

Washington State Health Care Authority

Manufacturer Data Submission Guide

Drug Price Transparency – RCW 43.71C Version 3.0

Effective Date: 3/1/2023



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About

In 2019, the Washington State Legislature passed a law (<u>Chapter 43.71C Revised Code of Washington</u>) which created the Drug Price Transparency (DPT) program at Health Care Authority (HCA). The law requires issuers of health insurance, pharmacy benefit managers (PBMs), manufacturers, and pharmacy service administrative organizations (PSAOs), to submit data on drug costs and pricing to HCA. HCA will use the data to create annual reports that demonstrate the overall impact that drug costs, rebates, and other discounts have on health care premiums.

You may visit HCA website for more information about the Drug Price Transparency program.

https://www.hca.wa.gov/about-hca/clinical-collaboration-and-initiatives/prescription-drug-cost-transparency

HCA developed this submission guide with input from stakeholders, which allowed stakeholders to review and comment on the draft data submission guide, prior to publishing the final guide. HCA has final approval authority over the data submission guides and all subsequent changes.

For recent updates about the Drug Price Transparency (DPT) program, please see the link below:

https://www.hca.wa.gov/billers-providers-partners/prescription-drug-cost-transparency-update

Contacts

Compliance Questions or General Program Questions

For compliance questions or general questions about the Drug Price Transparency program, not related to technical data submissions, please contact the program staff by sending an email to:

drugtransparency@hca.wa.gov

Technical Support

For technical assistance related to questions about data definitions, formatting, or the data submission process, please contact the technical support staff by sending an email to:

HCADPTTechSupport@hca.wa.gov

Definitions

"Authority" means the Health Care Authority.

"Calendar days" means the same as in Washington Administrative Code 182-526-0010.

"Calendar year" means the period from January 1 to December 31 of each year.

"Covered drug" means any prescription drug that:

- (a) A covered manufacturer intends to introduce to the market in Washington State at a wholesale acquisition cost of ten thousand dollars or more for a course of treatment lasting less than one month or a thirty-day supply, whichever period is longer; or
- (b) Meets all of the following:
 - (i) Is currently on the market in Washington state;
 - (ii) Is manufactured by a covered manufacturer; and



- (iii) Has a wholesale acquisition cost of more than one hundred dollars for a course of treatment lasting less than one month or a thirty-day supply, and, taking into account only price increases that take effect after July 28, 2019, the manufacturer increases the wholesale acquisition cost such that:
 - (A) The new wholesale acquisition cost is twenty percent higher than the wholesale acquisition cost on the same day of the month, twelve months before the date of the proposed increase; or
 - (B) The new wholesale acquisition cost is fifty percent higher than the wholesale acquisition cost on the same day of the month, thirty-six months before the date of the proposed increase.

"Covered manufacturer" means a person, corporation or other entity engaged in the manufacture of prescription drugs sold in or into Washington state. "Covered manufacturer" does not include a private label distributor or retail pharmacy that sells a drug under the retail pharmacy's store label, or a prescription drug repackager.

"Data" means all data provided to the authority under RCW 43.71C.020 through 43.71C.080 and any analysis prepared by the authority.

"Data submission guide" means the document that identifies the required data to be reported under RCW 43.71C and provides instructions for submitting this data to the authority, including guidance on required format.

"Food and drug administration (FDA) approval date" means the deadline for the FDA to review applications for new drugs or new biologics after the new drug application or biologic application is accepted by the FDA as complete in accordance with the Prescription Drug User Fee Act of 1992 (106 Stat. 4491; P.L. 102-571).

"Introduced to market" means marketed in Washington State.

"Pipeline drug" means a drug or biologic product, not yet approved by the Food and Drug Administration, for which a manufacturer intends to seek initial approval from the Food and Drug Administration under an original new drug application under 21 U.S.C. Sec. 355(b) or under a biologics license application under 42 U.S.C. Sec. 262to be marketed in Washington State.

"Prescription drug" means a drug regulated under chapter 69.41 or 69.50 RCW, including generic, brand, specialty, and biological products that are prescribed for outpatient use and distributed in a retail setting.

"Rebate" means negotiated price concessions, discounts, however characterized, that accrue directly or indirectly to a reporting entity in connection with utilization of prescription drugs by reporting entity members including, but is not limited to, rebates, administrative fees, market share rebates, price protection rebates, performance-based price concessions, volume-related rebates, other credits, and any other negotiated price concessions or discounts that are reasonably anticipated to be passed through to a reporting entity during a coverage year, and any other form of price concession prearranged with a covered manufacturer, dispensing pharmacy, pharmacy benefit manager, rebate aggregator, group purchasing organization, or other party which are paid to a reporting entity and are directly attributable to the utilization of certain drugs by reporting entity members.

"Reporting entity" means carriers, covered manufacturers, health carriers, health plans, pharmacy benefit managers, and pharmacy services administrative organizations, which are required to or voluntarily submit data according to chapter 43.71C RCW.

"Wholesale acquisition cost" means, with respect to a prescription drug, the manufacturer's list price for the drug to wholesalers or direct purchasers in the United States, excluding any discounts, rebates, or reductions in price, for the most recent month for which the information is available, as reported in wholesale acquisition cost guides or other publications of prescription drug pricing.



Submission Schedule

The table below describes the schedule of submissions that are required for you to be in compliance with the DPT program.

Report Type	Submission Due Date	Description
Covered Drugs with a qualifying price increase	(a)Sixty days in advance of a qualifying prices increase for a covered drug marketed in Washington State; or Within thirty days of a new covered drug's introduction to market in Washington State.	A covered manufacturer must submit to the authority all data specified in RCW 43.71C.050 and 43.71C.070, following the guidelines set forth in this data submission guide, for each newly marketed covered drug or a covered drug that had a qualifying price increase on or after October 16, 2020, as follows: (a) Sixty days in advance of a qualifying prices increase for a covered drug marketed in Washington State; or (b) Within thirty days of a new covered drug's introduction to market in Washington State.
New Covered Drugs	(b)Within thirty days of a new covered drug's introduction to market in Washington State.	A covered manufacturer must submit to the authority all data specified in RCW 43.71C.050 and 43.71C.070, following the guidelines set forth in this data submission guide, for each newly marketed covered drug or a covered drug that had a qualifying price increase on or after October 16, 2020, as follows: (a) Sixty days in advance of a qualifying prices increase for a covered drug marketed in Washington State; or (b) Within thirty days of a new covered drug's introduction to market in Washington State.
New Drug Application (notice from FDA that drug will be reviewed by deadline)	Within sixty calendar days of the manufacturer receiving the FDA approval date.	A manufacturer must submit to the authority all data specified in RCW 43.71C.060(1), following the guidelines set in the authority's applicable data submission guide for all new drug applications or biologic license applications for pipeline drugs submitted on or after October 16, 2020, within sixty calendar days of the manufacturer receiving the FDA approval date.

How to Register

In order to submit data to HCA, you must first complete the registration process and receive credentials for the Secure File Transfer (SFT) service offering hosted by Washington Technology Solutions (WATECH).

To register, you must complete and submit the registration form to HCA. You can access the form at the link below. Once you've completed the required information in the form, click the "Submit" button to generate an email.

Registering thirty days in advance of a reporting due date for this program is strongly encouraged, in order to ensure ample time to be added to the system. Once your registration is processed, you will receive a user ID and password from HCA to access the SFT service to submit data to HCA.

https://www.hca.wa.gov/assets/billers-and-providers/13-0051-drug-price-transparency-submitter-registration.pdf

Please email <u>DrugTransparency@hca.wa.gov</u> for any questions or concerns about the form and the registration process.



How to Submit

You must sign up for a Secure Access Washington Account (SAW). The email address used for this SAW account must match the DPT primary or secondary contact email address. Please note you are only allowed 2 (two) contacts. This is a change from the previous system.

SAW Instructions

Portal Instructions (also listed at the end of this document)

Submission Specifications

Data Validation

Data validation is a two-step process and at any time submissions may be rejected. Each submitted file undergoes technical and program validation to ensure that the data meets the requirements of RCW 43.71C and is compatible with HCA's reporting software. These primarily cover verification of data types (number vs. string) and formats (2023-01-01 vs. 01/01/2023). The program validation process is performed by program staff after technical validation and includes additional checks of the files to complete the data validation process.

If your report is rejected during Program validation, you will need to resubmit a corrected report within 10 business days of receipt of the rejection notice.

- Step 1 Technical validation You will receive immediate confirmation whether your submission passed or failed Technical Validation. If your submission passed Technical Validation a message indicating your submission was successfully uploaded will appear on the screen. If your submission failed, you would see an error log noting a list of all errors that must be corrected. All errors must be corrected prior to clicking the submit button. If you do not receive an email notification of either success or failure within 72 hours of submitting your report, please contact DPT program staff at drugtransparency@hca.wa.gov to confirm that your
- Step 2 Program validation An analyst will validate information submitted in ensure it meets program requirements. You will receive an approval email or a rejection email. This email will be sent to the email provided when you registered. If your report is rejected, you will need to resubmit within 10-days.

Each submitted file undergoes technical and program validation to ensure that the data meets the requirements of RCW 43.71C and is compatible with HCAs reporting software. The technical validation process is automated and applied shortly after submission to ensure that the data meets all of the technical rules described in the Table Specifications. These primarily cover verification of data types (number vs. string) and formats (2021-01-01 vs. 01/01/2021). The program validation process is performed by program staff after technical validation and includes additional checks of the files to complete the data validation process.

If you need help understanding your error log, the <u>Data Submission FAQ</u> clarifies the meaning of the error and provides guidance on corrections, or you may submit your questions to <u>HCADPTTechSupport@hca.wa.gov</u> for assistance.

