrected data you submitted to the Module.

3.4.2 Upload Restate Financial Data

Follow these steps to upload restate financial data:

1. From the Medicare Part B Average Sales Price homepage, click the **Financial Data** tab; then select **Upload Financial Data for Prior Quarters**. Refer to *Figure 77*.

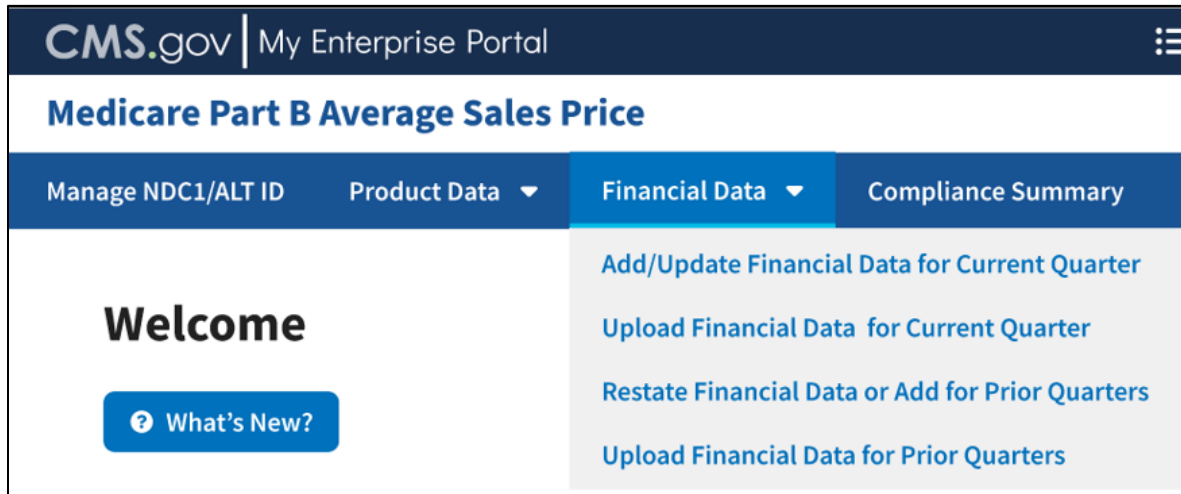


Figure 77: Financial Data - Main Drop-down

The **Upload Financial Data for Prior Quarters** page opens, listing the financial quarter and year for the upcoming reporting period. Refer to *Figure 78*.

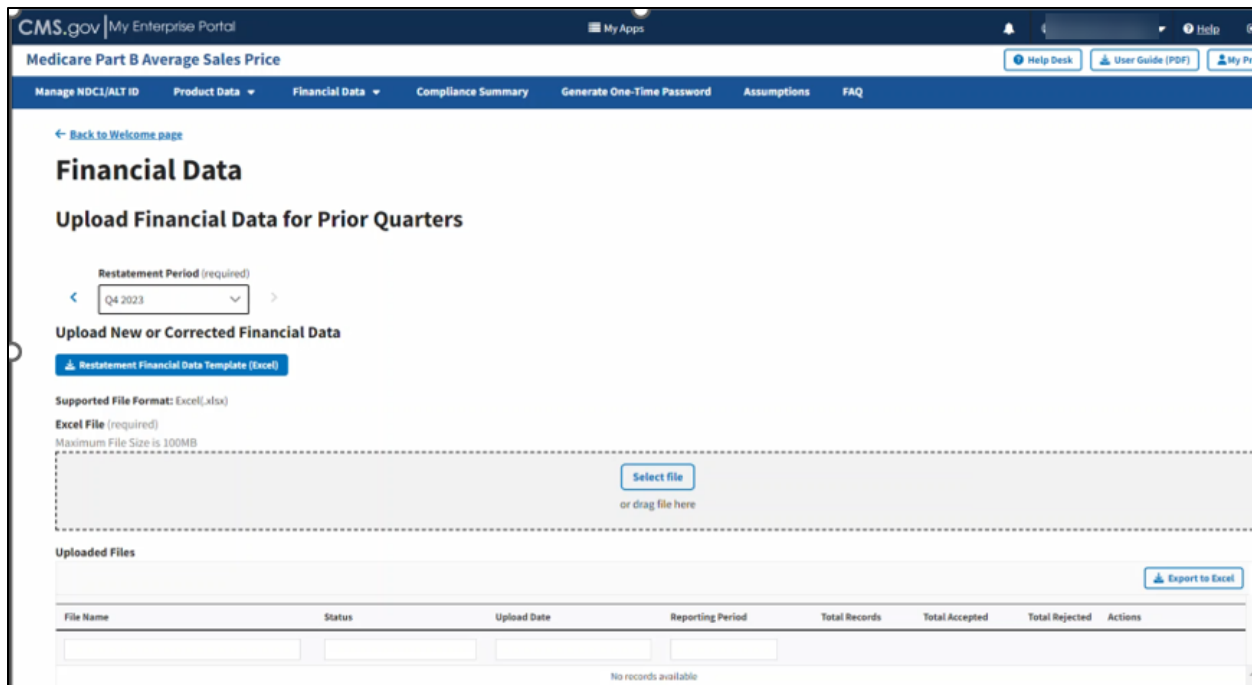


Figure 78: Upload Financial Data for Prior Quarters Restate Financial Data

Note: Under **Upload New or Corrected Financial Data**, there is a **Restatement Financial Data Template (Excel)** available for download. Click the button to download a desktop copy.

- Upon preparing your **.xlsx file (required)** and verifying your information for accuracy, click **Select File** to browse your desktop and upload the file to the Module. You may also drag the file into the **Select File** box. Refer to *Figure 79*.

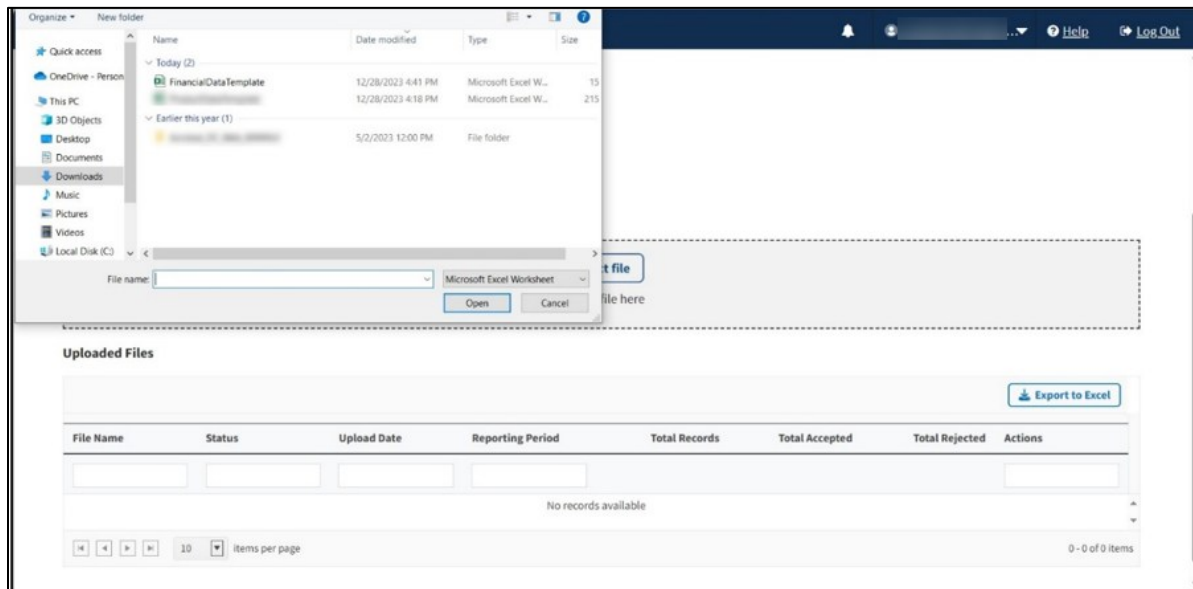


Figure 79: Upload Financial Data for Prior Quarters - Uploading Files From Desktop

A download bar displays as your file uploads. A message displays confirming you have successfully uploaded your **.xlsx** file. Refer to *Figure 80*.

Note: If the Module cannot process your file, an error message displays, and a **New Report** generates under **Uploaded Files**.

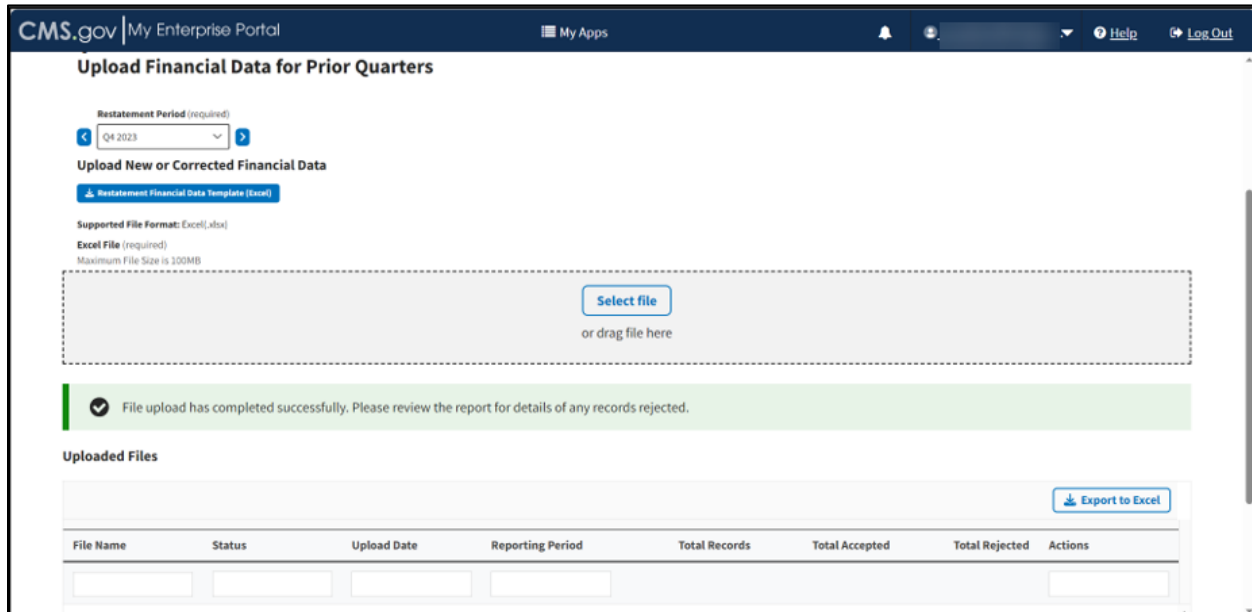
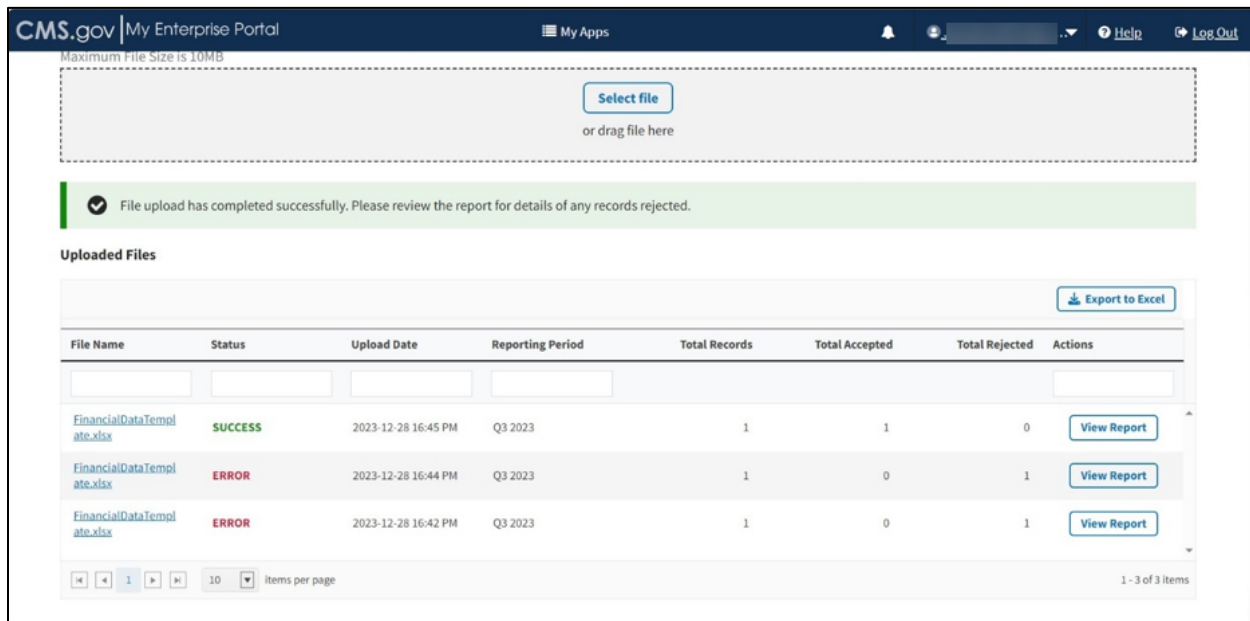


Figure 80: Upload Financial Data for Prior Quarters - New File Successfully Uploaded

The **Uploaded Files** section displays files you uploaded recently as well as previous files still in the Module. Refer to *Figure 81*.