Resubmissions

Failed Program Validations

In the event that your submission is rejected, you have 10 days after you receive the initial rejection notice to make necessary corrections and resubmit. You may <u>request an extension</u> of the due date subject to HCA approval. If you fail to comply with reporting requirements after receiving a rejection notice, the authority may assess a fine as allowed under WAC 182-51-1300.



To ensure HCA receives the resubmission, use the same file name (including the YYYYMMDD value) used on the first submission.

For example, if you submitted the file 'manufacturer_covered_drugs_2023_M12345_20230301.csv', and received a rejection, after making corrections you should resubmit the file 'manufacturer_covered_drugs_2023_M12345_20230301.csv' with the same name as it was originally submitted under, even if the date of resubmission is a different date.

Corrective Submissions

In the event that you find an error in your approved submission, you will need to fill out the <u>Resubmission</u> form which can be found on our portal prior to resubmitting your report. You will need to let HCA know which report you will be resubmitting and the specific reasons why you request to resubmit. HCA will review your request and approve or deny your request within 5 business days. In the event your resubmission is rejected during validation, you would be subject to the 10 day limit for correcting rejected resubmissions.

File Specifications

All files submitted must be text files with comma-separated values (CSV). The text should be encoded using the UTF-8 standard. Line endings in UNIX ("\n") or Windows ("\r\n") format are both acceptable. The header row must be included in every file. For detailed technical guidance, see the <u>Library of Congress CSV Definition</u>.

Appropriately formatted files can also be generated via Microsoft Excel by saving a spreadsheet in CSV format. This will remove many of the features included in Excel, such as formatting, formulas, and "sheets", so you may want to save a copy in Excel format for your own reference in the future. We recommend using Microsoft Excel 2016 or earlier for the submission guide templates. Using Microsoft Excel 2019 or Microsoft 365 can cause formatting issues when saving as a CSV file and result in errors.

File names should follow the naming scheme specified for the specific data that you are submitting. See Table Specifications section for more information.

Data Specifications

Nullable: <u>All fields are required</u>, unless otherwise indicated in the table specification. A field that is not required, will be indicated with the word "Nullable" in the specification. In those cases, you must leave that field blank. Do NOT provide the value as "NULL", or otherwise provide a special indicator of a null value. In all other cases, providing a blank value will result in a rejection by the automated validation.

Date Formats: Unless otherwise specified, all dates should be reported in <u>ISO-8601</u> format with hyphens between years, months, and days: "YYYY-MM-DD". For example, December 1, 2022, would be recorded as "2022-12-01".

Important note about Excel version: We recommend using Microsoft Excel 2016 or earlier for the submission guide templates. Using Microsoft Excel 2019 can cause formatting issues when saving as a CSV file and result in the file being rejected.

Table Specifications

Covered Drugs with a Qualifying Price Increase



This report contains all of the fields necessary to comply with the notification of a price increase and covered drug as described in RCW 43.71C.050 and 43.71C.070. Files submitted for manufacturer covered drugs should be named using the following schema, where ID is the manufacturer ID assigned to you by HCA during the registration process (Washington DPT Number), YYYY is the current calendar year, and YYYYMMDD is a placeholder for the submission date. In the case of a resubmission after file rejection, please use the same value for YYYYMMDD as the file that was rejected. Do not replace "manufacturer" with your organizations name, this will result in your submission being rejected.

File naming schema: manufacturer_price_increase_{YYYY}_{ID}_{YYYYMMDD}.csv Example: manufacturer_price_increase_2023_M12345_20230301.csv (Please use the submission due date, not the date the report was prepared)

For example:

manufacturer_price_increase_2021_M12345_20210301.csv or manufacturer_price_increase_2022_M12345_20220301.csv or manufacturer_price_increase_2023_M12345_20230301.csv

Please see the Submission Schedule for details regarding the timelines for submitting reports for covered drugs with a qualifying price increase.

Specification	Description	
Name: Washington DPT Number Type: String Max Length: 6 characters Format: ABCDE		icy (DPT) assigned unique submitter identifier Health Care Authority Drug Price
	MXXXXX, SXXXXX or PXXXX	rou and follows a format of either CXXXXX, KX where C, M, S and P indicate whether you r, PSAO or PBM. The X's are numeric digits
	Entity Type	Washington DPT Number
	Carrier	C12345
	Manufacturer	M12345
	PSAO	S12345
	PBM	P12345
Name: Manufacturer Name Type: String Max Length: 80 characters Format: ABCDE	•	o markets the drug. This entity has the de in the following data field.
Name: Labeler Code Type: Numeric Format: 00000 Max Length: 5 digits	_	by Food and Drug Administration (FDA) These first 5 digits of all submitted NDCs in this



Name: NDC Type: Numeric

Format: 00000000000 Max Length: 11 digits Min Length: 11 digits A three-segment code maintained by the Federal Food and Drug Administration that includes a labeler code, a product code, and a

package code for a drug product.

Example: 00012345678

NOTE: The NDC field must be eleven digits long and maintain leading

Name of the drug for the NDC reported. Only include ingredient name.

zeros.

Name Drug Name

Type: String

Max Length: 100 characters

Format: ABCDE

For example:

NDC	Drug Name	Drug Product Name	Label Name
0000000000	EFAVIRENZ-	EFAVIRENZ-	ATRIPLA
	EMTRICITABINE-	EMTRICITABINE-	
	TENOFOVIR	TENOFOVIR	
	DISOPROXIL	DISOPROXIL	
	FUMARATE	FUMARATE	
		10MG TABLET	
0000000000	ADALIMUMAB	ADALIMUMAB	HUMIRA
		PEN INJ	
		40MG/0.8	
0000000000	ADALIMUMAB	ADALIMUMAB	HUMIRA
		PEN INJ	CD/UC/HS
		CD/UC/HS	STARTER
0000000000	AMOXICILLIN	AMOXICILLIN 500	AMOXICILLIN
		MG TABLET	
0000000000	AMOXICILLIN	AMOXICILLIN 500	AMOXICILLIN
		MG CAPSULE	

NOTE: Special characters, hyphens, symbols, or slashes are allowed.



Name: Drug Product Name

Type: String

Max Length: 100 characters

Format: ABCDE

Name of the drug product for the NDC reported, to include ingredient name as reported in standardized drug databases. This name should include ingredient, salt form, dosage form, strength, and any other information specific to the NDC.

NDC	Drug Name	Drug Product Name	Label Name
0000000000	EFAVIRENZ- EMTRICITABINE- TENOFOVIR DISOPROXIL FUMARATE	EFAVIRENZ- EMTRICITABINE- TENOFOVIR DISOPROXIL FUMARATE 10MG TABLET	ATRIPLA
00000000000	ADALIMUMAB	ADALIMUMAB PEN INJ 40MG/0.8	HUMIRA
00000000000	ADALIMUMAB	ADALIMUMAB PEN INJ CD/UC/HS	HUMIRA CD/UC/HS STARTER
0000000000	AMOXICILLIN	AMOXICILLIN 500 MG TABLET	AMOXICILLIN
0000000000	AMOXICILLIN	AMOXICILLIN 500 MG CAPSULE	AMOXICILLIN

NOTE: Special characters, hyphens, symbols, or slashes are allowed.

Proprietary or legal name as marketed by manufacturer.

NDC	Drug Name	Drug Product Name	Label Name
0000000000	EFAVIRENZ-	EFAVIRENZ-	ATRIPLA
	EMTRICITABINE-	EMTRICITABINE-	
	TENOFOVIR	TENOFOVIR	
	DISOPROXIL	DISOPROXIL	
	FUMARATE	FUMARATE	
		10MG TABLET	
0000000000	ADALIMUMAB	ADALIMUMAB	HUMIRA
		PEN INJ	
		40MG/0.8	
0000000000	ADALIMUMAB	ADALIMUMAB	HUMIRA
		PEN INJ	CD/UC/HS
		CD/UC/HS	STARTER
0000000000	AMOXICILLIN	AMOXICILLIN 500	AMOXICILLIN
		MG TABLET	
0000000000	AMOXICILLIN	AMOXICILLIN 500	AMOXICILLIN
		MG CAPSULE	

NOTE: Special characters, hyphens, symbols, or slashes are allowed.

Name: Label Name

Type: String

Max Length: 100 characters

Format: ABCDE



	Health Care Muthority
Name: Drug Type	Drug Type is one of following values:
Type: Choice	
Choices: S, N, I	Single Source (S) – Drugs with an FDA New Drug Application (NDA), or biologics with a Biologics License Application (BLA), and for drugs, there are no generic alternatives available on the market. Non-Innovator Multiple-Source (N) – Drugs with an FDA Abbreviated
	New Drug Application (ANDA).
	Innovator Multiple-Source (I) – Drugs with an NDA and no longer have
	patent exclusivity.
Name: Unit of Measure	Unit of Measure for Wholesale Acquisition Cost (WAC) defined as one
Type: Choice	of the following values:
Choices: AHF, CAP, SUP, GM, ML, TAB,	
TDP, EA	AHF: Anti-hemophilia factor
	CAP: Capsule
	SUP: Suppository
	GM: Gram
	ML: Milliliter
	TAB: Tablet
	TDP: Transdermal patch
	EA: Each
Name: Day Supply	Indicate estimated day supply in relation to package size.
Type: Numeric	Everyler Dackage size of 100 used once deily will equal a 100
Max Length: 100 characters Format: 99999	Example: Package size of 100 used once daily will equal a 100.
Format: 99999	Package supply of 100 used 5 x's a day will equal a 20. (If the drug used to treat an acute condition with a maximum dose per day, then use that
	maximum dosage in estimating the day supply).
Name: Package Size	The package size identifies the number of billing units (as specified by
Type: Numeric	the labeled quantity) in the package the pharmacist uses to dispense;
Format: 9999999999999	for example, 100 tablets, 1000 capsules, or 20 ml vial. The package
Max Length: 14 digits	quantity complies with the National Council of Prescription Drug
-	Programs (NCPDP) Billing Unit Standard.
Name: Maximum Unit	Maximum unit per day based on max dose on FDA label.
Type: Numeric	
Format: 999	
Name: Course of Treatment	Is the complete course of treatment expected to be less than one
Type: Choice	month or a 30-day supply.
Format: Y, N	Choice:
	Choice.
	Y – Course of Treatment is expected to be less than one month or a 30-
	day supply.
	N – Course of Treatment is not expected to be less than one month or a
	30-day supply.
Name: Minimum Day Supply	What is the minimum day supply for a course of treatment.
Type: Numeric	
Format: 999	
Rule: Required if field Course of	Note: Fill out minimum and maying and development of the course the
Treatment is "Y"	Note: Fill out minimum and maximum day supply even if they are the same number.
	same number.



	Health Care Kuthority
Name: Maximum Day Supply Type: Numeric Format: 999	What is the maximum day supply for a course of treatment.
Rule: Required if field Course of Treatment is "Y"	Note: Fill out minimum and maximum day supply even if they are the same number.
Name: Qualifying Price Increase Type: Choice Choices: Y, N	Indicator for qualifying price increase. Manufacturer must use this field as 'yes' or 'no' to indicate if the drug meets the criteria of a qualifying price increase as defined in RCW 43.71C.010(8).
	If you are looking to report for a new covered drug use the New Covered Drug report.
Name: WAC - Current (Unit Price)	The wholesale acquisition cost per unit of measure on the date of the
Type: Numeric	submission (60 days prior to the effective date of the WAC increase). If
Format: 999999999999	the covered drug report is for a drug being introduced to the market,
Max Length: 14 digits	then leave blank.
	NOTE: Do not include the dollar sign or commas.
Name: WAC - Current (Package Price)	The wholesale acquisition cost per package on the date of the
Type: Numeric	submission (60 days prior to the effective date of the WAC increase). If
Format: 999999999999	the covered drug report is for a drug being introduced to the market,
Max Length: 14 digits	then leave blank. NOTE: Do not include the dollar sign or commas.
Name: WAC – New (Unit Price)	The new wholesale acquisition cost (WAC) per unit of measure on the
Type: Numeric	WAC effective date. If the covered drug report is for a new covered
Format: 999999999999	drug being introduced to the market, then this field should be the WAC
Max Length: 14 digits	on the date the product is first available.
Rule: Required when "WAC Type" field is	
"Unit" or "Both"	NOTE: Do not include the dollar sign or commas.
Nullable if WAC Type = "Package"	
Name: WAC – New (Package Price)	The new wholesale acquisition cost (WAC) per package on the WAC
Type: Numeric	effective date. If the covered drug report is for a new covered drug
Format: 999999999999	being introduced to the market, then this field should be the WAC on
Max Length: 14 digits	the date the product is first available.
Rule: Required when "WAC Type" field is	
"Package" or "Both"	NOTE: Do not include the dollar sign or commas.
Nullable if WAC Type = "Unit"	
Name: WAC - Effective Date	Effective date of the wholesale acquisition cost increase for the drug
Type: Date	product. If the covered drug report is for a new covered drug being
Format: YYYY-MM-DD Min Year: 1900	introduced to the market, then this field should be the date the product will first be available.
Max Year: 2100	will first be available.
Name: WAC - Type	Manufacturer must indicate if reporting by package, unit price or both.
	Package – Complete WAC Increase (Package Price) and WAC – New
Name: WAC - Type Type: Choice	Package – Complete WAC Increase (Package Price) and WAC – New (Package Price) fields. Unit – Complete WAC Increase (Unit Price) and WAC – New (Unit Price)
Name: WAC - Type Type: Choice	Package – Complete WAC Increase (Package Price) and WAC – New (Package Price) fields.



Manufacturer of Drug Type: Choice Choices: Y, N Choice: Y – New manufacturer who has previously sold this drug. Name: Previous Manufacturer of Drug Type: Choice Choices: Y, N Name: Previous Manufacturer of Drug Type: Choice Choices: Y, N Mark "Y" if the drug has been manufactured by the manufacturer the previous 5 years. If "Y", the WAC for the previous 5 years must reported. Mark "N" if the drug has been manufactured by the manufacturer less than 5 years. The WAC for the previous 5 years is not required Choice: Y – Have manufactured the drug for the previous 5 years N – Have not manufactured the drug for the previous 5 years N – Have not manufactured the drug for the previous 5 years N – Have not manufactured the drug for the previous 5 years Indicate "Yes" or "No" if they are a new manufactured when "Yes" or "No" if they are a new manufactured the drug previously. Mark "Y" if the drug has been manufactured by the manufactured less than 5 years. The WAC for the previous 5 years is not required the drug for the previous 5 years N – Have not manufactured the drug for the previous 5 years Amount of wholesale acquisition cost increase per unit of measured the drug product. Express this as a dollar amount up to 5 decimal places. If the covered drug report is for a new drug being introduction the market, leave blank. Rule: Required when "WAC Type" field is	r for d.
Choices: Y, N Choices: Y – New manufacturer that has not sold this drug previously. N – Existing manufacturer who has previously sold this drug. Name: Previous Manufacturer of Drug Type: Choice the previous 5 years. If "Y", the WAC for the previous 5 years must reported. Mark "N" if the drug has been manufactured by the manufactured less than 5 years. The WAC for the previous 5 years is not required. Choice: Y – Have manufactured the drug for the previous 5 years N – Have not manufactured the drug for the previous 5 years Name: WAC - Increase (Unit Price) Type: Numeric the drug product. Express this as a dollar amount up to 5 decimal places. If the covered drug report is for a new drug being introduct the market, leave blank.	r for d.
Choice: Y – New manufacturer that has not sold this drug previously. N – Existing manufacturer who has previously sold this drug. Mark "Y" if the drug has been manufactured by the manufacturer the previous 5 years. If "Y", the WAC for the previous 5 years must reported. Mark "N" if the drug has been manufactured by the manufacturer less than 5 years. The WAC for the previous 5 years is not require. Choice: Y – Have manufactured the drug for the previous 5 years N – Have not manufactured the drug for the previous 5 years Name: WAC - Increase (Unit Price) Type: Numeric Format: 999999.9999 Max Length: 11 digits Choice: Y – Have manufactured the drug for the previous 5 years Amount of wholesale acquisition cost increase per unit of measure the drug product. Express this as a dollar amount up to 5 decimal places. If the covered drug report is for a new drug being introduction the market, leave blank.	r for d.
Name: Previous Manufacturer of Drug Type: Choice Choices: Y, N Mark "N" if the drug has been manufactured by the manufacturer less than 5 years. The WAC for the previous 5 years is not required Choice: Y - Have manufactured the drug for the previous 5 years N - Have not manufactured the drug for the previous 5 years Name: WAC - Increase (Unit Price) Type: Numeric Format: 999999.99999 Max Length: 11 digits Name: Wac - Increase (Increase Increase	r for d.
N – Existing manufacturer who has previously sold this drug. Name: Previous Manufacturer of Drug Type: Choice Choices: Y, N Mark "N" if the drug has been manufactured by the manufacturer the previous 5 years. If "Y", the WAC for the previous 5 years must reported. Mark "N" if the drug has been manufactured by the manufacturer less than 5 years. The WAC for the previous 5 years is not require Choice: Y – Have manufactured the drug for the previous 5 years N – Have not manufactured the drug for the previous 5 years N – Have not manufactured the drug for the previous 5 years Type: Numeric Format: 999999.99999 Max Length: 11 digits Name: WAC - Increase (Unit Price) Type: Numeric Format: 999999.99999 Max Length: 11 digits	r for d.
Name: Previous Manufacturer of Drug Type: Choice Choices: Y, N Mark "N" if the drug has been manufactured by the manufactured the previous 5 years. If "Y", the WAC for the previous 5 years must reported. Mark "N" if the drug has been manufactured by the manufactured less than 5 years. The WAC for the previous 5 years is not required. Choice: Y - Have manufactured the drug for the previous 5 years N - Have not manufactured the drug for the previous 5 years Amount of wholesale acquisition cost increase per unit of measured the drug product. Express this as a dollar amount up to 5 decimal places. If the covered drug report is for a new drug being introduct the market, leave blank.	r for d.
Type: Choice Choices: Y, N Mark "N" if the drug has been manufactured by the manufacture less than 5 years. The WAC for the previous 5 years is not require Choice: Y – Have manufactured the drug for the previous 5 years N – Have not manufactured the drug for the previous 5 years Name: WAC - Increase (Unit Price) Type: Numeric Format: 999999.99999 Max Length: 11 digits the previous 5 years. If "Y", the WAC for the previous 5 years must reported. Mark "N" if the drug has been manufactured by the manufacture less than 5 years. The WAC for the previous 5 years is not required. Amount actually the drug for the previous 5 years Amount of wholesale acquisition cost increase per unit of measured the drug product. Express this as a dollar amount up to 5 decimal places. If the covered drug report is for a new drug being introduction the market, leave blank.	r for d.
Choices: Y, N Mark "N" if the drug has been manufactured by the manufacture less than 5 years. The WAC for the previous 5 years is not require Choice: Y – Have manufactured the drug for the previous 5 years N – Have not manufactured the drug for the previous 5 years Name: WAC - Increase (Unit Price) Type: Numeric Format: 999999.9999 Type: Numeric Format: 999999.9999 Max Length: 11 digits reported. Mark "N" if the drug has been manufactured by the manufactured less than 5 years. The WAC for the previous 5 years on the drug for the previous 5 years Amount of wholesale acquisition cost increase per unit of measured the drug product. Express this as a dollar amount up to 5 decimal places. If the covered drug report is for a new drug being introduction the market, leave blank.	r for d. e for
Mark "N" if the drug has been manufactured by the manufacture less than 5 years. The WAC for the previous 5 years is not require Choice: Y – Have manufactured the drug for the previous 5 years N – Have not manufactured the drug for the previous 5 years N – Have not manufactured the drug for the previous 5 years Amount of wholesale acquisition cost increase per unit of measure the drug product. Express this as a dollar amount up to 5 decimal places. If the covered drug report is for a new drug being introduct the market, leave blank.	d. e for
less than 5 years. The WAC for the previous 5 years is not require Choice: Y – Have manufactured the drug for the previous 5 years N – Have not manufactured the drug for the previous 5 years Amount of wholesale acquisition cost increase per unit of measure the drug product. Express this as a dollar amount up to 5 decimal places. If the covered drug report is for a new drug being introduction market, leave blank.	d. e for
less than 5 years. The WAC for the previous 5 years is not require Choice: Y – Have manufactured the drug for the previous 5 years N – Have not manufactured the drug for the previous 5 years Amount of wholesale acquisition cost increase per unit of measure the drug product. Express this as a dollar amount up to 5 decimal places. If the covered drug report is for a new drug being introduction market, leave blank.	d. e for
Choice: Y – Have manufactured the drug for the previous 5 years N – Have not manufactured the drug for the previous 5 years Name: WAC - Increase (Unit Price) Type: Numeric Format: 999999.99999 Max Length: 11 digits Choice: Y – Have manufactured the drug for the previous 5 years Amount of wholesale acquisition cost increase per unit of measure the drug product. Express this as a dollar amount up to 5 decimal places. If the covered drug report is for a new drug being introducted the market, leave blank.	e for
Y – Have manufactured the drug for the previous 5 years N – Have not manufactured the drug for the previous 5 years Name: WAC - Increase (Unit Price) Type: Numeric Format: 999999.9999 Max Length: 11 digits Y – Have manufactured the drug for the previous 5 years Amount of wholesale acquisition cost increase per unit of measure the drug product. Express this as a dollar amount up to 5 decimal places. If the covered drug report is for a new drug being introduction the market, leave blank.	
Y – Have manufactured the drug for the previous 5 years N – Have not manufactured the drug for the previous 5 years Name: WAC - Increase (Unit Price) Type: Numeric Format: 999999.9999 Max Length: 11 digits Y – Have manufactured the drug for the previous 5 years Amount of wholesale acquisition cost increase per unit of measure the drug product. Express this as a dollar amount up to 5 decimal places. If the covered drug report is for a new drug being introduction the market, leave blank.	
N – Have not manufactured the drug for the previous 5 years Name: WAC - Increase (Unit Price) Type: Numeric Format: 999999.99999 Max Length: 11 digits N – Have not manufactured the drug for the previous 5 years Amount of wholesale acquisition cost increase per unit of measure the drug product. Express this as a dollar amount up to 5 decimal places. If the covered drug report is for a new drug being introduction the market, leave blank.	
Name: WAC - Increase (Unit Price) Type: Numeric Format: 999999.99999 Max Length: 11 digits Amount of wholesale acquisition cost increase per unit of measure the drug product. Express this as a dollar amount up to 5 decimal places. If the covered drug report is for a new drug being introduction the market, leave blank.	
Type: Numeric the drug product. Express this as a dollar amount up to 5 decimal places. If the covered drug report is for a new drug being introduction the market, leave blank.	
Format: 999999.99999 places. If the covered drug report is for a new drug being introduction the market, leave blank.	ed to
Max Length: 11 digits the market, leave blank.	ed to
,	
Rule: Required when "WAC Type" field is	
"Unit" or "Both",	
Nullable if WAC Type = "Package" NOTE: Do not include the dollar sign or commas.	
Name: WAC - Increase (Package Price) Amount of wholesale acquisition cost increase per package for the	_
Type: Numeric product. Express this as a dollar amount up to 5 decimal places. If	
Format: 999999.99999 covered drug report is for a new drug being introduced to the ma	rket,
Max Length: 11 digits leave blank.	
Rule: Required when "WAC Type" field	
is "Package" or "Both"	
Nullable if WAC Type = "Unit" NOTE: Do not include the dollar sign or commas.	
Name: WAC - 1 Year Prior (Unit Price) Wholesale acquisition cost per unit of measure 12 months prior to	WAC
Type: Numeric Effective Date.	
Format: 99999999.99999	м Г - ···
Max Length: 14 digits This field must be populated if you have manufactured this drug for more years.	1 5 or
Rule: Existing Manufacturer Drug "Y", more years.	
and Pule: Pequired when "WAC Type" field	
Rule: Required when "WAC Type" field is "Unit" or "Both", value must be	
greater than zero,	
Nullable if WAC Type = "Package" NOTE: Do not include the dollar sign or commas.	
Name: WAC - 2 Year Prior (Unit Price) Wholesale acquisition cost per unit of measure 24 months prior to	WAC
Type: Numeric Effective Date.	
Format: 999999999999999999999999999999999999	
Max Length: 14 digits This field must be populated if you have manufactured this drug for	r 5 or
Rule: Existing Manufacturer Drug "Y", more years.	- - -
and	
Rule: Required when "WAC Type" field	
is "Unit" or "Both", value must be	
greater than zero,	
Nullable if WAC Type = "Package" NOTE: Do not include the dollar sign or commas.	