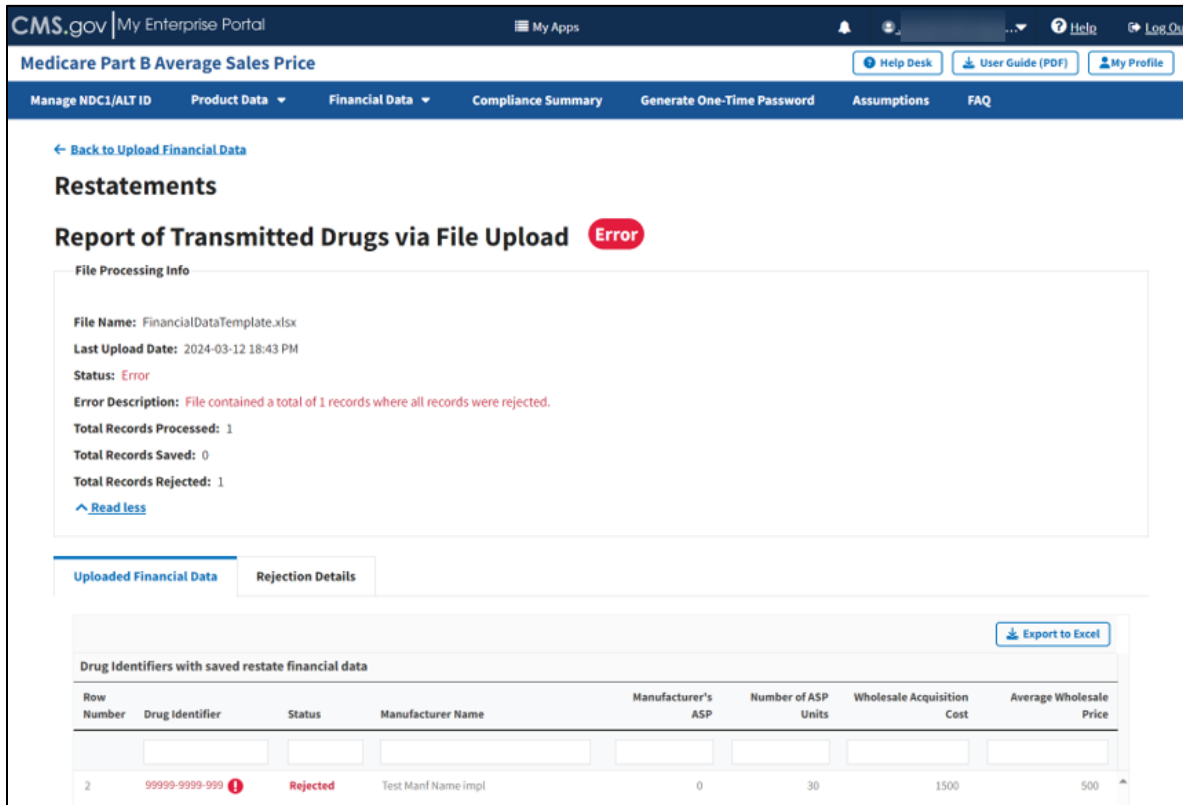
The screenshot shows the CMS.gov My Enterprise Portal interface. At the top, there are navigation links for 'My Apps', a bell icon, a help icon, and 'Log Out'. Below the header, a large dashed box area is labeled 'Maximum File Size is 10MB' and contains a 'Select file' button and a placeholder text 'or drag file here'. A green success message box at the top states: 'File upload has completed successfully. Please review the report for details of any records rejected.' Below this, a section titled 'Uploaded Files' lists three entries in a table format. The table columns are: File Name, Status, Upload Date, Reporting Period, Total Records, Total Accepted, Total Rejected, and Actions. The first file, 'FinancialDataTemplate.xlsx', has a 'Status' of 'SUCCESS', was uploaded on '2023-12-28 16:45 PM', and is for 'Q3 2023'. It has 1 total record, 1 accepted, and 0 rejected. The 'Actions' column contains a 'View Report' button. The next two files, 'FinancialDataTemplate.xlsx' and 'FinancialDataTemplate.xlsx', both have a 'Status' of 'ERROR', were uploaded on '2023-12-28 16:44 PM' and '2023-12-28 16:42 PM' respectively, and are for 'Q3 2023'. They both have 1 total record, 0 accepted, and 1 rejected. The 'Actions' column for these files also contains a 'View Report' button. At the bottom of the table, there are navigation buttons for page navigation and a dropdown for 'items per page'.

Figure 81: Upload Financial Data for Prior Quarters - Uploaded Files

Each uploaded file displays the **File Name**, **Status**, **Upload Date**, **Reporting Period**, **Total Records**, **Total Accepted**, **Total Rejected**, and **Actions** categories submitted to the Module.

3. Click **View Report** under **Actions** in the **Uploaded Files** section to view the full report for a submitted file.

The report opens on the next page. Refer to *Figure 82*.



The screenshot shows the CMS.gov My Enterprise Portal interface. The top navigation bar includes links for "My Apps", "Help", "User Guide (PDF)", and "Log Out". Below the navigation, a sub-menu for "Medicare Part B Average Sales Price" is open, showing options like "Manage NDC1/ALT ID", "Product Data", "Financial Data", "Compliance Summary", "Generate One-Time Password", "Assumptions", and "FAQ". A "Back to Upload Financial Data" link is visible. The main content area is titled "Restatements" and "Report of Transmitted Drugs via File Upload" with an "Error" status. A "File Processing Info" box displays the following data:

File Name:	FinancialDataTemplate.xlsx
Last Upload Date:	2024-03-12 18:43 PM
Status:	Error
Error Description:	File contained a total of 1 records where all records were rejected.
Total Records Processed:	1
Total Records Saved:	0
Total Records Rejected:	1

Below this is a "Read less" link. At the bottom of the "File Processing Info" box are two tabs: "Uploaded Financial Data" (selected) and "Rejection Details". The "Rejection Details" tab shows a table titled "Drug Identifiers with saved restate financial data". The table has the following columns:

Row Number	Drug Identifier	Status	Manufacturer Name	Manufacturer's ASP	Number of ASP Units	Wholesale Acquisition Cost	Average Wholesale Price
2	99999-9999-9999	Rejected	Test Manf Name impl	0	30	1500	500

An "Export to Excel" button is located at the top right of the "Rejection Details" table. The entire screenshot is framed by a light gray border.

Figure 82: Upload Financial Data for Prior Quarters - Report of Transmitted Drugs

The report lists all drug identifiers with saved restate financial data in the ASP system. The Module organizes the full list by row number and includes each drug identifier, status, and other previously submitted information from the **Add Product Data** sections.

Note: The Module highlights errors in red. Hover over the red text to display information about the specific error.

4. Click the **Rejection Details** tab.

A listing of drug identifiers with rejected restate financial data displays. Refer to *Figure 83*.

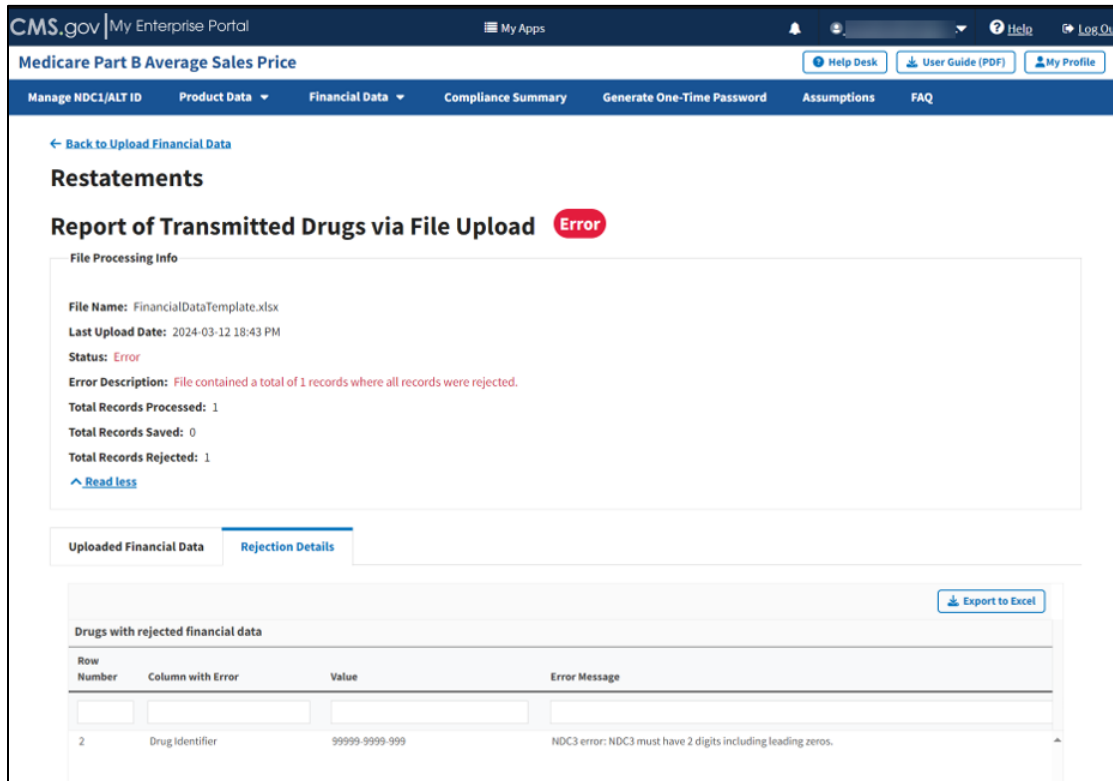


Figure 83: Upload Financial Data for Prior Quarters - Reported Rejection Details

The Module lists all errors found in submitted data by **Row Number**, **Column with Error**, **Value** and **Error Message** under **Drugs with rejected financial data**.

5. Return to the **Add/Update Financial Data** section of the Module to request any changes to your product data.
6. Contact your Certifier to recertify the corrected data you submitted to the Module.

3.5 Compliance Summary

The features in the **Compliance Summary** section allow drug manufacturers to determine if their products meet the current submission reporting requirements.

The **Compliance Summary** consists of the following sections:

- **Missing:** Displays drug products that are missing financial data for the selected reporting period.
- **Pending:** Displays drug products that are both pending certification and pending restatement certification, combined under one tab.
- **Certified:** Displays previously certified drug products for the selected reporting period.

Note: Financial data will be suppressed for prior quarters.

- **New:** Displays drug products with a first marketing date in the same reporting period.
- **Off Cycle:** Displays drug products added on or after the first day of the submission window of the current quarter.

- **Expired:** Displays drug products that have an expired date of final lot sold. A drug product that expired in an earlier quarter will continue to show in subsequent quarters.

Follow these steps to navigate the **Compliance Summary** section:

1. **From the Medicare Part B Average Sales Price homepage, click the **Compliance Summary** tab.**

The **Compliance Summary** page opens. The page displays the status for each submitted drug product regarding the drug manufacturer's compliance for the selected reporting period. The page automatically defaults to the **Missing** tab. Refer to *Figure 84*.

Note: *Figure 84* shows an alert message under **Reporting Period** stating that there are drug products in need of attention.

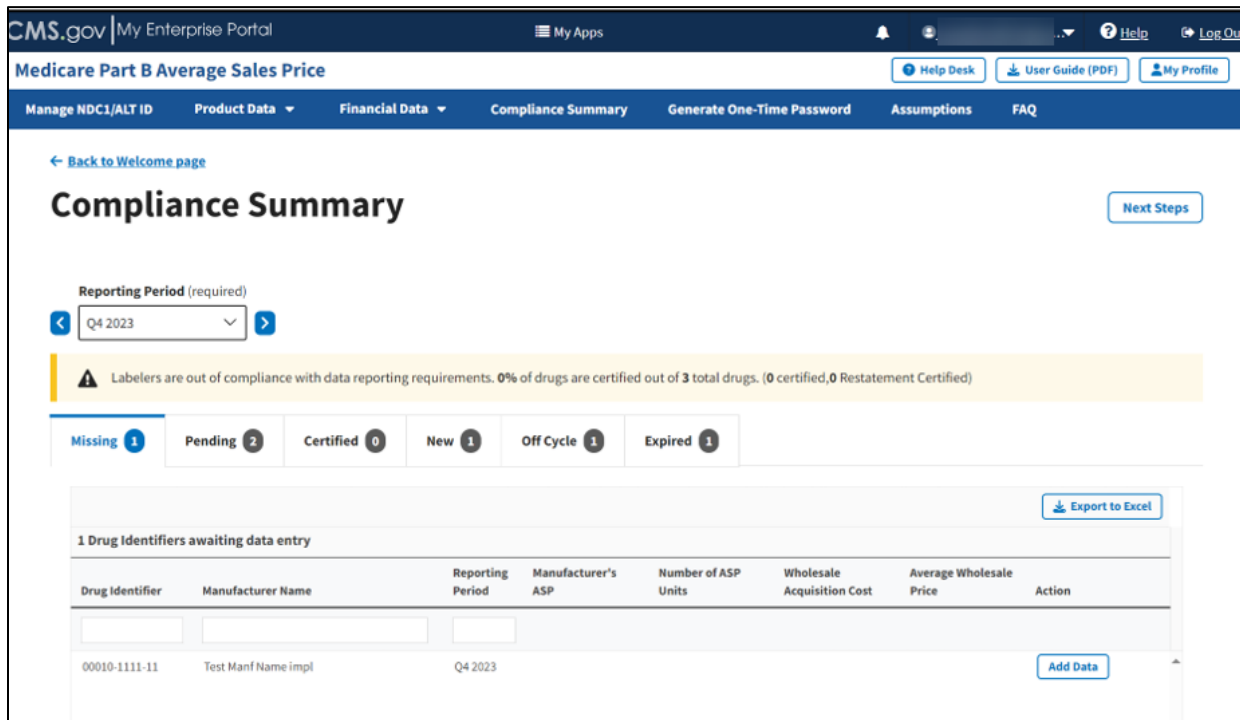

 The screenshot shows the CMS Compliance Summary page. At the top, there is a reporting period dropdown set to "Q4 2023" with arrows for navigation. A yellow alert box states: "Labelers are out of compliance with data reporting requirements. 0% of drugs are certified out of 3 total drugs. (0 certified, 0 Restatement Certified)". Below this, a navigation bar includes tabs for Missing (1), Pending (2), Certified (0), New (1), Off Cycle (1), and Expired (1). A "Next Steps" button is in the top right. The main content area shows a table for "1 Drug Identifiers awaiting data entry". The columns are: Drug Identifier, Manufacturer Name, Reporting Period, Manufacturer's ASP, Number of ASP Units, Wholesale Acquisition Cost, Average Wholesale Price, and Action. One row is visible: "00010-1111-11" and "Test Manf Name impl" under "Manufacturer Name", with "Q4 2023" under "Reporting Period". A "Export to Excel" button is in the top right of the table area.

Figure 84: Compliance Summary

Note: Click the **Reporting Period (required)** tab in the top left to scroll through previous quarters. Click the blue arrows to navigate to a previous quarter starting with the most recent or next quarter.

3.5.1 Missing

Follow these steps to add data in the **Missing** tab of the **Compliance Summary**:

1. Under **Drug Identifiers waiting for data entry**, review and identify the missing fields or incorrect financial information to address; confirm the accuracy of all the necessary financial information listed on the page.

The Module organizes the full list by **Drug Identifier** and **Manufacturer Name**, and includes **Reporting Period**, **Manufacturer's ASP**, **Number of ASP Units**, **Wholesale Acquisition Cost**, and **Average Wholesale Price** fields.

Note: Click the **Export to Excel** button to download all products under the **Missing** tab.

2. Click the **Add Data** tab next to the appropriate drug product.

An **Add Financial Data** window opens. Refer to *Figure 85*.

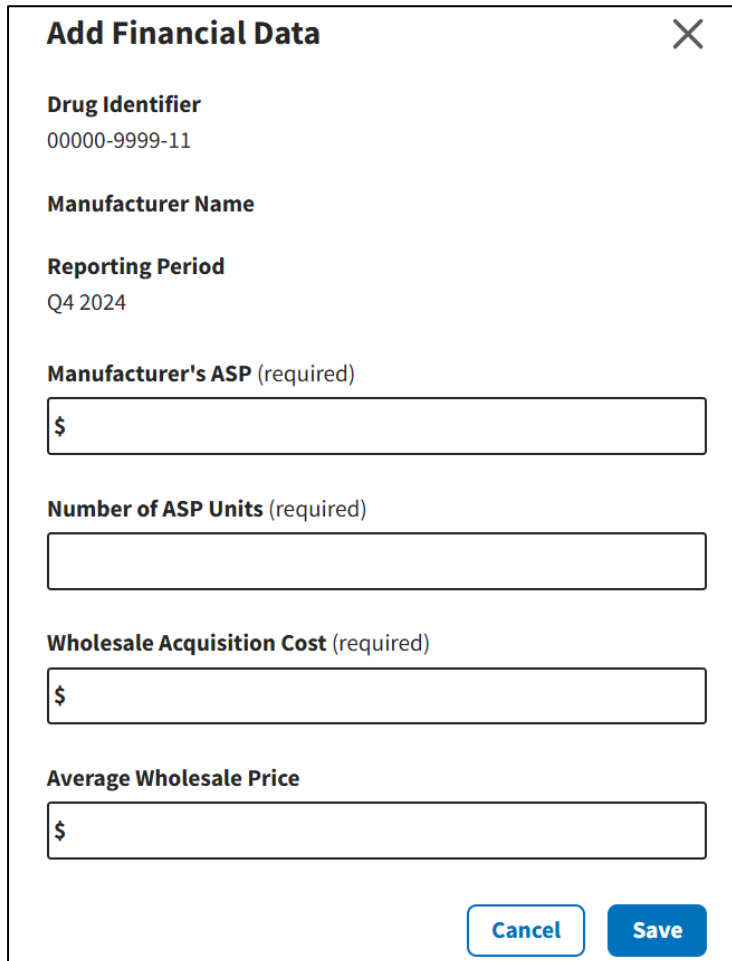

 A screenshot of the 'Add Financial Data' window. The window has a light gray background and a white content area. At the top left is the title 'Add Financial Data' and a close button 'X' at the top right. Below the title are five input fields with labels: 'Drug Identifier' (value: 00000-9999-11), 'Manufacturer Name' (empty), 'Reporting Period' (value: Q4 2024), 'Manufacturer's ASP (required)' (empty), 'Number of ASP Units (required)' (empty), 'Wholesale Acquisition Cost (required)' (empty), and 'Average Wholesale Price' (empty). Each input field is a horizontal text box with a '\$' symbol at the beginning. At the bottom right are two buttons: 'Cancel' in a light blue box and 'Save' in a dark blue box.

Figure 85: Compliance Summary - Add Data Screen

3. Type the requested information in the empty **Manufacturer's ASP (required)**, **Number of ASP Units (required)**, **Wholesale Acquisition Cost (required)**, and **Average Wholesale Price (required)** fields.
4. Click **Save** to submit your information to the Module.

A message displays confirming you have successfully added your data, and that your product is now pending certification. Refer to *Figure 86*.

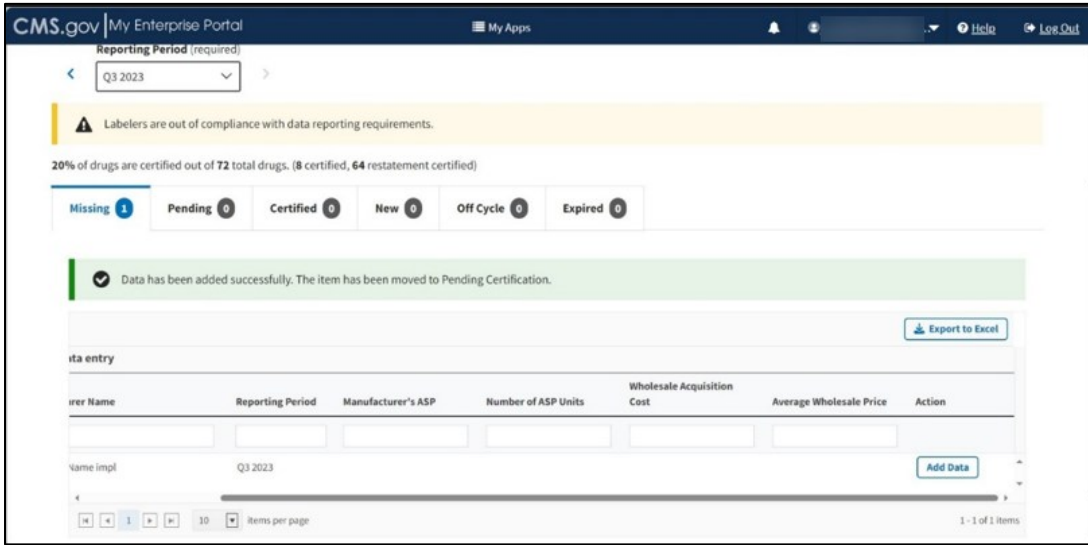

 A screenshot of the CMS.gov My Enterprise Portal. The top navigation bar includes 'My Apps', 'Help', and 'Log Out'. The reporting period is set to 'Q3 2023'. A yellow warning bar states: 'Labelers are out of compliance with data reporting requirements. 20% of drugs are certified out of 72 total drugs. (8 certified, 64 restatement certified)'. Below this, a success message says: 'Data has been added successfully. The item has been moved to Pending Certification.' A table titled 'Data entry' shows columns for 'Manufacturer Name', 'Reporting Period', 'Manufacturer's ASP', 'Number of ASP Units', 'Wholesale Acquisition Cost', 'Average Wholesale Price', and 'Action'. One row is visible: 'Name impl' under Manufacturer Name, 'Q3 2023' under Reporting Period, and 'Add Data' under Action. The bottom of the table shows pagination: '1-1 of 1 items'.

Figure 86: Compliance Summary - Successfully Saved

3.5.2 Pending

Follow these steps to review your data in the **Pending** tab of the **Compliance Summary**:

1. From the default **Compliance Summary** page, click the **Pending** tab.

The **Pending** tab displays. Refer to *Figure 87*.

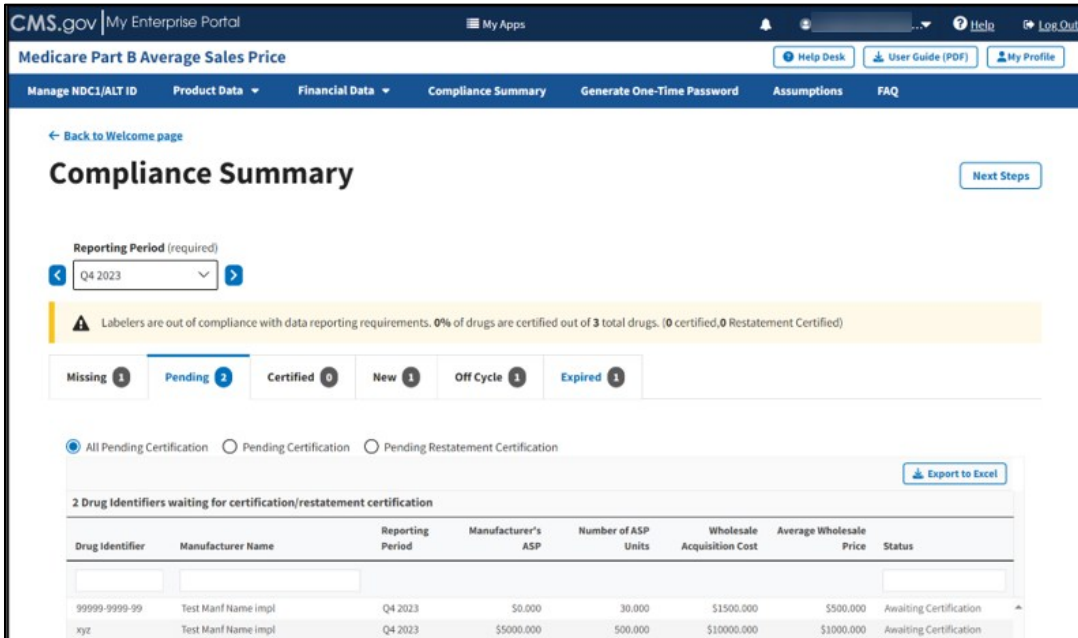

 A screenshot of the CMS.gov My Enterprise Portal. The top navigation bar includes 'My Apps', 'Help Desk', 'User Guide (PDF)', 'My Profile', and 'Log Out'. The reporting period is set to 'Q4 2023'. A yellow warning bar states: 'Labelers are out of compliance with data reporting requirements. 0% of drugs are certified out of 3 total drugs. (0 certified, 0 Restatement Certified)'. Below this, a success message says: 'All Pending Certification' is selected. A table titled '2 Drug Identifiers waiting for certification/restatement certification' shows columns for 'Drug Identifier', 'Manufacturer Name', 'Reporting Period', 'Manufacturer's ASP', 'Number of ASP Units', 'Wholesale Acquisition Cost', 'Average Wholesale Price', and 'Status'. Two rows are visible: '99999-9999-99' and 'xyz', both under 'Test Manf Name impl', with 'Q4 2023' under Reporting Period and '\$0.000' under Wholesale Acquisition Cost. The bottom of the table shows pagination: '1-2 of 2 items'.

Figure 87: Compliance Summary - All Pending Certification

The Module automatically selects the **All Pending Certification** radio button, and the page displays the drug identifiers waiting for certification/restatement certification.

Note: Click the **Export to Excel** button to download all products under the **Pending** tab.

- Under **Drug Identifiers Waiting for Certification/Restatement Certification**, review your information in the appropriate boxes previously submitted in *Section 3.2 - Product Data* and *Section 3.3 - Financial Data*.

The Module organizes the full list by **Drug Identifier** and **Manufacturer Name**, and includes **Reporting Period**, **Manufacturer's ASP**, **Number of ASP Units**, **Wholesale Acquisition Cost**, **Average Wholesale Price**, and **Status** fields.

- Click the **Pending Certification** radio button to filter only for drugs pending certification. Refer to *Figure 88*.

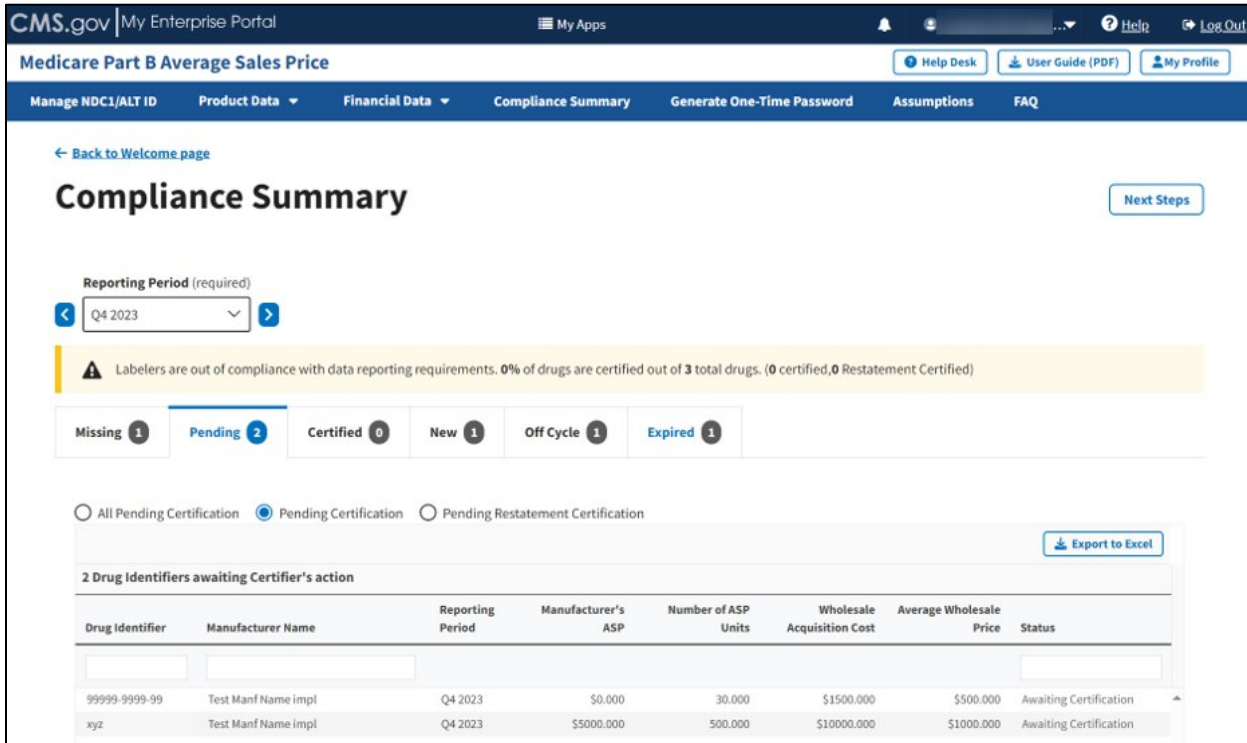

 A screenshot of the CMS ASP Module Enterprise Portal. The top navigation bar includes "My Enterprise Portal", "My Apps", "Help Desk", "User Guide (PDF)", "My Profile", and "Log Out". Below the navigation is a sub-menu for "Medicare Part B Average Sales Price" with links for "Manage NDC1/ALT ID", "Product Data", "Financial Data", "Compliance Summary" (which is the active tab), "Generate One-Time Password", "Assumptions", and "FAQ". A breadcrumb trail shows "← Back to Welcome page". The main content area is titled "Compliance Summary" with a "Next Steps" button. A reporting period dropdown is set to "Q4 2023". A yellow warning box states: "Labelers are out of compliance with data reporting requirements. 0% of drugs are certified out of 3 total drugs. (0 certified, 0 Restatement Certified)". Below this are five status buttons: "Missing 1", "Pending 2" (highlighted in blue), "Certified 0", "New 1", and "Off Cycle 1". A radio button group below shows "All Pending Certification" (unselected), "Pending Certification" (selected with a blue outline), and "Pending Restatement Certification" (unselected). An "Export to Excel" button is located in the top right of the data table. The table lists "2 Drug Identifiers awaiting Certifier's action" with columns: Drug Identifier, Manufacturer Name, Reporting Period, Manufacturer's ASP, Number of ASP Units, Wholesale Acquisition Cost, Average Wholesale Price, and Status. Two rows are shown: "99999-9999-99" and "xyz", both with "Test Manf Name impl" as the manufacturer name, "Q4 2023" as the reporting period, "\$0.000" as the manufacturer's ASP, "30.000" and "\$1500.000" as the number of ASP units and wholesale acquisition cost, "\$500.000" as the average wholesale price, and "Awaiting Certification" as the status.