Name: WAC - 3 Year Prior (Unit Price) Wholesale acquisition cost per unit of measure 36 months prior to WAC Type: Numeric Effective Date. Format: 999999999.99999 Max Length: 14 digits This field must be populated if you have manufactured this drug for 5 or Rule: Existing Manufacturer Drug "Y", more years. and Rule: Required when "WAC Type" field is "Unit" or "Both", value must be greater than zero, Nullable if WAC Type = "Package" NOTE: Do not include the dollar sign or commas. Name: WAC - 4 Year Prior (Unit Price) Wholesale acquisition cost per unit of measure 48 months prior to WAC Type: Numeric Effective Date. Format: 999999999.99999 Max Length: 14 digits This field must be populated if you have manufactured this drug for 5 or Rule: Existing Manufacturer Drug "Y", more years. and Rule Required when "WAC Type" field is "Unit" or "Both", value must be greater than zero, Nullable if WAC Type = "Package" NOTE: Do not include the dollar sign or commas. Name: WAC - 5 Year Prior (Unit Price) Wholesale acquisition cost per unit of measure 60 months prior to WAC Type: Numeric Effective Date. Format: 999999999.99999 Max Length: 14 digits This field must be populated if you have manufactured this drug for 5 or Rule: Existing Manufacturer Drug "Y", more years. and Rule: Required when "WAC Type" field is "Unit" or "Both", value must be greater than zero, Nullable if WAC Type = "Package" NOTE: Do not include the dollar sign or commas. Name: WAC - 1 Year Prior (Package Wholesale acquisition cost per package 12 months prior to WAC Effective Date. Price) Type: Numeric Format: 99999999.99999 This field must be populated if you have manufactured this drug for 5 or Max Length: 14 digits more years. Rule: Existing Manufacturer Drug "Y", and Rule: Required when "WAC Type" field is "Package" or "Both", value must be

NOTE: Do not include the dollar sign or commas.

greater than zero,

Nullable if WAC Type = "Unit"



Name: WAC - 2 Year Prior (Package Wholesale acquisition cost per package 24 months prior to WAC Price) Effective Date. Type: Numeric Format: 999999999.99999 This field must be populated if you have manufactured this drug for 5 or Max Length: 14 digits more years. Rule: Existing Manufacturer Drug "Y", and Rule: Required when "WAC Type" field is "Package" or "Both", value must be greater than zero, Nullable if WAC Type = "Unit" NOTE: Do not include the dollar sign or commas. Name: WAC - 3 Year Prior (Package Wholesale acquisition cost per package 36 months prior to WAC Effective Date. Price) Type: Numeric Format: 999999999.99999 This field must be populated if you have manufactured this drug for 5 or Max Length: 14 digits more years. Rule: Existing Manufacturer Drug "Y", and Rule: Required when "WAC Type" field is "Package" or "Both", value must be greater than zero, Nullable if WAC Type = "Unit" NOTE: Do not include the dollar sign or commas. Name: WAC - 4 Year Prior (Package Wholesale acquisition cost per package 48 months prior to WAC Price) Effective Date. Type: Numeric Format: 999999999.99999 This field must be populated if you have manufactured this drug for 5 or Max Length: 14 digits more years. Rule: Existing Manufacturer Drug "Y", and Rule: Required when "WAC Type" field is "Package" or "Both", value must be greater than zero, Nullable if WAC Type = "Unit" NOTE: Do not include the dollar sign or commas. Name: WAC - 5 Year Prior (Package Wholesale acquisition cost per package 60 months prior to WAC Price) Effective Date. Type: Numeric Format: 999999999.99999 This field must be populated if you have manufactured this drug for 5 or Max Length: 14 digits more years. Rule: Existing Manufacturer Drug "Y", and Rule: Required when "WAC Type" field is "Package" or "Both", value must be greater than zero, Nullable if WAC Type = "Unit" NOTE: Do not include the dollar sign or commas. Name: Change/Improvement A narrative description of any change or improvement in the drug that Description necessitates the WAC increase. Type: String Max Length: 5000 characters Format: ABCDE Rule: value is populated when column "Qualifying Price Increase" is equal to Y