Figure 88: Compliance Summary - Pending Certification

Note: Click the **Export to Excel** box to download all products under the **Pending** tab.

- Review the information previously submitted in *Section 3.2 - Product Data*.

The Module organizes the full list by **Drug Identifier** and **Manufacturer Name**, and includes **Reporting Period**, **Manufacturer's ASP**, **Number of ASP Units**, **Wholesale Acquisition Cost**, **Average Wholesale Price**, and **Status** fields.

- Click the **Pending Restatement Certification** radio button to filter only for drugs that are pending restatement certification. Refer to *Figure 89*.

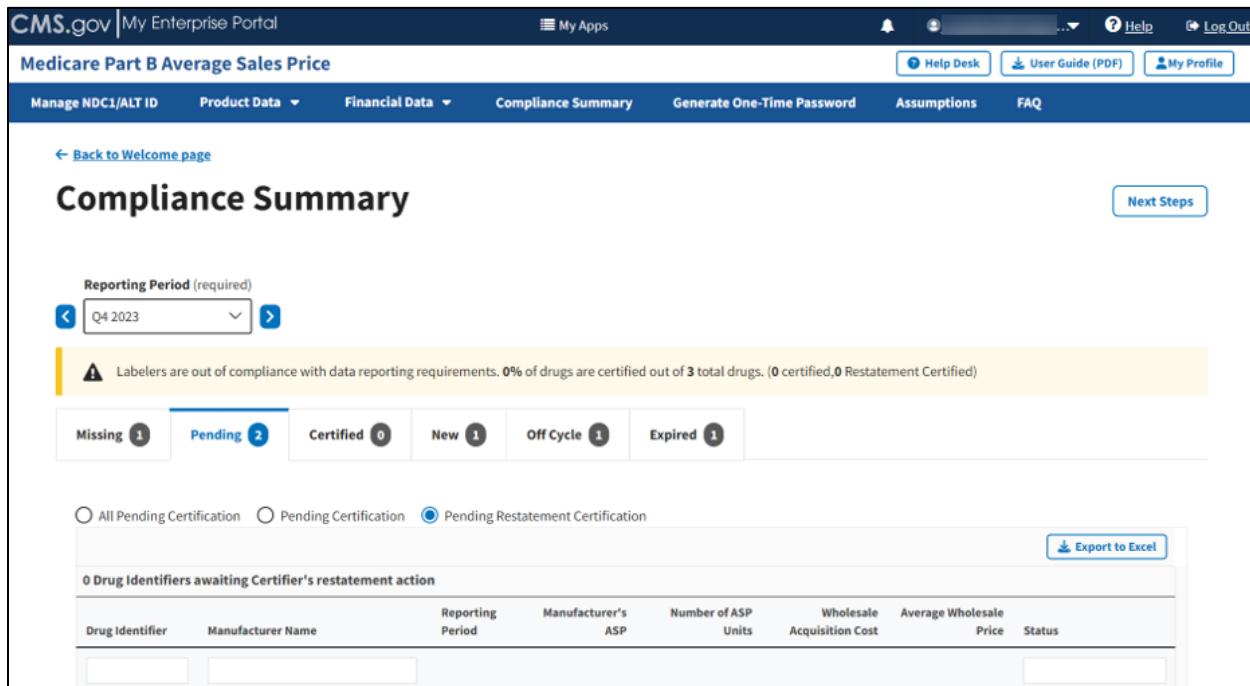


Figure 89: Compliance Summary - Pending Restatement Certification

Note: Click the **Export to Excel** button to download all products under the **Pending** tab.

6. Review all your information in the appropriate boxes previously submitted in *Section 3.2 - Product Data* and *Section 3.3 - Financial Data*.

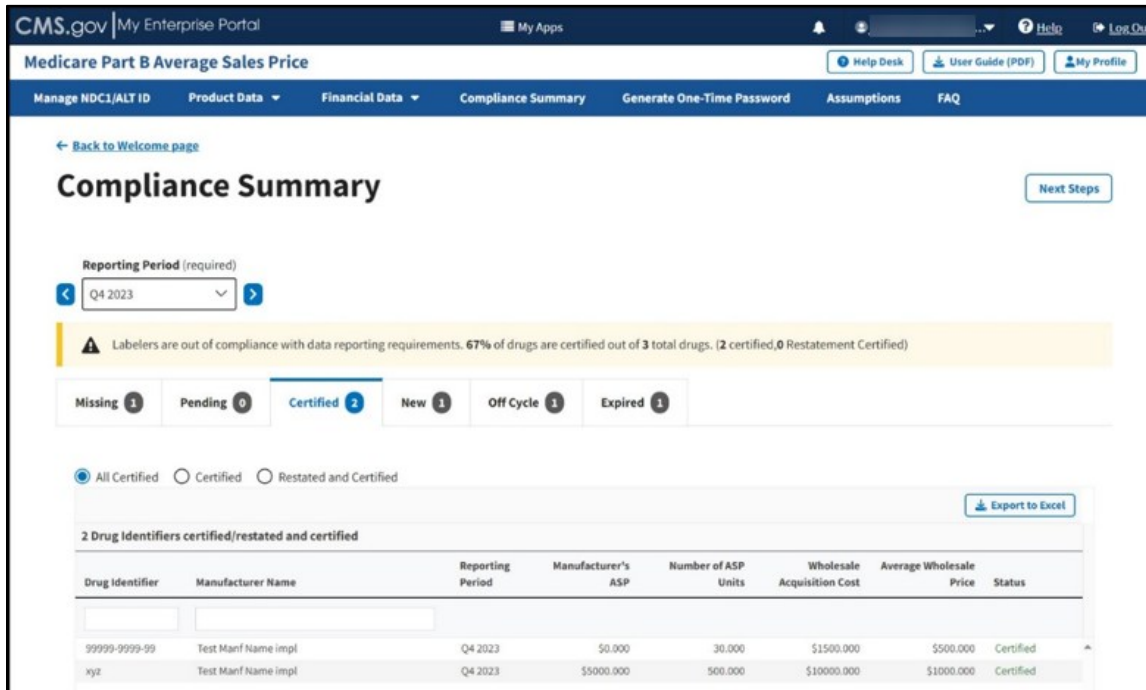
The Module organizes the full list by **Drug Identifier** and **Manufacturer Name**, and includes **Reporting Period**, **Manufacturer's ASP**, **Number of ASP Units**, **Wholesale Acquisition Cost**, and **Average Wholesale Price**, and **Status** fields.

3.5.3 Certified

Follow these steps to review your data in the **Certified** tab of the **Compliance Summary**:

1. From the default **Compliance Summary** page, click the **Certified** tab.

The **Certified** page displays. The Module automatically selects the **All Certified** radio button. Refer to *Figure 90*.



Reporting Period (required)
Q4 2023

⚠ Labelers are out of compliance with data reporting requirements. 67% of drugs are certified out of 3 total drugs. (2 certified, 0 Restatement Certified)

All Certified Certified Restated and Certified

2 Drug Identifiers certified/restated and certified

Drug Identifier	Manufacturer Name	Reporting Period	Manufacturer's ASP	Number of ASP Units	Wholesale Acquisition Cost	Average Wholesale Price	Status
99999-9999-99	Test Manf Name impl	Q4 2023	\$0.000	30.000	\$1500.000	\$500.000	Certified
xyz	Test Manf Name impl	Q4 2023	\$5000.000	500.000	\$10000.000	\$1000.000	Certified

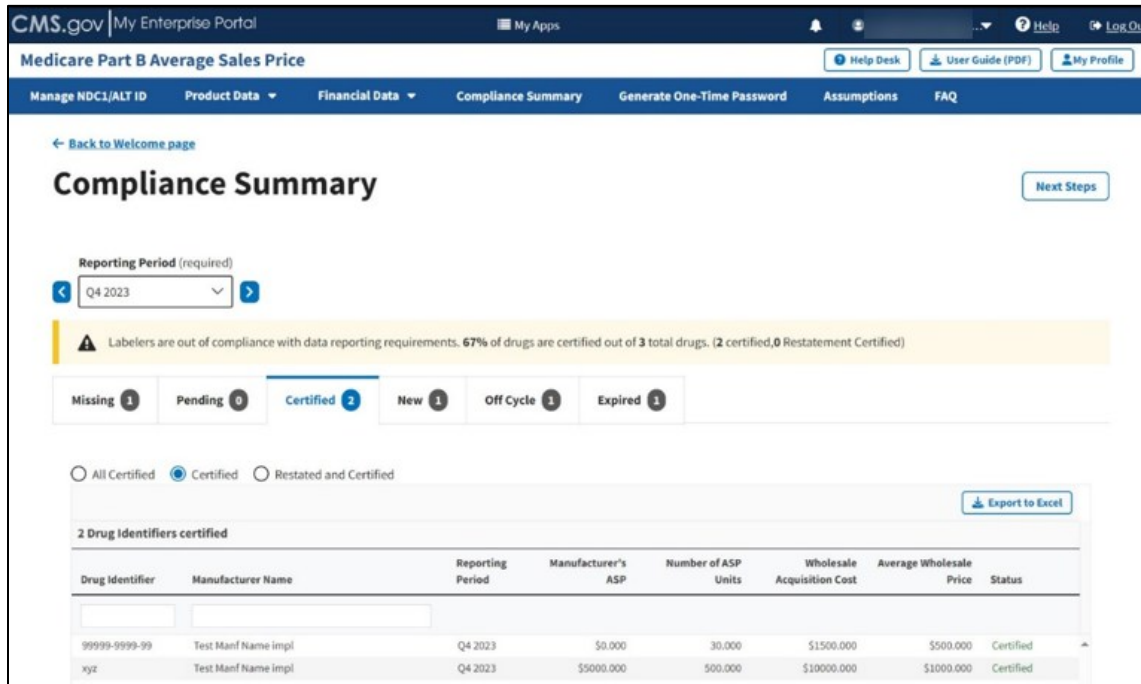
Figure 90: Compliance Summary - All Certified

Note: Click the **Export to Excel** button to download all products under the **Certified** tab.

2. Review all your information in the appropriate boxes previously submitted in *Section 3.2 - Product Data* and *Section 3.3 - Financial Data*.

The Module organizes the full list by **Drug Identifier** and **Manufacturer Name**, and includes **Reporting Period**, **Manufacturer's ASP**, **Number of ASP Units**, **Wholesale Acquisition Cost**, and **Average Wholesale Price**, and **Status** fields.

3. Click the **Certified** radio button to filter only for certified drugs. Refer to *Figure 91*.



The screenshot shows the CMS ASP Module Submitter User Guide. The top navigation bar includes links for 'My Apps', 'Help Desk', 'User Guide (PDF)', 'My Profile', 'Log Out', and 'Help'. The main menu has options like 'Manage NDC1/ALT ID', 'Product Data', 'Financial Data', 'Compliance Summary', 'Generate One-Time Password', 'Assumptions', and 'FAQ'. A 'Back to Welcome page' link is also present. The 'Compliance Summary' section is active, showing a reporting period of 'Q4 2023'. A warning message states: 'Labelers are out of compliance with data reporting requirements. 67% of drugs are certified out of 3 total drugs. (2 certified, 0 Restatement Certified)'. Below this, a navigation bar has tabs for 'Missing' (1), 'Pending' (0), 'Certified' (2), 'New' (1), 'Off Cycle' (1), and 'Expired' (1). The 'Certified' tab is selected. A radio button group allows filtering by 'All Certified', 'Certified' (which is selected), and 'Restated and Certified'. An 'Export to Excel' button is located at the top right of the data table. The table itself has columns for 'Drug Identifier', 'Manufacturer Name', 'Reporting Period', 'Manufacturer's ASP', 'Number of ASP Units', 'Wholesale Acquisition Cost', 'Average Wholesale Price', and 'Status'. It lists two entries: '99999-9999-99' and 'xyz', both from 'Test Manf Name impl' with a reporting period of 'Q4 2023'. The 'Status' column shows 'Certified' for both entries.

Figure 91: Compliance Summary - Certified

Note: Click the **Export to Excel** button to download all products under the **Certified** tab.

4. Review your information for accuracy.
The Module organizes the full list by **Drug Identifier** and **Manufacturer Name**, and includes **Reporting Period**, **Manufacturer's ASP**, **Number of ASP Units**, **Wholesale Acquisition Cost**, and **Average Wholesale Price**, and **Status** fields.
5. Click the **Restated and Certified** radio button to filter only for restated and certified drugs. Refer to *Figure 92*.

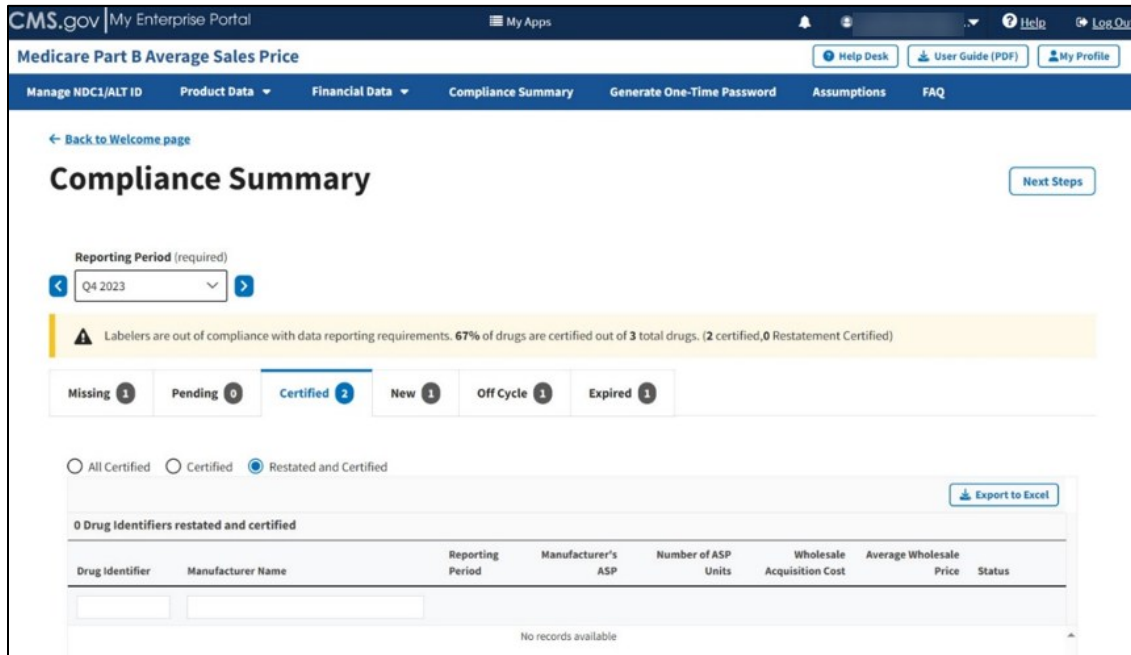

 The screenshot shows the CMS.gov My Enterprise Portal. The top navigation bar includes "My Apps", "Help", and "Log Out". Below the navigation is a menu bar with "Manage NDC/ALT ID", "Product Data", "Financial Data", "Compliance Summary", "Generate One-Time Password", "Assumptions", and "FAQ". A "Help Desk", "User Guide (PDF)", and "My Profile" link are also present. The main content area is titled "Compliance Summary". It features a reporting period selector set to "Q4 2023". A yellow warning box states: "Labelers are out of compliance with data reporting requirements. 67% of drugs are certified out of 3 total drugs. (2 certified, 0 Restatement Certified)". Below this are tabs for "Missing" (1), "Pending" (0), "Certified" (2), "New" (1), "Off Cycle" (1), and "Expired" (1). The "Certified" tab is selected. A radio button group below shows "All Certified" (unselected), "Certified" (selected), and "Restated and Certified" (selected). An "Export to Excel" button is available. A table below shows 0 drug identifiers restated and certified. The table has columns: Drug Identifier, Manufacturer Name, Reporting Period, Manufacturer's ASP, Number of ASP Units, Wholesale Acquisition Cost, Average Wholesale Price, and Status. The table is empty, showing "No records available".

Figure 92: Compliance Summary - Restated and Certified

Note: Click the **Export to Excel** box if you need to download all products under the **Certified** tab.

6. Review your information for accuracy.

The Module organizes the full list by **Drug Identifier** and **Manufacturer Name**, and includes **Reporting Period**, **Manufacturer's ASP**, **Number of ASP Units**, **Wholesale Acquisition Cost**, and **Average Wholesale Price**, and **Status** fields.

3.5.4 New

Follow these steps to review your data in the **New** tab of the **Compliance Summary**:

1. From the default **Compliance Summary** page, click the **New** tab.

The **New** page displays. Refer to *Figure 93*.

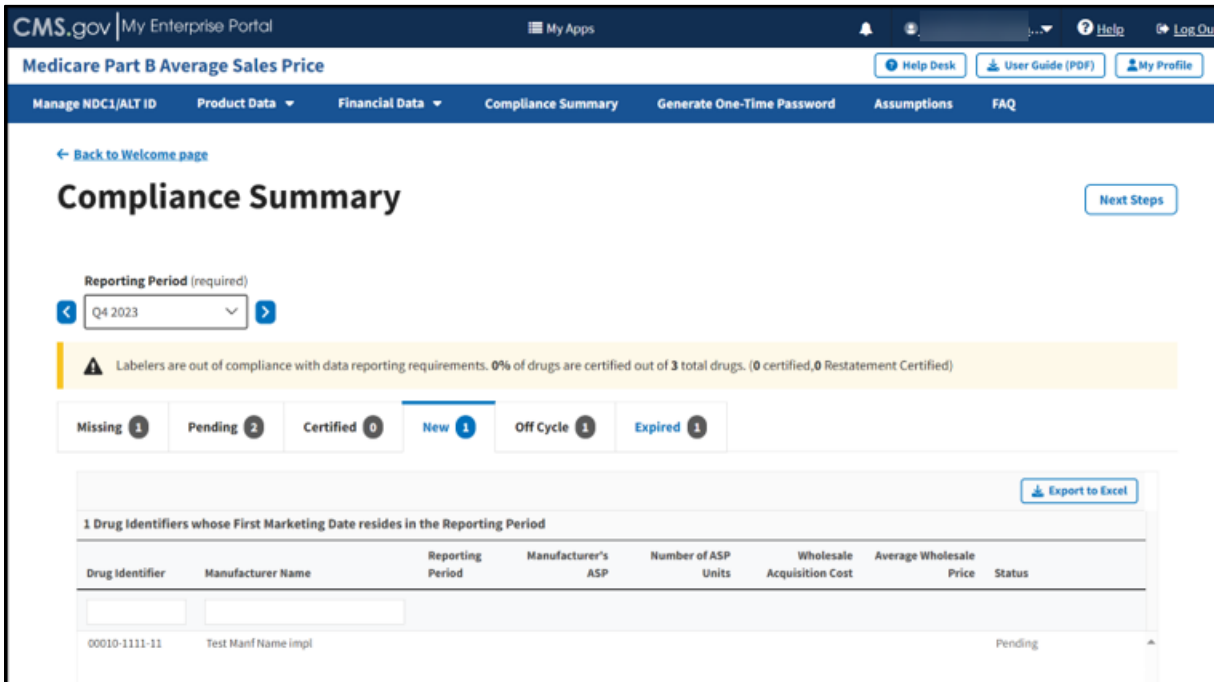

 A screenshot of the CMS ASP Module Submitter User Guide. The top navigation bar includes links for "My Apps", "Help", "User Guide (PDF)", and "My Profile". The main content area is titled "Compliance Summary" and shows a reporting period dropdown set to "Q4 2023". A warning message states: "Labelers are out of compliance with data reporting requirements. 0% of drugs are certified out of 3 total drugs. (0 certified, 0 Restatement Certified)". Below this are tabs for "Missing" (1), "Pending" (2), "Certified" (1), "New" (1), "Off Cycle" (1), and "Expired" (3). The "New" tab is selected. A table titled "1 Drug Identifiers whose First Marketing Date resides in the Reporting Period" is displayed, with columns for Drug Identifier, Manufacturer Name, Reporting Period, Manufacturer's ASP, Number of ASP Units, Wholesale Acquisition Cost, Average Wholesale Price, and Status. One row is shown: "00010-1111-11" and "Test Manf Name impl" under "Manufacturer Name", with "Pending" under "Status". An "Export to Excel" button is located at the top right of the table.

Figure 93: Compliance Summary - New

Note: Click the **Export to Excel** button to download all products under the **New** tab.

2. Review your information for accuracy.

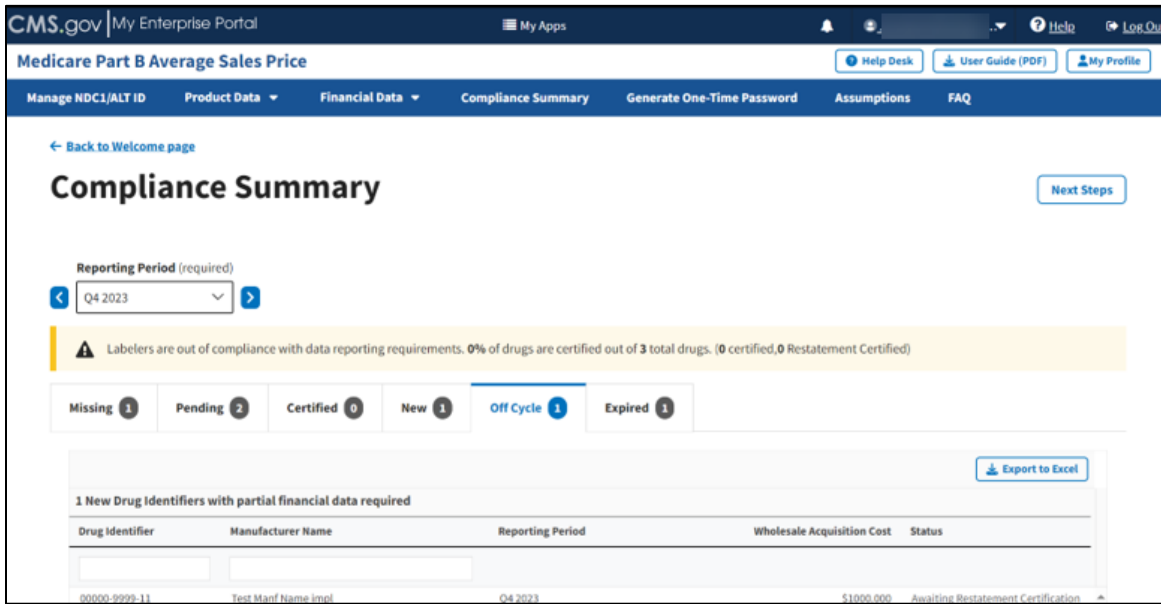
The Module organizes the full list by **Drug Identifier** and **Manufacturer Name**, and includes **Reporting Period**, **Manufacturer's ASP**, **Number of ASP Units**, **Wholesale Acquisition Cost**, and **Average Wholesale Price**, and **Status** fields.

3.5.5 Off Cycle

Follow these steps to review your data in the **Off Cycle** tab of the **Compliance Summary**:

1. From the default **Compliance Summary** page, click the **Off Cycle** tab.

The **Off Cycle** page displays. Refer to *Figure 94*.



The screenshot shows the CMS.gov My Enterprise Portal with the 'Medicare Part B Average Sales Price' module selected. The 'Compliance Summary' tab is active. A reporting period dropdown shows 'Q4 2023'. A yellow warning box states: 'Labelers are out of compliance with data reporting requirements. 0% of drugs are certified out of 3 total drugs. (0 certified, 0 Restatement Certified)'. Below are tabs for Missing (1), Pending (2), Certified (0), New (3), Off Cycle (1), and Expired (1). A table lists one product: '00000-9999-11' (Test Manf Name impl), 'Q4 2023', '\$1000.000', and 'Awaiting Restatement Certification'. An 'Export to Excel' button is at the top right of the table.

Figure 94: Compliance Summary - Off Cycle

Note: Click the **Export to Excel** button to download all products under the **Off Cycle** tab.

2. Review your information for accuracy.

The Module organizes the full list by **Drug Identifier** and **Manufacturer Name**, and includes **Reporting Period**, **Wholesale Acquisition Cost**, and **Status** fields.

3.5.6 Expired

Follow these steps to review your data in the **Expired** tab of the **Compliance Summary**:

1. From the default **Compliance Summary** page, click the **Expired** tab.
2. The **Expired** page displays. Refer to *Figure 95*.

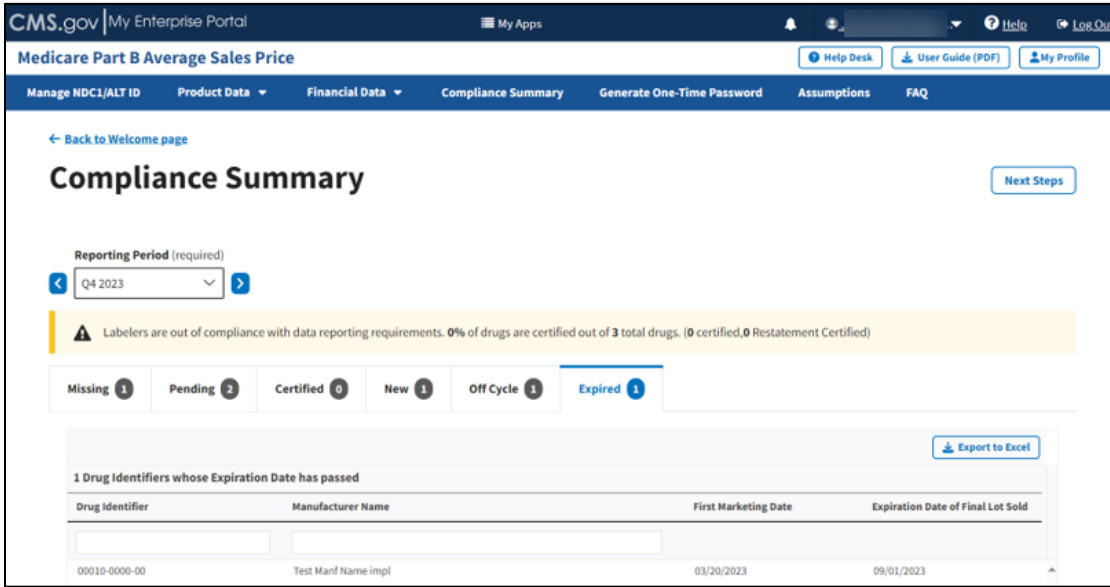

 A screenshot of the CMS.gov Compliance Summary page. The top navigation bar includes links for "My Apps", "Help", "User Guide (PDF)", "My Profile", "Help Desk", and "Logout". The main menu has options like "Manage NDC1/ALT ID", "Product Data", "Financial Data", "Compliance Summary", "Generate One-Time Password", "Assumptions", and "FAQ". A "Back to Welcome page" link is also present. The "Compliance Summary" section shows a reporting period of "Q4 2023". A yellow warning box states: "Labelers are out of compliance with data reporting requirements. 0% of drugs are certified out of 3 total drugs. (0 certified, 0 Restatement Certified)". Below this, a horizontal bar shows the count of drugs in various states: Missing (1), Pending (2), Certified (0), New (1), Off Cycle (1), and Expired (1). A table below lists "1 Drug Identifiers whose Expiration Date has passed". The table has columns for "Drug Identifier", "Manufacturer Name", "First Marketing Date", and "Expiration Date of Final Lot Sold". An example row shows "00010-0000-00", "Test Manf Name impl", "03/20/2023", and "09/01/2023". A "Next Steps" button is in the top right, and an "Export to Excel" button is in the middle right.