Name: Financial Factors A narrative description of the specific financial factors used to make the Type: String decision to set the WAC for a new Covered Drug or to increase the Max Length: 5000 characters wholesale acquisition cost of an existing Covered Drug. Format: ABCDE Rule: value is populated when column "Qualifying Price Increase" is equal to Y Note: Do not include hard returns. A narrative description of the specific non-financial used to make the Name: Non-financial factors decision to set the WAC for a new Covered Drug or to increase the Type: String Max Length: 5000 characters wholesale acquisition cost of an existing Covered. Format: ABCDE Rule: value is populated when column "Qualifying Price Increase" is equal to Y Note: Do not include hard returns. **Name: Patent Expiration Date** The date when all patents on the drug product will expire. Patents Type: Date owned by the manufacturer (i.e., originator or the inventor). Blanks are acceptable if the drug type field is "N" or "I". Format: YYYY-MM-DD Min Year: 1900 Max Year: 2100 Rule: Must be populated if "Drug Type = S Name: Market Entry Date The date the drug was Introduced to Market in Washington state. Type: Date Format: YYYY-MM-DD Min Year: 1900 Max Year: 2100 Name: WAC - Market Entry Type Manufacturer must indicate if reporting by package, unit price or both. Type: Choice Choice: Package, Unit or Both Choice: **Package** Unit Both Name: WAC - Unit Market Entry The wholesale acquisition cost per unit of measure for the existing Type: Numeric Covered Drug on the Market Entry Date of that Covered Drug. For new to market Covered Drugs, leave blank. Format: 999999999.99999 Max Length: 14 digits Rule: value is populated when column "Market Entry Date" is populated, and WAC Market Entry Type indicates "Unit" or "Both" Nullable if WAC Market Entry = "Package" NOTE: Do not include the dollar sign or commas. The wholesale acquisition cost per package for the existing Covered Drug Name: WAC - Package Market Entry Type: Numeric on the Market Entry Date of that Covered Drug. For new to market Format: 999999999.99999 Covered Drugs, leave blank. Max Length: 14 digits Rule: value is populated when column "Market Entry Date" is populated, and WAC Market Entry Type indicates "Package" or "Both" Nullable if WAC Market Entry = "Unit" NOTE: Do not include the dollar sign or commas.



Name: Reporting Period From Type: Date Format: YYYY-MM-DD Min Year: 1900	The starting date of the period being used to report for annual manufacturing, marketing, and advertising costs. Report the most recent completed calendar year.
Max Year: 2100	For example, if the effective date of the WAC increase is January 1, 2023, through February 28, 2023, report calendar year 2021. If the effective date of the WAC increase is March 1, 2023, through December 31, 2023, report calendar year 2022.
Name: Reporting Period To Type: Date Format: YYYY-MM-DD Min Year: 1900	The ending date of the period being used to report for annual manufacturing, marketing, and advertising costs. Report the most recent completed calendar year.
Max Year: 2100	For example, if the effective date of the WAC increase is January 1, 2023, through February 28, 2023, report calendar year 2021. If the effective date of the WAC increase is March 1, 2023, through December 31, 2023, report calendar year 2022.
Name: Manufacturing Costs Type: Numeric Format: 999999999999999999999999999999999999	The total cost to produce the number of units manufactured in most recent completed calendar year prior to the WAC Effective Date.
Max Length: 17 digits	For example, if the effective date of the WAC increase is January 1, 2023, through February 28, 2023, report calendar year 2021. If the effective date of the WAC increase is March 1, 2023, through December 31, 2023, report calendar year 2022, report the total cost to manufacture the drug product in calendar year 2022.
	For new to market covered drugs, fill with zeros.
	NOTE: Do not include the dollar sign or commas.
Name: Marketing and Advertising Costs Type: Numeric Format: 999999999999999999999999999999999999	Amount spent on marketing and advertising, in the most recent completed calendar year prior to the WAC Effective Date, including but not limited to direct-to-consumer marketing (television, radio print, digital, etc.), salaries for sales representatives, salaries for medical liaisons, hosted CE events and provider education, and provider detailing.
	For example, if the effective date of the WAC increase is January 1, 2023, through February 28, 2023, report calendar year 2021. If the effective date of the WAC increase is March 1, 2023, through December 31, 2023, report calendar year 2022, report calendar year 2022.
	For new to market covered drugs, leave blank.
	NOTE: Do not include the dollar sign or commas.
Name: Clinical Trials Costs Type: Numeric Format: 999999999999999999999999999999999999	Total costs for all clinical trials for the covered drug.
Max Length: 17 digits	NOTE: Do not include the dollar sign or commas.



	Health Care Muthority
Name: Research and Development Costs	Total expenditure on research and development prior to Market Entry Date.
Type: Numeric	
Format: 999999999999999999999999999999999999	
Max Length: 17 digits	NOTE: Do not include the dollar sign or commas.
Name: Regulation Costs	All costs paid by the manufacturer to the FDA and any other regulatory
Type: Numeric	body for considering their drug application and bringing the drug to
Format: 999999999999999999999999999999999999	market.
Max Length: 17 digits	
	NOTE: Do not include the dollar sign or commas.
Name: Acquired from Previous	Indicator for whether the drug was acquired from another
Manufacturer	manufacturer. Manufacturer must use this field as 'yes' or 'no' to
Type: Choice	indicate if the drug meets the criteria in RCW 43.71C.050(4).
Choices: Y, N	
Name: Previous Owner's Name	The legal name of entity who sold the covered drug to the manufacturer.
Type: String	
Max Length: 80 characters	
Format: ABCDE	
Rule: value is populated when column	
"Acquired from Previous	
Manufacturer" is equal to Y	
Nullable if Acquired from Previous	
Manufacturer is "N"	
Name: Previous Manufacturer ID	If the drug product was purchased from another manufacturer,
Type: Numeric	repackager, or private label distributor, the labeler code as assigned by
Format: 00000	Food and Drug Administration (FDA). If previous owner does not have a
Max Length: 5 digits	labeler ID fill with 5 zeros.
Rule: value is populated when column	
"Acquired from Previous	
Manufacturer" is equal to Y	
Nullable if Acquired from Previous	
Manufacturer is "N"	
Name: Previous NDC	The NDC that was used by the original or previous manufacturer. For
Type: Numeric	new drug products that do not have a previous NDC fill with eleven
Format: 0000000000	zeros.
Max Length: 11 digits	
Min Length: 11 digits	
Rule: value is populated when column	
"Acquired from Previous	
Manufacturer" is equal to Y	
Nullable if Acquired from Previous	NOTE. The NDC field worther already digital are and prejutating leading
Manufacturer is "N"	NOTE: The NDC field must be eleven digits long and maintain leading
Name: Purchase Price	If the drug product was not developed by the manufacturer, the amount
Type: Numeric	the manufacturer paid to acquire the drug.
Format: 999999999999999999999999999999999999	the manufacturer paid to acquire the drug.
Max Length: 17 digits	
-	
Rule: value is populated when column	
"Acquired from Previous	
Manufacturer" is equal to Y	
Nullable if Acquired from Previous	NOTE. Do not include the dellawaign or common
Manufacturer is "N"	NOTE: Do not include the dollar sign or commas.



Name: Currency of Purchase

Type: String

Max Length: 50 characters

Format: ABCDE

Rule: value is populated when column

"Acquired from Previous Manufacturer" is equal to Y

Nullable if Acquired from Previous

Manufacturer is "N"

Name: Acquisition Date

Type: Date

Format: YYYY-MM-DD

Min Year: 1900 Max Year: 2100

Rule: value is populated when column

"Acquired from Previous Manufacturer" is equal to Y

Nullable if Acquired from Previous

Manufacturer is "N"

Name: WAC - Acquisition Type

Type: Choice

Choice: Package, Unit or Both

Rule: value is populated when column

"Acquired from Previous Choice:

Manufacturer" is equal to Y Package

Nullable if Acquired from Previous Unit

Manufacturer is "N" Both

Name: WAC - Acquisition (Unit Price)

Type: Numeric

Format: 999999999.99999 Max Length: 14 digits

Rule: value is populated when column

"Acquired from Previous

Manufacturer" is equal to Y and WAC Acquisition Type indicates "Unit" or

"Both"

Nullable if WAC Acquisition Type = "Package" or Acquired from Previous

Manufacturer is "N"

The country of acquisition and type currency used to acquire the drug

e.g., USD, EUR, GBP, CAD, JPY, AUD, INR, CNY, MXN, etc.

If the drug product was not developed by the manufacturer, the date the

manufacturer acquired the drug.

Manufacturer must indicate if reporting by package, unit, or both.

The wholesale acquisition cost per unit of measure for the drug product

on the acquisition date.

NOTE: Do not include the dollar sign or commas.



The wholesale acquisition cost per package for the drug product on the

Manufacturer must indicate if reporting by package, unit, or both.

Unit of Measure for WAC (prior to acquisition) defined as one of the

Name: WAC - Acquisition (Package

Price)

Type: Numeric

Format: 999999999.99999 Max Length: 14 digits

Rule: value is populated when column

"Acquired from Previous

Manufacturer" is equal to Y and WAC Acquisition Type indicates "Package" or

"Both"

Nullable if WAC Acquisition Type = "Unit" or Acquired from Previous

Manufacturer is "N" NOTE: Do not include the dollar sign or commas.

acquisition date.

Name: WAC - Prior to Acquisition Type

Type: Choice

Choice: Package, Unit or Both

Rule: value is populated when column

"Acquired from Previous **Choice:** Manufacturer" is equal to Y **Package Nullable if Acquired from Previous** Unit Manufacturer is "N" **Both**

Name: WAC - Prior to Acquisition (Unit

Price)

The wholesale acquisition cost per unit of measure for the drug product 12 months prior to the acquisition date.

Type: Numeric

Format: 999999999.99999 Max Length: 14 digits

Rule: value is populated when column

"Acquired from Previous

Manufacturer" is equal to Y and WAC Prior to Acquisition Type indicates

"Unit" or "Both"

Nullable if WAC Prior to Acquisition Type = "Package" or Acquired from

Previous Manufacturer is "N" NOTE: Do not include the dollar sign or commas.

Name: Unit of Measure - Prior to

Acquisition Type: Choice

Choices: AHF, CAP, SUP, GM, ML, TAB,

TDP, EA

Rule: value is populated when column

"WAC - Prior to Acquisition" is equal to

any non-zero value

Nullable or Acquired from Previous

Manufacturer is "N"

AHF: Anti-hemophilia factor

CAP: Capsule **SUP**: Suppository

following values:

GM: Gram ML: Milliliter

TAB: Tablet

TDP: Transdermal patch

EA: Each



Name: WAC - Prior to Acquisition (Package Price)

Type: Numeric

Rule: value is populated when column

"Acquired from Previous

Manufacturer" is equal to Y and WAC Prior to Acquisition Type indicates

"Package" or "Both"

Nullable if WAC Prior to Acquisition

Type = "Unit" or Acquired from

Previous Manufacturer is "N"
Name: Financial Assistance Program

CostsType: Numeric

Max Length: 17 digits

Rule: greater than or equal to 0

The wholesale acquisition cost per package for the drug product 12 months prior to the acquisition date.

NOTE: Do not include the dollar sign or commas.

Total cost of all financial assistance programs including financial assistance for uninsured individuals, compassionate use, patient assistance, charity care, donated drug product, etc., associated with the drug product for the calendar year prior to the WAC Effective Date. For example, if the WAC Effective Date is March 1, 2020, report the total amount spent on financial assistance programs in calendar year 2019. If no financial assistance was provided fill with zeros.

NOTE: Do not include the dollar sign or commas.

Name: Rebates
Type: Numeric

Max Length: 17 digits

Rule: greater than or equal to 0

Total amount of rebates paid out associated with the NDC in the calendar year prior to the WAC Effective Date.

For example, if the effective date of the WAC increase is between and including January 1, 2022, through February 28, 2022, report calendar year 2020. If the WAC Effective Date is March 1, 2022, report the total amount of rebates paid to any entity in calendar year 2021. If no rebates were provided fill with zeros.

NOTE: Do not include the dollar sign or commas.

Name: Cost Share Assistance

Type: Numeric

Max Length: 17 digits

Rule: greater than or equal to 0

Total amount of money paid toward lowering an <u>insured individual's</u> out of pocket expenditure for the drug product in the calendar year prior to the WAC Effective Date.

For example, if the effective date of the WAC increase is between and including January 1, 2022, through February 28, 2022, report calendar year 2020. If the WAC Effective Date is March 1, 2022, report the total amount spent on cost share assistance in calendar year 2021. If no financial assistance was provided fill with zeros.

NOTE: Do not include the dollar sign or commas.



Name: Other Financial Assistance	Total amount of all other financial assistance paid out associated with
Amount	the NDC in the calendar year prior to the WAC Effective Date.
Type: Numeric	
Format: 999999999999999999999999999999999999	For example, if the effective date of the WAC increase is between and
Max Length: 17 digits	including January 1, 2022, through February 28, 2022, report calendar
Rule: greater than or equal to 0	year 2020. I the WAC Effective Date is March 1, 2022, report the total amount of all other financial assistance paid to any entity in calendar year 2021. If no other financial assistance was provided fill with zeros.
	NOTE: Do not include the dollar sign or commas.
Name: General Comments	Any additional information you would like to submit or provide to
Type: String	explain your responses.
Format: ABCDE	
Max Length: 5000 characters	
Nullable	Note: Do not include hard returns.

Covered Drugs

This report contains all of the fields necessary to comply with the notification of a price increase and covered drug as described in RCW 43.71C.050 and 43.71C.070. Files submitted for manufacturer covered drugs should be named using the following schema, where ID is the manufacturer ID assigned to you by HCA during the registration process (Washington DPT Number), YYYY is the current calendar year, and YYYYMMDD is a placeholder for the submission date. In the case of a resubmission after file rejection, please use the same value for YYYYMMDD as the file that was rejected. Do not replace "manufacturer" with your organizations name, this will result in your submission being rejected.

File naming schema: manufacturer_new_covered_drugs_{YYYY}_{ID}_{YYYYMMDD}.csv Example: manufacturer_new_covered_drugs_2023_M12345_20230301.csv (Please use the submission due date, not the date the report was prepared)

For example:

```
manufacturer_new_covered_drugs_2021_M12345_20210301.csv or
manufacturer_new_covered_drugs_2022_M12345_20220301.csv or
manufacturer_new_covered_drugs_2023_M12345_20230301.csv
```

Please see the Submission Schedule for details regarding the timelines for submitting reports for new covered drugs.

0 10 11	
Specification	Description



Name: Washington DPT Number

Type: String

Max Length: 6 characters

Format: ABCDE

WA Drug Price Transparency (DPT) assigned unique submitter identifier upon registration with the Health Care Authority Drug Price

Transparency program.

This number is unique to you and follows a format of either CXXXXX, MXXXXX, SXXXXX or PXXXXX where C, M, S and P indicate whether you are a carrier, manufacturer, PSAO or PBM. The X's are numeric digits

e.g. 12345.

Example:

Entity Type	Washington DPT Number
Carrier	C12345
Manufacturer	M12345
PSAO	S12345
PBM	P12345

Name: Manufacturer Name

Type: String

Max Length: 80 characters

Format: ABCDE

Labeler name of entity who markets the drug. This entity has the corresponding Labeler Code in the following data field.

Name: NDC
Type: Numeric

Format: 000000000000

Max Length: 11 digits Min Length: 11 digits A three-segment code maintained by the Federal Food and Drug Administration that includes a labeler code, a product code, and a

package code for a drug product.

Example: 00012345678

NOTE: The NDC field must be eleven digits long and maintain leading

zeros.



Name: Drug Name

Type: String

Max Length: 100 characters

Format: ABCDE

Name of the drug for the NDC reported. Only include ingredient name.

For example:

NDC	Drug Name	Drug Product Name	Label Name
0000000000	EFAVIRENZ-	EFAVIRENZ-	ATRIPLA
	EMTRICITABINE-	EMTRICITABINE-	
	TENOFOVIR	TENOFOVIR	
	DISOPROXIL	DISOPROXIL	
	FUMARATE	FUMARATE	
		10MG TABLET	
0000000000	ADALIMUMAB	ADALIMUMAB	HUMIRA
		PEN INJ	
		40MG/0.8	
0000000000	ADALIMUMAB	ADALIMUMAB	HUMIRA
		PEN INJ	CD/UC/HS
		CD/UC/HS	STARTER
0000000000	AMOXICILLIN	AMOXICILLIN 500	AMOXICILLIN
		MG TABLET	
0000000000	AMOXICILLIN	AMOXICILLIN 500	AMOXICILLIN
		MG CAPSULE	

NOTE: Special characters, hyphens, symbols, or slashes are allowed.

Name of the drug product for the NDC reported, to include ingredient name as reported in standardized drug databases. This name should include ingredient, salt form, dosage form, strength, and any other information specific to the NDC.

NDC	Drug Name	Drug Product Name	Label Name
0000000000	EFAVIRENZ-	EFAVIRENZ-	ATRIPLA
	EMTRICITABINE-	EMTRICITABINE-	
	TENOFOVIR	TENOFOVIR	
	DISOPROXIL	DISOPROXIL	
	FUMARATE	FUMARATE	
		10MG TABLET	
0000000000	ADALIMUMAB	ADALIMUMAB	HUMIRA
		PEN INJ	
		40MG/0.8	
0000000000	ADALIMUMAB	ADALIMUMAB	HUMIRA
		PEN INJ	CD/UC/HS
		CD/UC/HS	STARTER
0000000000	AMOXICILLIN	AMOXICILLIN 500	AMOXICILLIN
		MG TABLET	
0000000000	AMOXICILLIN	AMOXICILLIN 500	AMOXICILLIN
		MG CAPSULE	

NOTE: Special characters, hyphens, symbols, or slashes are allowed.

Name: Drug Product Name

Type: String

Max Length: 100 characters

Format: ABCDE



Name: Label Name

Type: String

Max Length: 100 characters

Format: ABCDE

Proprietary or legal name as marketed by manufacturer.

NDC	Drug Name	Drug Product Name	Label Name
0000000000	EFAVIRENZ-	EFAVIRENZ-	ATRIPLA
	EMTRICITABINE-	EMTRICITABINE-	
	TENOFOVIR	TENOFOVIR	
	DISOPROXIL	DISOPROXIL	
	FUMARATE	FUMARATE	
		10MG TABLET	
0000000000	ADALIMUMAB	ADALIMUMAB	HUMIRA
		PEN INJ	
		40MG/0.8	
0000000000	ADALIMUMAB	ADALIMUMAB	HUMIRA
		PEN INJ	CD/UC/HS
		CD/UC/HS	STARTER
0000000000	AMOXICILLIN	AMOXICILLIN 500	AMOXICILLIN
		MG TABLET	
0000000000	AMOXICILLIN	AMOXICILLIN 500	AMOXICILLIN
		MG CAPSULE	

NOTE: Special characters, hyphens, symbols, or slashes are allowed.

Name: Drug Type

Type: Choice Choices: S, N, I Drug Type is one of following values:

Single Source (S) – Drugs with an FDA New Drug Application (NDA), or biologics with a Biologics License Application (BLA), and for drugs, there

are no generic alternatives available on the market.

Non-Innovator Multiple-Source (N) – Drugs with an FDA Abbreviated

New Drug Application (ANDA).

Innovator Multiple-Source (I) – Drugs with an NDA and no longer have

patent exclusivity.

Name: Unit of Measure

Type: Choice

Choices: AHF, CAP, SUP, GM, ML, TAB,

TDP, EA

Unit of Measure for Wholesale Acquisition Cost (WAC) defined as one

of the following values:

AHF: Anti-hemophilia factor

CAP: Capsule **SUP**: Suppository

GM: Gram **ML**: Milliliter **TAB**: Tablet

TDP: Transdermal patch

EA: Each

Name: Day Supply

Type: Numeric

Max Length: 100 characters

Format: 999

Indicate estimated day supply in relation to package size.

Example: Package size of 100 used once daily will equal a 100.