Figure 95: Compliance Summary - Expired

Note: Click the **Export to Excel** box if you need to download all products under the **Expired** tab.

3. Review your information for accuracy.

The Module organizes the full list by **Drug Identifier** and **Manufacturer Name** and includes **First Marketing Date** and **Expiration Date of Final Lot Sold** fields.

3.6 Generate One-Time Password

Once you successfully enter product and financial data in the ASP Module, you can generate a one-time password (OTP) for each manufacturer name. Note the following about OTPs:

- OTPs protect sensitive information and product specific drug-data from tampering or alterations by others outside of the Submitter or Certifier.
- The OTP is a one-time authentication step to link a Submitter to a Certifier within the system. This step does not need to take place during every submission. There can only be one active Certifier per manufacturer. If the Certifier changes, the Submitter must create and share a new OTP with the new Certifier.
- The Submitter and Certifier cannot be the same person within your organization.
- You can share the OTP with the Certifier. This passcode will remain the same for as long as the Certifier is the same person in your organization who uses the ASP Module.
- If the OTP expires, you can generate another OTP and provide it to the Certifier again.

Note: Refer to the Certifier User Guide for more information about the Certifier role.

Follow these steps to generate an OTP:

1. From the Medicare Part B ASP Homepage, click the **Generate One-Time Password** tab.

The **Generate One-Time Password** page opens. Refer to *Figure 96*.



Figure 96: Generate One-Time Password

2. Click the **-Select-** box under **Manufacturer Name (required)** to expand the list. Refer to *Figure 97*.

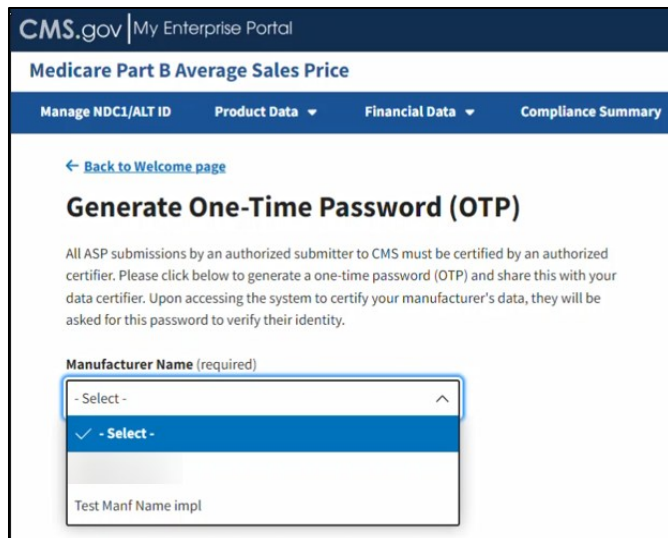
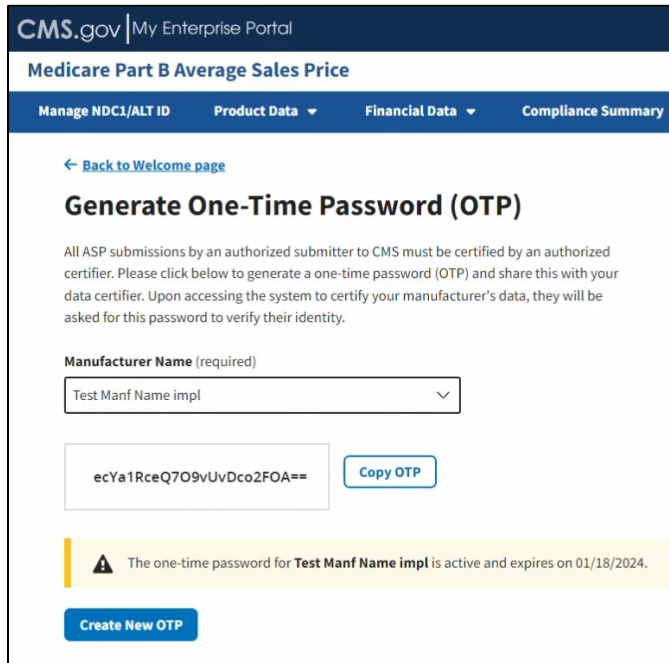


Figure 97: Generate One-Time Password - Manufacturer Name

3. Select the appropriate manufacturer name.

A new OTP displays. Refer to *Figure 98*.



Generate One-Time Password (OTP)

All ASP submissions by an authorized submitter to CMS must be certified by an authorized certifier. Please click below to generate a one-time password (OTP) and share this with your data certifier. Upon accessing the system to certify your manufacturer's data, they will be asked for this password to verify their identity.

Manufacturer Name (required)

Test Manf Name impl

ecYa1RceQ7O9vUvDco2FOA== Copy OTP

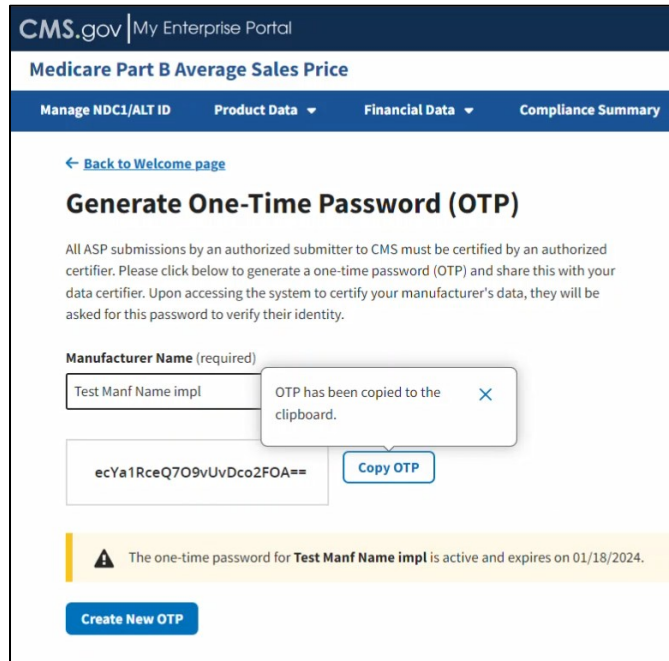
⚠ The one-time password for **Test Manf Name impl** is active and expires on 01/18/2024.

Create New OTP

Figure 98: Generate One-Time Password - Password Created

4. Click **Copy OTP** to copy your OTP.

Hover text indicates that you have successfully copied the new password. Refer to *Figure 99*.



Generate One-Time Password (OTP)

All ASP submissions by an authorized submitter to CMS must be certified by an authorized certifier. Please click below to generate a one-time password (OTP) and share this with your data certifier. Upon accessing the system to certify your manufacturer's data, they will be asked for this password to verify their identity.

Manufacturer Name (required)

Test Manf Name impl

OTP has been copied to the clipboard. X

ecYa1RceQ7O9vUvDco2FOA== Copy OTP

⚠ The one-time password for **Test Manf Name impl** is active and expires on 01/18/2024.

Create New OTP

Figure 99: Generate One-Time Password - Password Copied

5. Copy the OTP and send it to your Certifier. You must recreate the OTP if the Certifier cannot confirm the OTP on the Module, or if it expires.

Note: A message displays at the bottom of the window noting the expiration date for your new password. The Certifier must log in to the ASP Module to use that OTP before the noted expiration date.

Note: An OTP is only valid for seven days. After seven days, you must generate a new OTP.

3.7 Assumptions

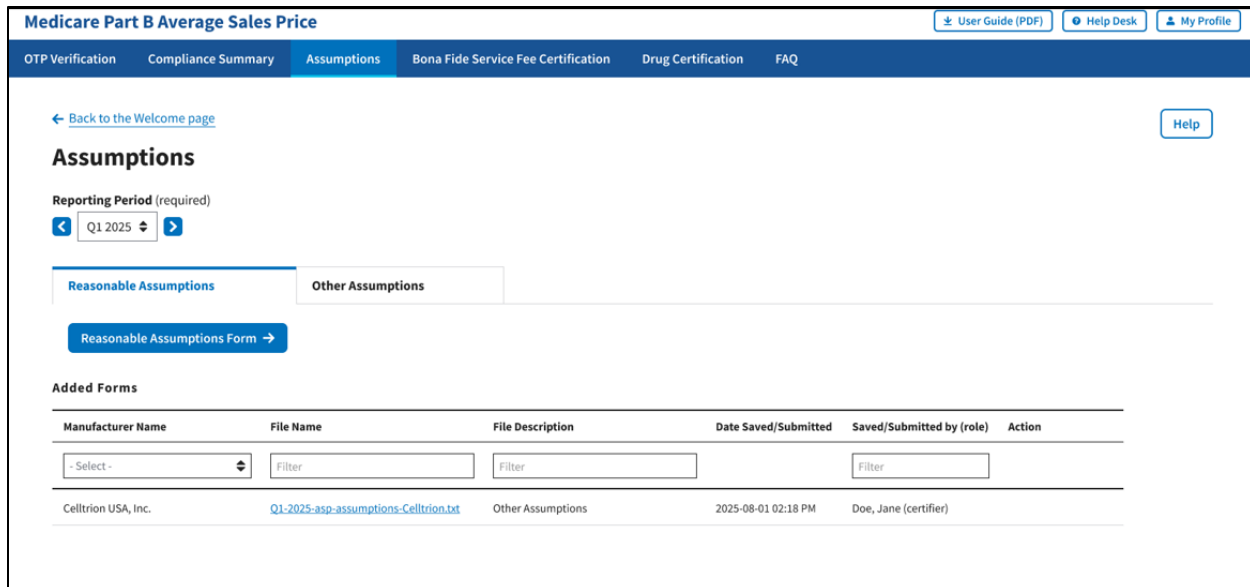
Drug manufacturers can submit comments regarding their certifications to CMS. Manufacturers may submit these comments for either the current or prior reporting periods. Each quarter, manufacturers will submit these comments for the current reporting period, or they may submit assumptions for any previous quarters they are restating and resubmitting. Submitters can enter assumptions, but certifiers must complete the assumptions form before certification.

3.7.1 Reasonable Assumptions

Follow these steps to submit certification assumptions to CMS:

1. From the **Medicare Part B Average Sales Price** homepage, click the **Assumptions** tab.

The **Assumptions** page opens, and defaults to the current quarter and year. Select the appropriate reporting period before clicking the **Reasonable Assumptions** tab. Refer to *Figure 100*.



Manufacturer Name	File Name	File Description	Date Saved/Submitted	Saved/Submitted by (role)	Action
- Select -	Q1-2025-asp-assumptions-Celltrion.txt	Other Assumptions	2025-08-01 02:18 PM	Doe, Jane (certifier)	<input type="button" value="Filter"/>

Figure 100: Assumptions

Note: Click the **Reporting Period (Required)** tab in the top left to scroll through previous quarters.

2. Click the **Reasonable Assumptions Form** button.

The **Reasonable Assumption Form** window displays. The Module automatically defaults to the **Reporting Period** selected on the **Assumptions** default page with a **Manufacturer Name (required)** drop-down menu and empty required response fields. Refer to *Figure 101*.

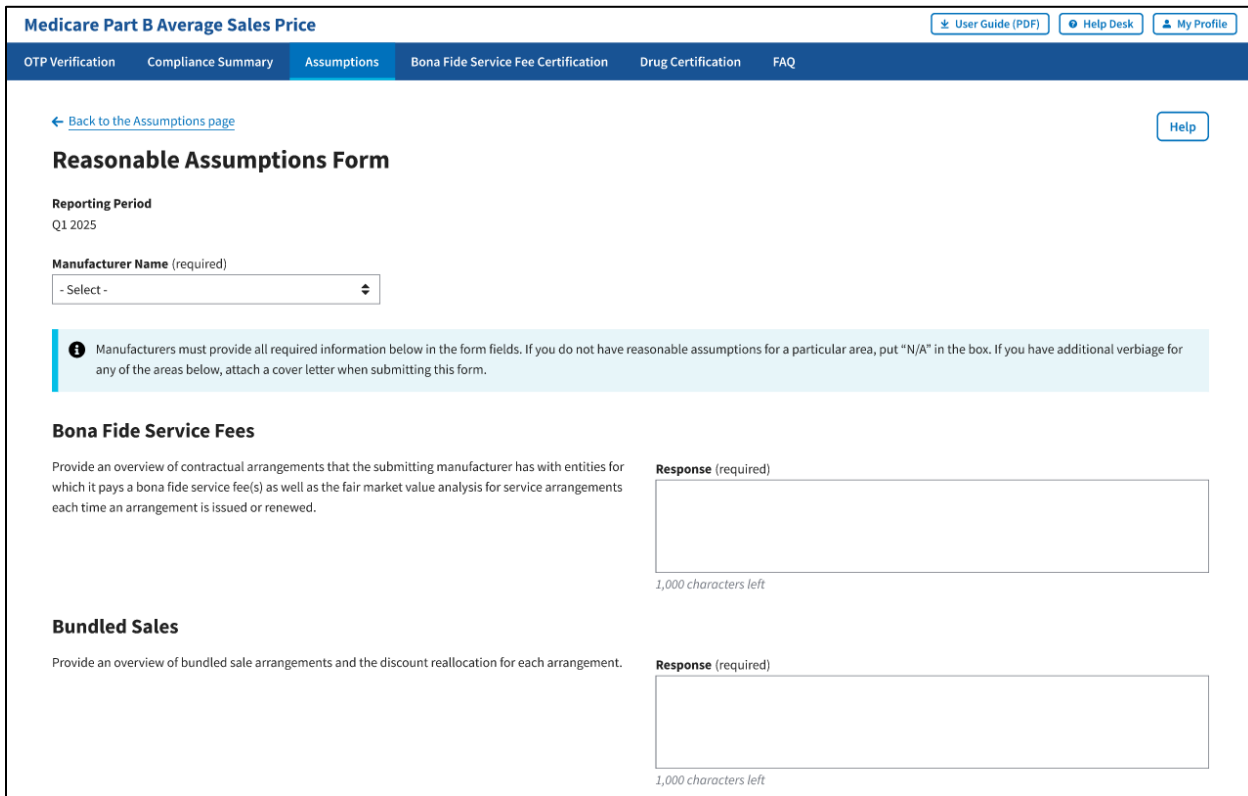

 A screenshot of the CMS ASP Module's Reasonable Assumptions Form. The top navigation bar includes links for User Guide (PDF), Help Desk, and My Profile. The main content area has a blue header 'Medicare Part B Average Sales Price' and a blue sub-header 'Reasonable Assumptions Form'. A 'Reporting Period' section shows 'Q1 2025'. A 'Manufacturer Name (required)' dropdown menu is open, showing the option '- Select -'. A note below the dropdown states: 'Manufacturers must provide all required information below in the form fields. If you do not have reasonable assumptions for a particular area, put "N/A" in the box. If you have additional verbiage for any of the areas below, attach a cover letter when submitting this form.' Below this are sections for 'Bona Fide Service Fees' and 'Bundled Sales', each with a text area labeled 'Response (required)' and a character count of '1,000 characters left'.