Package supply of 100 used 5 x's a day will equal a 20. (If the drug used to treat an acute condition with a maximum dose per day, then use that

maximum dosage in estimating the day supply).



	Health Care Muthority
Name: Package Size	The package size identifies the number of billing units (as specified by
Type: Numeric	the labeled quantity) in the package the pharmacist uses to dispense;
Format: 999999999999	for example, 100 tablets, 1000 capsules, or 20 ml vial. The package
Max Length: 14 digits	quantity complies with the National Council of Prescription Drug
	Programs (NCPDP) Billing Unit Standard.
Name: WAC - Effective Date	Effective date of the wholesale acquisition cost increase for the drug
Type: Date	product. If the covered drug report is for a new covered drug being
Format: YYYY-MM-DD	introduced to the market, then this field should be the date the product
Min Year: 1900	will first be available.
Max Year: 2100	
Name: WAC - Type	Manufacturer must indicate if reporting by package, unit price or both.
Type: Choice	manaracturer mast mateute in reporting by package, and price of both.
Choices: Package, Unit or Both	Package – Complete WAC Increase (Package Price) and WAC – New
Choices. Fackage, Offic of Both	
	(Package Price) fields.
	Unit – Complete WAC Increase (Unit Price) and WAC – New (Unit Price)
	fields.
	Both – Complete WAC Increase (Package Price), WAC Increase (Unit
	Price), WAC - New (Package Price) and WAC – New (Unit Price).
Name: WAC - New (Unit Price)	The new wholesale acquisition cost (WAC) per unit of measure on the
Type: Numeric	WAC effective date. If the covered drug report is for a new covered drug
Format: 9999999999999	being introduced to the market, then this field should be the WAC on
Max Length: 14 digits	the date the product is first available.
Rule: Required when "WAC Type" field	the date the product is mist available.
is "Unit" or "Both"	
	NOTE: Do not include the dellar size or common
Nullable if WAC Type = "Package"	NOTE: Do not include the dollar sign or commas.
Name: WAC - New (Package Price)	The new wholesale acquisition cost (WAC) per package on the WAC
Type: Numeric	effective date. If the covered drug report is for a new covered drug
Format: 999999999999	being introduced to the market, then this field should be the WAC on
Max Length: 14 digits	the date the product is first available.
Rule: Required when "WAC Type" field	
is "Package" or "Both"	
Nullable if WAC Type = "Unit"	NOTE: Do not include the dollar sign or commas.
Name: Financial Factors	A narrative description of the specific financial factors used to make the
Type: String	decision to set the WAC for a new Covered Drug or to increase the
Max Length: 5000 characters	wholesale acquisition cost of an existing Covered Drug.
Format: ABCDE	misicsare acquisition cost of all existing covered brug.
Rule: value is populated when column	
The state of the s	Note: Do not include hard returns
"Qualifying Price Increase" is equal to Y	Note: Do not include hard returns.
Name: Non-financial Factors	A narrative description of the specific non-financial used to make the
Type: String	decision to set the WAC for a new Covered Drug or to increase the
Max Length: 5000 characters	wholesale acquisition cost of an existing Covered.
Format: ABCDE	
Rule: value is populated when column	
"Qualifying Price Increase" is equal to Y	Note: Do not include hard returns.
Name: Patent Expiration Date	The date when all patents on the drug product will expire. Patents
Type: Date	owned by the manufacturer (i.e., originator or the inventor). Blanks are
Format: YYYY-MM-DD	acceptable if the drug type field is "N" or "I".
Min Year: 1900	,
Max Year: 2100	
Rule: Must be populated if "Drug Type	
hale. Must be populated it brug Type	
= S	



Name: Market Entry Date	The date the drug was Introduced to Market in Washington state.
Type: Date	
Format: YYYY-MM-DD	
Min Year: 1900	
Max Year: 2100	
Name: WAC - Market Entry Type	Manufacturer must indicate if reporting by package, unit price or both.
Type: Choice	
Choice: Package, Unit or Both	Choice:
	Package
	Unit
	Both
Name: General Comments	Any additional information you would like to submit or provide to
Type: String	explain your responses.
Format: ABCDE	
Max Length: 5000 characters	
Nullable	Note: Do not include hard returns.

New Drug Application

This report contains all of the data fields necessary to comply with reporting a New Drug Application to HCA, per RCW 43.71C.060.

Files submitted for manufacturer new drugs should be named using the following schema, where: ID is the manufacturer ID assigned to you by HCA during the registration process (Washington DPT Number), YYYY is the current reporting period, and YYYYMMDD is a placeholder for the submission date. In the case of a resubmission after file rejection, please use the same value for YYYYMMDD as the file that was rejected. Do not replace "manufacturer" with your organizations name, this will result in your submission being rejected.

File naming schema: manufacturer_new_drugs_{YYYY}_{ID}_{YYYYMMDD}.csv Example: manufacturer_new_drugs_2023_M12345_20230301.csv (Please use the submission due date, not the date the report was prepared)

For example:

manufacturer_new_drugs_2021_M12345_20210301.csv or manufacturer_new_drugs_2022_M12345_20220301.csv or manufacturer_new_drugs_2023_M12345_20230301.csv

Please see the Submission Schedule for details regarding the timelines for submitting reports for covered drugs.

Description	cification
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Name: Washington DPT Number

Type: String

Max Length: 6 characters

Format: ABC123

WA Drug Price Transparency (DPT) assigned unique submitter identifier upon registration with the Health Care Authority Drug Price

Transparency program.

This number is unique to you and follows a format of either CXXXXX, MXXXXXX, SXXXXXX or PXXXXXX where C, M, S and P indicate whether you are a carrier, manufacturer, PSAO or PBM. The X's are numeric digits e.g. 12345.

Example:

Entity Type	Washington DPT Number
Carrier	C12345
Manufacturer	M12345
PSAO	S12345
PBM	P12345

Name: Manufacturer Name

Type: String

Max Length: 80 characters

Format: ABCDE

Name: Labeler Code

Type: Numeric Format: 00000 Max Length: 5 digits Labeler code as assigned by Food and Drug Administration (FDA)

Labeler name of entity who manufactures and markets the drug.

Name: Drug Name

Type: String

Max Length: 100 characters

Format: ABCDE

Name of the drug for the NDC reported. Only include ingredient name.

For example:

NDC	Drug Name	Drug Product Name	Label Name
0000000000	EFAVIRENZ-	EFAVIRENZ-	ATRIPLA
	EMTRICITABINE-	EMTRICITABINE-	
	TENOFOVIR	TENOFOVIR	
	DISOPROXIL	DISOPROXIL	
	FUMARATE	FUMARATE	
		10MG TABLET	
0000000000	ADALIMUMAB	ADALIMUMAB	HUMIRA
		PEN INJ	
		40MG/0.8	
0000000000	ADALIMUMAB	ADALIMUMAB	HUMIRA
		PEN INJ	CD/UC/HS
		CD/UC/HS	STARTER
0000000000	AMOXICILLIN	AMOXICILLIN 500	AMOXICILLIN
		MG TABLET	
0000000000	AMOXICILLIN	AMOXICILLIN 500	AMOXICILLIN
		MG CAPSULE	

NOTE: Special characters, hyphens, symbols, or slashes are allowed.



Name: Drug Product Name

Type: String

Max Length: 100 characters

Format: ABCDE

Nullable

Name of the drug product for the NDC reported, to include ingredient name as reported in standardized drug databases. This name should include ingredient, salt form, dosage form, strength, and any other information specific to the NDC.

For example, "fluoxetine HCL 20 mg tablets" is acceptable.

NDC	Drug Name	Drug Product Name	Label Name
0000000000	EFAVIRENZ-	EFAVIRENZ-	ATRIPLA
	EMTRICITABINE-	EMTRICITABINE-	
	TENOFOVIR	TENOFOVIR	
	DISOPROXIL	DISOPROXIL	
	FUMARATE	FUMARATE	
		10MG TABLET	
0000000000	ADALIMUMAB	ADALIMUMAB	HUMIRA
		PEN INJ	
		40MG/0.8	
0000000000	ADALIMUMAB	ADALIMUMAB	HUMIRA
		PEN INJ	CD/UC/HS
		CD/UC/HS	STARTER
0000000000	AMOXICILLIN	AMOXICILLIN 500	AMOXICILLIN
		MG TABLET	
0000000000	AMOXICILLIN	AMOXICILLIN 500	AMOXICILLIN
		MG CAPSULE	

NOTE: Special characters, hyphens, symbols, or slashes are allowed.

Proprietary or legal name as marketed by manufacturer.

Name: Label Name or Pipeline Drug Name

Type: String

Max Length: 100 characters

Format: ABCDE

Nullable

If not approved by the FDA, then enter the name of the Pipeline Drug. For example, "AAA600".

NDC	Drug Name	Drug Product Name	Label Name
00000000000	EFAVIRENZ-	EFAVIRENZ-	ATRIPLA
	EMTRICITABINE-	EMTRICITABINE-	
	TENOFOVIR	TENOFOVIR	
	DISOPROXIL	DISOPROXIL	
	FUMARATE	FUMARATE	
		10MG TABLET	
0000000000	ADALIMUMAB	ADALIMUMAB	HUMIRA
		PEN INJ	
		40MG/0.8	
00000000000	ADALIMUMAB	ADALIMUMAB	HUMIRA
		PEN INJ	CD/UC/HS
		CD/UC/HS	STARTER
00000000000	AMOXICILLIN	AMOXICILLIN 500	AMOXICILLIN
		MG TABLET	
00000000000	AMOXICILLIN	AMOXICILLIN 500	AMOXICILLIN
		MG CAPSULE	

NOTE: Special characters, hyphens, symbols, or slashes are allowed.



	Health Care Authority		
Name: Drug Type Type: Choice	Drug Type is one of following values:		
Choices: S, N, I	Single Source (S) – Drugs that having an FDA New Drug Application (NDA), or biologics having a Biologics License Application (BLA), and there are no generic alternatives available on the market. Non-Innovator Multiple-Source (N) – Drugs that have an FDA Abbreviated New Drug Application (ANDA). Innovator Multiple-Source (I) – Drugs that have an NDA and no longer have patent exclusivity.		
Name: Application Type Type: Choice	Application Type is one of following values:		
Choices: BLA, NDA, ANDA	New Drug Application (NDA) – Drug is a pipeline drug and was submitted as a New Drug Application to the FDA.		
	Biologics License Application (BLA) – Drugs is a pipeline drug and was submitted as a Biologics License Application to the FDA.		
	Abbreviate New Drug Application (ANDA) – contains data which is submitted to FDA for the review and potential approval of a generic drug.		
Name: Regulatory Pathway Type: Choice Choices: 505(b)(1), 351(a), Other	Regulatory pathway for approval by the Food and Drug Administration. Acceptable values are $505(b)(1)$, $351(a)$ or Other.		
	If choosing "Other" please list the regulatory pathway this product was approved in General Comments.		
Name: Application Number Type: String Format: ABC123 Max Length: 6 digits Min Length: 6 digits Nullable	The application number assigned by the Food and Drug Administration. For application numbers less than 6 digits, the application number should be preceded using zeros.		
Name: Application Supplement Number Type: String Format: AB12 Max Length: 4 digits Min Length: 4 digits Nullable	The supplemental application number assigned by the Food and Drug Administration. For application numbers less than 4 digits, the supplemental application number should be preceded using zeros.		
Name: Significant Impact on State Expenditures Type: Choice Choices: Y, N	Indicator of whether the pipeline drug will cost Washington State government agencies at least \$50,000 per biennium in any future biennium. HCA believes that drugs costing at least \$50,000 per biennium for Washington State government agencies to qualify as a significant impact on state expenditures. HCA may request from the manufacturer the information in the remaining fields if HCA believes the drug will have a significant impact on state expenditures and require manufacturers to resubmit with information for all of the following fields. If manufacturers believe drugs to meet or exceed this threshold, the following fields may be completed. WAC 182-51-0700(3)		



	Health Care Muthority		
Name: Proposed Indication	The proposed indication or indications submitted on the application to		
Type: String	the FDA. Use the SNOMED CT disease term listed on the application.		
Max Length: 5000 characters Format: ABCDE	Use a semi-colon to separate multiple indications.		
Nullable	Manufacturers may submit this information voluntarily if the pipeline drug is expected to cost Washington State at least \$50,000 per biennium WAC 182-51-0700(3).		
Name: Area of Study Type: String Max Length: 5000 characters Format: ABCDE Nullable	A list of diseases, conditions, and therapeutic areas being studied for this drug and whether the chemical drug has received an indication in the FDA approved labeling for use in these diseases, conditions, or therapeutic areas.		
	Manufacturers may submit this information voluntarily if the pipeline drug is expected to cost Washington State at least \$50,000 per biennium WAC 182-51-0700(3).		
Name: Route of Administration Type: String Max Length: 5000 characters Format: ABCDE	List each route of administration being studied for this drug, including any differences between immediate-release and extended-release formulations.		
Nullable	Manufacturers may submit this information voluntarily if the pipeline drug is expected to cost Washington State at least \$50,000 per biennium WAC 182-51-0700(3).		
Name: Clinical Comparator Type: String Max Length: 5000 characters Format: ABCDE	All clinical comparators including dosage regimen being used for which to evaluate the comparative differences in safety, efficacy, effectiveness, costs, value, or any other outcomes in clinical trials.		
Nullable	Manufacturers may submit this information voluntarily if the pipeline drug is expected to cost Washington State at least \$50,000 per biennium WAC 182-51-0700(3).		
Name: PDUFA Date	Prescription Drug User Fee Act (PDUFA) date assigned by the FDA.		
Type: Date Format: YYYY-MM-DD Min Year: 1900 Max Year: 2100 Nullable	Manufacturers may submit this information voluntarily if the pipeline drug is expected to cost Washington State at least \$50,000 per biennium WAC 182-51-0700(3).		
Name: Rare Disease Indication Type: Choice Choices: Y, N	Indicator of whether the FDA assigned the drug as being defined as a treatment for a rare disease.		
Nullable	Manufacturers may submit this information voluntarily if the pipeline drug is expected to cost Washington State at least \$50,000 per biennium WAC 182-51-0700(3).		
Name: Orphan Drug Status Type: Choice Choices: Y, N	Indicator of whether the FDA assigned the drug as having an Orphan designation.		
Nullable	Manufacturers may submit this information voluntarily if the pipeline drug is expected to cost Washington State at least \$50,000 per biennium WAC 182-51-0700(3).		



	Health Care Muthority				
Name: Orphan Designation Number	Orphan designation number assigned by the FDA. For Orphan				
Type: Numeric Format: 000000	Designation numbers less than 6 digits, the supplemental application				
Max Length: 6 digits	number should be preceded using zeros.				
Min Length: 6 digits	Manufacturers may submit this information voluntarily if the pipeline				
Nullable	drug is expected to cost Washington State at least \$50,000 per				
- Tunasic	biennium WAC 182-51-0700(3).				
Name: Pediatric Indication	Indicator of whether the indication is for use in individuals under 18				
Type: Choice	years of age.				
Choices: Y, N					
Nullable	Manufacturers may submit this information voluntarily if the pipeline				
	drug is expected to cost Washington State at least \$50,000 per				
	biennium WAC 182-51-0700(3).				
Name: Fast Track Status	Indicator of whether the FDA assigned the drug as having fast track				
Type: Choice	status.				
Choices: Y, N					
Nullable	Manufacturers may submit this information voluntarily if the pipeline drug is expected to cost Washington State at least \$50,000 per				
	biennium WAC 182-51-0700(3).				
Name: Breakthrough Therapy Status	Indicator of whether the FDA assigned the drug as having breakthrough				
Type: Choice	therapy status.				
Choices: Y, N					
Nullable	Manufacturers may submit this information voluntarily if the pipeline				
	drug is expected to cost Washington State at least \$50,000 per				
	biennium WAC 182-51-0700(3).				
Name: Accelerated Approval Status	Indicator of whether the FDA assigned the drug as having accelerated				
Type: Choice	approval status.				
Choices: Y, N					
Nullable	Manufacturers may submit this information voluntarily if the pipeline				
	drug is expected to cost Washington State at least \$50,000 per biennium WAC 182-51-0700(3).				
Name: Priority Review Status	Indicator of whether the FDA assigned the drug as having priority				
Type: Choice	review status.				
Choices: Y, N	Teview status.				
Nullable	Manufacturers may submit this information voluntarily if the pipeline				
	drug is expected to cost Washington State at least \$50,000 per				
	biennium WAC 182-51-0700(3).				
Name: New Molecular Entity Status	Indicator of whether the FDA assigned the drug as having new				
Type: Choice	molecular entity status.				
Choices: Y, N					
Nullable	Manufacturers may submit this information voluntarily if the pipeline				
	drug is expected to cost Washington State at least \$50,000 per				
Names Consul Consul	biennium WAC 182-51-0700(3).				
Name: General Comments	Any additional information you would like to submit or provide to				
Type: String Format: ABCDE	explain your responses.				
Max Length: 5000 characters					
Nullable	Note: Do not include hard returns.				



Appendix A – ST Web Client User Guideline

Prerequisites

	Before vou can	log in to ST	Web Client and o	pen a session.	vou need:
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- A high-speed Internet connection
- A supported Internet browser:
 - Microsoft Internet Explorer 11
 - o Microsoft Edge latest version
 - o Mozilla Firefox latest version
 - o Apple Safari latest version
 - o Google Chrome latest version

Sign into portal

Step 1.

All entities will go to the following external portal link

https://support.hca.wa.gov/hcasupport

Step 2.

Click on "Public"

Login with your current SAW login in credentials. If you don't have a SAW account please click on "SIGN UP!"

Step 3.

Click on "Make a request".

You will now have access to all of your entities' SAW accounts.

Step 4.



First time registering – you will see "DPT Entity Registration" only

Important:

Primary and secondary contact emails must be for an individual and not a group or shared email.

Once registration is completed your entity will be assigned a unique HCA ID.

Once you have registered you will have the additional options of:

- DPT Registration Correction
 - o Update contact information
- DPT Template Submission
 - Submit reports
- DPT Re-submission/Extension
 - o Request an extension for your submission
 - o Request permission to resubmit a report that has previously been submitted (these reports have previously been accepted for both Technical Validation and Program Validation).

Update contact information click on "DPT Registration Correction Form"

You must know your Tax ID number.

You have the option of updating one or all of the following:

- Organization address
- Primary contact information
- Secondary contact information

Click on the first box you would like to update. When finished with that section click on the next section you would like to update.

Important!

You must click "Submit" when complete.

DPT Template Submission

Organization Type

- Carrier
- Manufacturer
- PBM



PSAO

Organization Name – Entity name will appear, click on it.

Template – Entity must choose which report they are uploading.

Reporting year – Entity must choose which year they are reporting for.

Click



You

will navigate to your report.

You will receive immediate feedback on errors, and may resubmit at anytime once you have corrected those errors.

row 1 col 15 (WAC Increase Rank Percent): Percent value is too large

You will also receive feedback stating "File successfully validated".

Important!

You must click "Submit" once you receive the file successfully validated in order for the report to be accepted into the Enterprise Data Warehouse (EDW).

DPT Re-submission/Extension

DPT Re-submission form is used when an entity finds an error in a report that has previously been submitted. This report has been accepted by HCA DPT for both technical and program validations. The entity is requested to resubmit this report.

Extension form is used when an entity will not be able to meet the due date of their required reports and is requesting additional time.

You first will need to choose the "Action".

- Resubmit
- Extension

Important!

You must click "Submit" in order to submit your request.