Figure 101: Reasonable Assumptions Form

3. From the **Manufacturer Name (required)** drop-down menu, click the **-Select-** drop-down menu to expand the list and select the manufacturer name.
4. Click **View All** to view all the required response fields. Refer to *Figure 102*.
 - Bona Fide Service Fees
 - Bundled Sales
 - Price Concessions and Discounts
 - Reporting of Products with Zero, Negative, or False Positive ASPs
 - Sales Excluded from Best Price
 - Sales to U.S. Territories
 - Time Value of Money
 - Free Goods Not Contingent on a Purchase Requirement
 - Value-Based Purchasing Agreements
 - Sales to 340B Covered Entities
 - Returned Goods
 - Billing Corrections

Sales to U.S. Territories

Confirm how the manufacturer considers sales to customers in United States territories in the calculation of ASP..

Response (required)

1,000 characters left

[View All](#)

Figure 102: View All Required Response Fields

5. Complete all the response fields. Enter **NA** if reasonable assumptions are not available for a particular field.

Note: Each required field allows for 1,000 characters of text to provide a summary of the assumption. If a response exceeds the character limit, please submit or upload the additional verbiage on the **Other Assumptions** tab. Refer to *Section 3.7.2 - Other Assumptions* for instructions.

6. Click the **Save Form** button located at the bottom of the form. Refer to *Figure 103*.

Returned Goods

Confirm how returned goods will be treated in the ASP calculation.

Response (required)

1,000 characters left

Billing Corrections

Confirm how you process transactional issues that may require a credit or rebill.

Response (required)

1,000 characters left

[Save Form](#)

Figure 103: Save Reasonable Assumptions Form

A message displays confirming you have successfully created your **Reasonable Assumptions**. The Module lists saved forms under **Added Forms**. Refer to *Figure 104*.

Medicare Part B Average Sales Price

[User Guide \(PDF\)](#) [Help Desk](#) [My Profile](#)

[OTP Verification](#) [Compliance Summary](#) [Assumptions](#) [Bona Fide Service Fee Certification](#) [Drug Certification](#) [FAQ](#)

[← Back to the Welcome page](#) [Help](#)

Assumptions

Reporting Period (required)

Q1 2025 < >

[Reasonable Assumptions](#) [Other Assumptions](#)

[Reasonable Assumptions Form →](#)

✓ Reasonable Assumptions Form has been saved successfully.

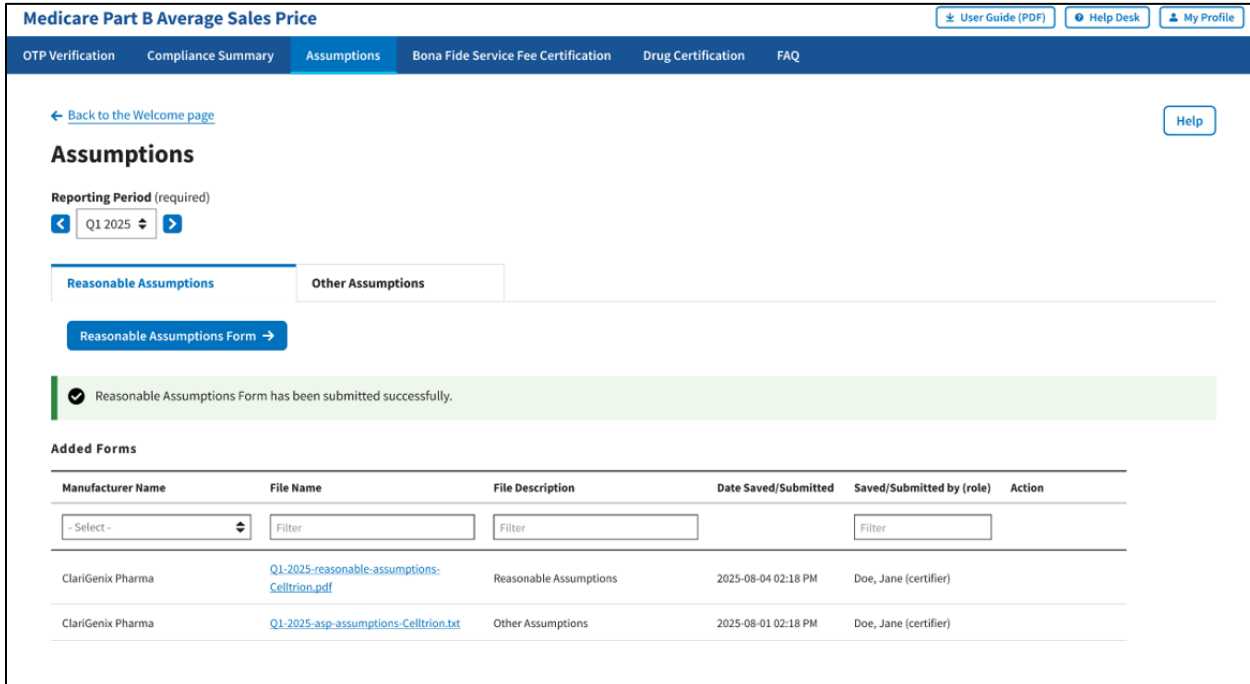
Added Forms

Manufacturer Name	File Name	File Description	Date Saved/Submitted	Saved/Submitted by (role)	Action
<input style="width: 150px; height: 25px; border: 1px solid #ccc; border-radius: 5px; padding: 2px; margin-bottom: 5px;" type="button" value="- Select -"/> Filter Filter Filter	Q1-2025-reasonable-assumptions-Celltrion.pdf	Reasonable Assumptions	2025-08-04 11:18 AM	Doe, Jane (certifier)	Edit Submit
<input style="width: 150px; height: 25px; border: 1px solid #ccc; border-radius: 5px; padding: 2px; margin-bottom: 5px;" type="button" value="ClariGenix Pharma"/> Filter Filter Filter	Q1-2025-asp-assumptions-Celltrion.txt	Other Assumptions	2025-08-01 02:18 PM	Doe, Jane (certifier)	

Figure 104: New Assumption Successfully Saved

7. To make any necessary revisions before submitting, click the **Edit** button.
8. If the submission does not require additional revisions, click the **Submit** button.

A message displays confirming you have successfully submitted your **Reasonable Assumptions**. Refer to *Figure 105*.


 A screenshot of the CMS ASP Module Submitter User Guide. The page title is "Medicare Part B Average Sales Price". The top navigation bar includes links for OTP Verification, Compliance Summary, Assumptions, Bona Fide Service Fee Certification, Drug Certification, and FAQ. On the far right of the top bar are links for "User Guide (PDF)", "Help Desk", and "My Profile". Below the top bar, there is a "Back to the Welcome page" link and a "Help" button. The main content area is titled "Assumptions". Under "Assumptions", there is a "Reporting Period (required)" dropdown set to "Q1 2025". Below the dropdown are two tabs: "Reasonable Assumptions" (which is selected and highlighted in blue) and "Other Assumptions". A blue button labeled "Reasonable Assumptions Form →" is visible. A green success message box contains the text: "Reasonable Assumptions Form has been submitted successfully." Below this message, there is a section titled "Added Forms" with a table. The table has columns for Manufacturer Name, File Name, File Description, Date Saved/Submitted, Saved/Submitted by (role), and Action. It lists two entries: one for ClariGenix Pharma with file name "Q1-2025-reasonable-assumptions-Celtrion.pdf" and another for ClariGenix Pharma with file name "Q1-2025-asp-assumptions-Celtrion.txt".

Manufacturer Name	File Name	File Description	Date Saved/Submitted	Saved/Submitted by (role)	Action
- Select -	Filter	Filter	Filter		
ClariGenix Pharma	Q1-2025-reasonable-assumptions-Celtrion.pdf	Reasonable Assumptions	2025-08-04 02:18 PM	Doe, Jane (certifier)	
ClariGenix Pharma	Q1-2025-asp-assumptions-Celtrion.txt	Other Assumptions	2025-08-01 02:18 PM	Doe, Jane (certifier)	

Figure 105: Reasonable Assumptions Successfully Submitted

3.7.2 Other Assumptions

This section provides instructions on how drug manufacturers can submit comments regarding their certifications to CMS via **Create Assumptions** or **Upload Assumptions**.

3.7.2.1 Create Assumptions

Follow these steps to create an assumption:

1. From the **Medicare Part B Average Sales Price** homepage, click the **Assumptions** tab. The Module automatically defaults to the **Reasonable Assumptions** tab. Click the **Other Assumptions** tab. Refer to *Figure 106*.

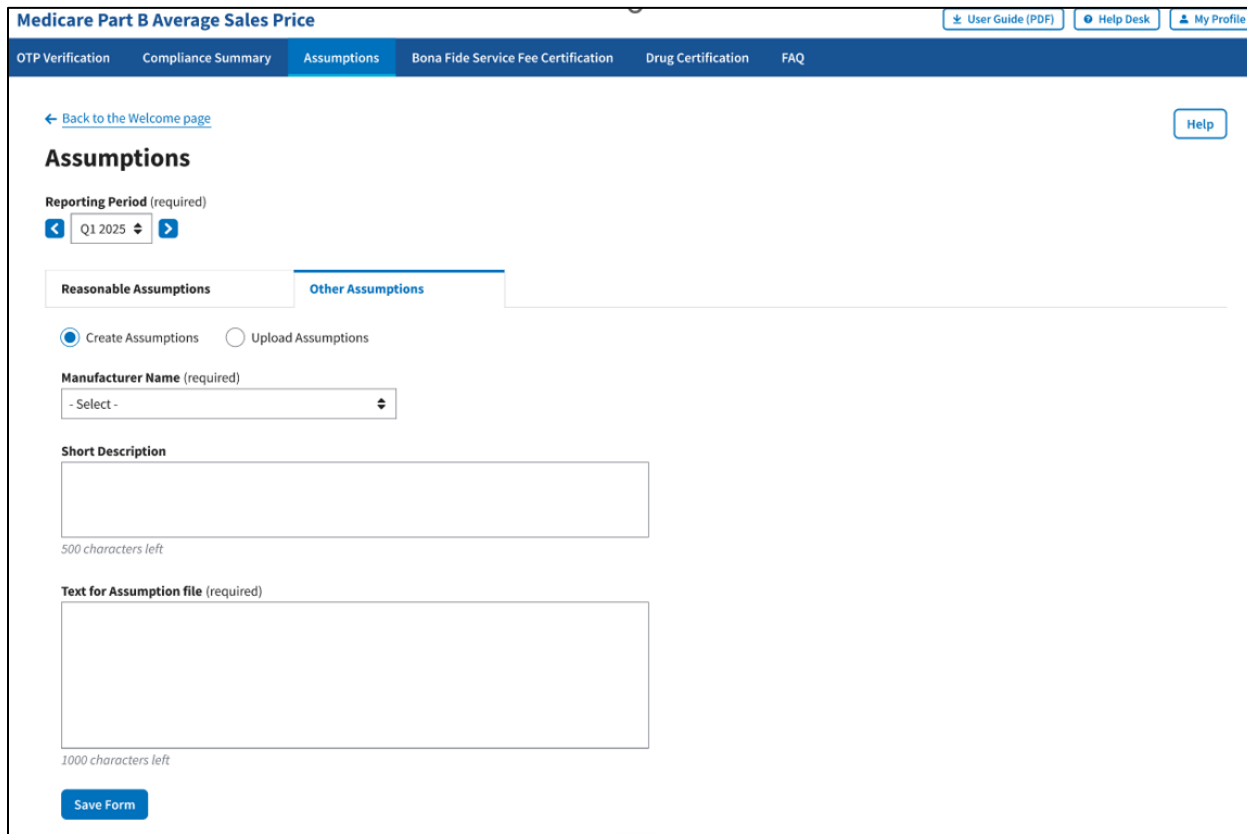

 A screenshot of a web page titled "Medicare Part B Average Sales Price". The top navigation bar includes links for "OTP Verification", "Compliance Summary", "Assumptions" (which is the active tab), "Bona Fide Service Fee Certification", "Drug Certification", and "FAQ". On the right side of the top bar are links for "User Guide (PDF)", "Help Desk", and "My Profile". Below the top bar, there is a "Back to the Welcome page" link and a "Help" button. The main content area is titled "Assumptions". It features a "Reporting Period (required)" section with a dropdown menu showing "Q1 2025". Below this are two tabs: "Reasonable Assumptions" (selected) and "Other Assumptions". Under "Other Assumptions", there are two radio buttons: "Create Assumptions" (selected) and "Upload Assumptions". A "Manufacturer Name (required)" dropdown menu is shown with the placeholder "- Select -". A "Short Description" text area is present with the note "500 characters left". A larger "Text for Assumption file (required)" text area is also present with the note "1000 characters left". At the bottom is a "Save Form" button.

Figure 106: Create Other Assumptions

Note: Click the **Reporting Period** tab in the top left to view previous quarters. Use the drop-down menu to navigate to select the appropriate quarter.

2. Click the **Other Assumptions** file button.

The Module automatically defaults to the **Create Assumptions** radio button with a **Manufacturer Name (required)** drop-down menu and empty **Short Description** and **Text for Assumption file** fields. Refer to *Figure 106*.

3. From the **Manufacturer Name (required)** drop-down menu, click the **-Select-** drop-down menu to expand the list and select the manufacturer name.
4. Complete the **Short Description** and **Text for Assumption file** fields.

Note: The **Short Description** field is optional and allows for 500 characters of text to provide a summary of the complete assumption you are submitting to CMS. The **Text for Assumption file** field is required and allows for 1,000 characters to provide as much detail as possible related to the selected period's financial submission.

5. Click the **Save Form** button.

A message displays confirming you have successfully created your **Assumption**. Refer to *Figure 107*.

Other Assumptions Form has been saved successfully.

Added Forms					
Manufacturer Name	File Name	File Description	Date Saved/Submitted	Saved/Submitted by (role)	Action
- Select -	<input type="button" value="Filter"/>	<input type="button" value="Filter"/>			<input type="button" value="Filter"/>
ClariGenix Pharma	Q1-2025-other-assumptions-ClariGenix-Pharma.txt	Other Assumptions	2025-08-04 02:18 PM	Doe, Jane (certifier)	
ClariGenix Pharma	Q1-2025-reasonable-assumptions-ClariGenix-Pharma.pdf	Reasonable Assumptions	2025-08-01 01:28 PM	Doe, Jane (certifier)	<input type="button" value="Edit"/> <input type="button" value="Submit"/>

Figure 107: Other Assumptions Saved Successfully

3.7.2.2 Upload Assumption File

Follow these steps to upload an assumption file to the Module:

1. Click the **Other Assumptions** file tab.

The **Create Assumption or Upload Assumption File** window displays. The Module automatically defaults to the **Create Assumption** radio button.

2. Click the **Upload Assumption File** radio button.

A **Manufacturer Name (required)** drop-down menu and empty **File Description (required)** field display. Refer to *Figure 108*.

[Back to the Welcome page](#) [User Guide \(PDF\)](#) [Help Desk](#) [My Profile](#)

Assumptions	
Reporting Period (required) <input type="button" value="Q1 2025"/> <input type="button" value="Next"/>	
<input type="radio"/> Create Assumptions	<input checked="" type="radio"/> Upload Assumptions
Manufacturer Name (required) <input type="button" value="- Select -"/>	
File Description (required) <div style="border: 1px solid #ccc; height: 40px; margin-bottom: 5px;"></div> <small>500 characters left</small>	
Upload .pdf, .docx, .txt, or .xlsx File (required) <small>Maximum file size is 10 MB</small> <div style="border: 1px dashed #ccc; height: 40px; margin-bottom: 5px;"></div> <small>Select File</small> <small>or drag file here</small>	

Figure 108: Upload Assumptions

3. From the **Manufacturer Name (required)** drop-down menu, click the **-Select-** drop-down menu to expand the list and select the manufacturer name.

4. In the **File Description** field, enter your assumption about a data submission. You have 500 characters of total text to comment about your submission in this section.
5. Click **Select File** to browse your desktop and upload your **Assumption File** to the Module. You may also drag your **Assumption File** into the **Select File** box.

A message opens to confirm you have successfully uploaded your **Assumption File**. Refer to *Figure 109*.

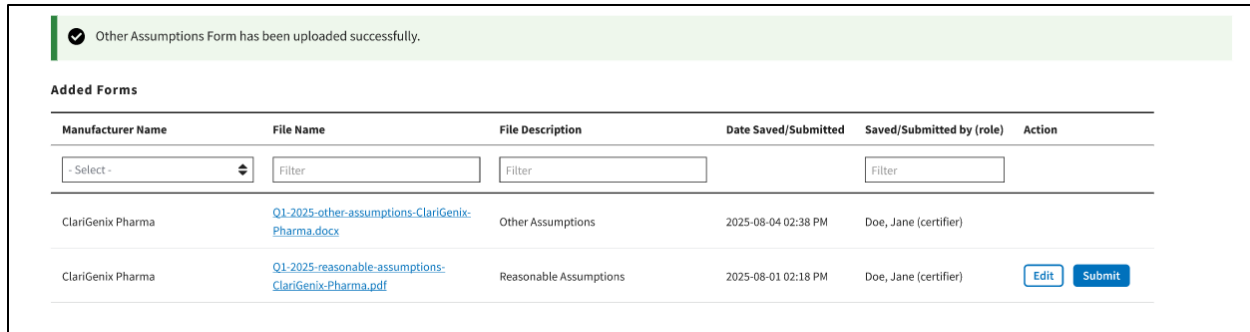

 A screenshot of a web-based application interface. At the top, a green banner displays a checkmark icon and the text 'Other Assumptions Form has been uploaded successfully.' Below this, a table titled 'Added Forms' lists two entries. The first entry is for 'ClariGenix Pharma' with a file named 'Q1-2025-other-assumptions-ClariGenix-Pharma.docx', described as 'Other Assumptions', saved on '2025-08-04 02:38 PM' by 'Doe, Jane (certifier)'. The second entry is for 'ClariGenix Pharma' with a file named 'Q1-2025-reasonable-assumptions-ClariGenix-Pharma.pdf', described as 'Reasonable Assumptions', saved on '2025-08-01 02:18 PM' by 'Doe, Jane (certifier)'. Each row has 'Edit' and 'Submit' buttons on the right.

Figure 109: Upload Assumption File - Successfully Added

4. Technical Support Contact Information

Contact the FFSDCS (ASP) Application Helpdesk for issues such as:

- Account unlock
- Password reset
- Registration process questions
- System availability escalations

Table 1 provides contact information for technical support.

Table 1: Technical Support Contacts

Email Address	Phone Number	Hours
ASPHelpDesk@dcca.com	1-844-876-0765	9:00 a.m. to 6:00 p.m. Eastern Standard Time (EST), Monday through Friday

Appendix A: Field Definitions

Table 2 provides an overview of field definitions for this document.

Table 2: Field Definitions

Column/Field Name	Format	Allowed/Sample Values	Required/Optional	Notes
Manufacturer Name	Alphanumeric	Maximum of 250 characters	Required	<ul style="list-style-type: none"> When entering product data for the same Manufacturer more than once, be sure the spelling matches. Special characters (comma, dash, period) allowed.
NDC1	5-digit number	e.g., 12345	Required	<ul style="list-style-type: none"> First segment of the National Drug Code (NDC) that identifies the labeler. Products that do not have an NDC should only use the Alternate ID column. Not required if the product has an Alternate ID. Leading zero allowed.
NDC2	4-digit number	e.g., 1234	Required	<ul style="list-style-type: none"> Not required if the product has an Alternate ID. The NDC2 is the sixth through the ninth digits of the 11-digit NDC that identifies the product.
NDC3	2-digit number	e.g., 12	Required	<ul style="list-style-type: none"> Not required if the product has an Alternate ID. The NDC3 is the last two digits of the 11-digit NDC that identify the package size.
Alternate ID	alphanumeric	maximum of 23 characters	Required	<ul style="list-style-type: none"> Not required if the product has an NDC. Must match product ID exactly as listed publicly on the manufacturer's website. Special characters (colon, dash, period) allowed.
Alternate ID Website URL	NA	e.g., http://www.medicare.gov	NA	Must have http:// or https:// prefix.

Column/Field Name	Format	Allowed/Sample Values	Required/Optional	Notes
Brand Name	Alphanumeric	Maximum of 250 characters	Optional	Enter strength and package size in their respective fields unless it is a part of the registered brand name.
Generic Name	Alphanumeric	Maximum of 250 characters	Required	Refer to valid values in Generic Name.
Volume Per Item	Numeric	NA	Required	For Alternate ID, report the volume amount in one item. (For instance, enter 10 for 10 ml in one vial, and enter 1 for powders, sheets, or patches.)
Unit for Volume per Item	NA	NA	NA	See valid value in Unit of Volume per Item. For example, for Alternate ID, select EACH for powders, sheets, or patches.
Number of Items Per NDC or Alternate ID	Numeric	Maximum of 9 digits and 2 decimal places	Required	<ul style="list-style-type: none"> For NDCs: Indicates the number units within the NDC package (for instance, enter 5 for 5 vials in a package). For Alternate IDs: Indicates the number of units within the Alternate ID. (for instance, enter 5 for 5 grafts in a package).
Package Type	Alphanumeric	2 characters	Required	Enter SD, MD, or NA. (SD = Single dose, MD = Multi dose, NA = Not Applicable)
Strength	Numeric	e.g., 300	Required	NA
Unit for Strength	NA	NA	NA	See valid values in Unit for Strength
FDA Application Number/Registration Number	Alphanumeric	Maximum of 6 characters	Required	<ul style="list-style-type: none"> Enter FDA Application Number for NDCs and Registration Number for Alternate IDs. Enter Facility Registration Number for Human Cells, Tissues, and Cellular and Tissue-Based Products (HCT/Ps).
FDA Application Supplement Number	Alphanumeric	Maximum of 9 characters	Optional	NA

Column/Field Name	Format	Allowed/Sample Values	Required/Optional	Notes
Additional FDA Application Number #1	Alphanumeric	Maximum of 6 characters	Optional	NA
Additional FDA Application Supplement Number #1	Alphanumeric	Maximum of 9 characters	Optional	NA
Additional FDA Application Number #2	Alphanumeric	Maximum of 6 characters	Optional	NA
Additional FDA Application Supplement Number #2	Alphanumeric	Maximum of 9 characters	Optional	NA
FDA Approval/Registration Date	MM/DD/YYYY	e.g., 01/01/2023	Required	Must be prior to the current submission period start date.
FDA Approval Type	NA	NA	Required	Refer to valid values in FDA Approval Type.
First Marketing Date	MM/DD/YYYY	e.g., 01/01/2023	Required	<ul style="list-style-type: none"> Must be on or after the FDA Approval Date. Must be prior to the current submission period start date. If the date is after the current submission period start date, it must be submitted as an off-cycle submission. NDC: For drugs marketed under an FDA-approved application (e.g., Abbreviated New Drug Application (ANDA), Biologics License Application (BLA), New Drug Application (NDA)), the earliest date the drug was first marketed under the application number by any labeler. If a drug was purchased or otherwise acquired from another labeler, the First Marketing Date should be equal to the First Marketing Date of the original product.

Column/Field Name	Format	Allowed/Sample Values	Required/Optional	Notes
First Marketing Date (continued)	MM/DD/YYYY	e.g., 01/01/2023	Required	<ul style="list-style-type: none"> Alternate ID: For products marketed under an FDA-approved application/registration (e.g., 510(k), HCT/P, Premarket Approval (PMA)), the earliest date the product was first marketed under the application/registration number by any labeler. If a product was purchased or otherwise acquired from another labeler, the date should be equal to the First Marketing Date of the original product.
Date of First Sale for this Product	MM/DD/YYYY	e.g., 01/01/2023	Required	<ul style="list-style-type: none"> Must be after the First Marketing Date. Must be prior to the current submission reporting period start date unless it is an off-cycle submission. NDC: The date of first sale of individual NDCs. Alternate ID: The date of first sale of individual Alternate IDs.

Appendix B: Revision History

Table 3 provides a revision history for this document.

Table 3: Revision History

Version Number	Date	Author/Editor	Description of Change
1.0	03/15/2024	Index Analytics/DCCA	Initial version of ASP Data Collection System Submitter User Guide
2.0	07/11/2025	Index Analytics/DCCA	<ul style="list-style-type: none">• Updated based on updates to the ASP Data Collection System.• Made various font, grammatical, punctuation, shading, formatting, date, version, pagination, glossary, and alignment corrections.
2.1	08/29/2025	Index Analytics/DCCA	<ul style="list-style-type: none">• Updated <i>Section 3.7</i>.• Made various font, grammatical, punctuation, shading, formatting, date, version, pagination, glossary, and alignment corrections.

Appendix C: Glossary

Table 4 provides a list of terms, acronyms, and definitions in this document.

Table 4: Glossary

Expanded Form	Acronym/Term	Definition
510(k)	NA	A 510(k) submission is the mechanism through which the majority of medical devices obtain U.S. marketing clearance. Such devices include catheters, contact lenses, and absorbable sutures.
Abbreviated New Drug Application	ANDA	An ANDA is an application for a U.S. generic drug approval for an existing licensed medication or approved drug. Authorized generics do not require ANDAs.
Average Sales Price	ASP	ASP refers to the price at which an organization typically sells a certain class of good or service. CMS uses manufacturer-reported ASPs, based on manufacturers' actual quarterly drug sales, to calculate provider payment amounts for these drugs. Federal law defines the price.
Biologics License Application	BLA	A BLA is used to request permission to introduce or deliver a biologic product into interstate commerce.
Center for Medicare Management	CMM	The CMM oversees the fee-for-service Medicare program.
Centers for Medicare & Medicaid Services	CMS	CMS is a federal agency within the U.S. Department of Health and Human Services that administers the Medicare program and works in partnership with state governments to administer Medicaid, the State Children's Health Insurance Program, and health insurance portability standards.
Consolidated Appropriations Act, 2021	CAA	The CAA establishes protections for consumers related to surprise billing and transparency in health care. The No Surprises Act (NSA) is part of the CAA.
Eastern Standard Time	EST	EST is the standard time in the 5th time zone west of Greenwich, reckoned at the 75th meridian. This time zone is in the eastern part of the United States.
Fee-for-Service Data Collection System	FFSDCS	The FFSDCS is an instrument to collect cost, revenue, utilization, and other information for FFS claims.
Human Cells, Tissues, and Cellular Products	HCT/P	HCT/Ps include human cells or tissue intended for implantation, transplantation, infusion, or transfer into a human recipient. The FDA Center for Biologics Evaluation and Research (CBER) regulates HCT/Ps.
Interactive Voice Response	IVR	IVR is a technology that allows a computer to detect voice and DTMF keypad inputs.

Expanded Form	Acronym/Term	Definition
Medicare	NA	Medicare is the federal system of health insurance for people over 65 years of age and for certain younger people with disabilities.
Medicare Part B	NA	Medicare Part B is the part of Medicare that covers doctor services, outpatient hospital care, and other medical services that Part A does not cover such as physical and occupational therapy, X-rays, medical equipment, or limited ambulance service.
New Drug Application	NDA	An NDA is the vehicle through which drug sponsors formally propose that the FDA approve a new pharmaceutical drug for sale and marketing.
Okta	NA	Okta is an enterprise-grade, identity management service, built for the cloud, but compatible with many on-premises applications.
One-Time Password	OTP	An OTP is a password that is valid for only one login session or transaction.
Premarket Approval	PMA	PMA is the FDA process of scientific and regulatory review to evaluate the safety and effectiveness of Class III medical devices. Such devices include implants, ventilators, and pacemakers.
Short Message Service	SMS	SMS is a text messaging service component of phone, web, or mobile communication systems. It uses standardized communication protocols to allow fixed-line or mobile phone devices to exchange short text messages.
Social Security Act	SSA	The SSA is a law that provides income to retired workers aged 65 or older.
Uniform Resource Locator	URL	The URL is a global address of documents and other resources on the World Wide Web.

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