# PROPOSED RULE MAKING



# CR-102 (June 2024) (Implements RCW 34.05.320)

Do NOT use for expedited rule making

#### **CODE REVISER USE ONLY**

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DATE: November 22, 2024

TIME: 4:55 PM

WSR 24-24-028

Agency: De	partme	nt of Health – Pharmacy Quality Assurance	Commission
☐ Original I	Notice		
Supplem	ental N	lotice to WSR <u>24-14-140</u>	
☐ Continua	nce of	WSR	
□ Preprope	osal Sta	atement of Inquiry was filed as WSR 23-2	<u>1-010</u> ; <b>or</b>
☐ Expedite	d Rule	MakingProposed notice was filed as W	SR; or
☐ Proposal	is exe	mpt under RCW 34.05.310(4) or 34.05.330	(1); or
□ Proposal	is exe	mpt under RCW	
Dialysis Pro 246-945-09 legend drug further amer sell, deliver,	grams. I, 246-9 s, includ nds WA posses	The Pharmacy Quality Assurance Commiss 945-092, and 246-945-093 to include manufacting dialysate, in home dialysis program rule C 246-945-090 to add the word "may" and li	I Dialysis Device Manufacturers and Wholesalers in Home ion (commission) is proposing to amend WAC 246-945-090, acturers and wholesalers of dialysis devices and approved as under the commission's jurisdiction. This supplemental set the dialysis devices manufacturers and wholesalers may d to amend WACs 246-945-091, 246-945-092, and 246-945-
location(s)	•		
Date:		Location: (be specific)	Comment:
02/06/2025	9:30	Physical Location:	The commission will hold a hybrid hearing. Attendees are
	am	Department of Labor & Industries 7273 Linderson Way SW Tumwater, WA 98501	welcome to attend either in-person at the physical location or virtual via Zoom.
		Virtual Location: Virtual:	
		To access the meeting on February 6, 2025	
		at 9:30 am, go to	
		https://us02web.zoom.us/j/86309299195 or	
		https://zoom.us/join and use the Webinar ID	
		863 0929 9195	
		The access options include one tap mobile:	
		+12532158782,,86309299195# US	
		(Tacoma)	
		+12532050468,,86309299195# US	
		Or Telephone: Dial (for higher quality, dial	
		number based on your current location): +1 253 215 8782 US (Tacoma)	
		+1 253 205 0468 US	
Date of inte	nded a	doption: 2/06/2025 (Note: This is NOT the	•
Submit writ	ten coi	mments to:	Assistance for persons with disabilities:

•						
Name Julia Katz			Contact Julia Katz			
	7852, Olympia, WA 98504-78	852	Phone 360-236-4946			
Email Fax 360-236-226	0		Fax 360-236-2260 TTY 711			
	ss.wa.gov/doh/policyreview/		Email PharmacyRules@doh.wa.g	IOV		
·	d time) The date and time of	f this filing	Other None			
By (date and time)	January 23, 2025 at 11:59	pm	By (date) January 23, 2025			
proposal is to allow drugs, including cor amendments to WA consultation, record	manufacturers and wholesal mmercially available dialysate Cs 246-945-091, 246-945-09	ers to sell, del e, and dialysis 92, and 246-94 nce practices.	ing any changes in existing rules ver, possess, and dispense prescridevices directly to home dialysis parts of the properties of the prope	ibed approved le atients. Conform vices are conside	egend ing ered in	
required two amend is to add a list of ap home dialysis patie	Following the public rules hearing held on August 22, 2024, the commission determined that the proposed rule language required two amendments to WAC 246-945-090. The first amendment is to reinstate the term "may." The second amendment is to add a list of approved dialysis devices that manufacturers and wholesalers may sell, deliver, possess, and dispense to home dialysis patients. Amendments were needed to WAC 246-945-091, 246-945-092, and 246-945-093 in association with the WAC 246-945-090 amendments.					
and 69.41.032 to er to dialysis patients	nsure manufacturers and who and granted the commission neasures for wholesalers and	olesalers may authority to ad	ed to implement SHB 1675, which distribute approved legend drugs a opt rules. Additionally, the propose s dispensing approved legend drug	nd dialysis devic d rules establish	es directly important	
and conforming rev 092, and 246-945-0 SHB 1675 directs the	isions regarding the list of dia 191. It was decided that addin ne commission to adopt rules	alysis devices on the word "motor to implement"	c rules hearing that "may," the list owere missing from WACs 246-945- ay" corrects a typographical error a the statutes, including a list of appr	090, 246-945-09 and clarifies inter	91, 246-945- nt, and that	
	for adoption: RCW 18.64.	·	7, and 69.41.032			
Is rule necessary	emented: RCW 18.64.257	3110 69.41.032				
Federal Law				□ Yes	⊠ No	
Federal Cou	t Decision?			□ Yes	⊠ No	
State Court [	Decision?			□ Yes	⊠ No	
If yes, CITATION:			4			
matters: None	s or recommendations, if ai	iy, as to stati	tory language, implementation,	enforcement, ai	nd fiscal	
	nt: (person or organization) t: □ Private. □ Public. ⊠ Go		ty Assurance Commission			
Name of agency p	ersonnel responsible for:					
	Name	Office Loca	ation	Phone		
Drafting	Julia Katz	111 Israel	Rd SE, Tumwater, WA 98501	360-236-4	946	
Implementation	Julia Katz	111 Israel	Rd SE, Tumwater, WA 98501	360-236-4	946	
Enforcement	Marlee B. O'Neill	111 Israel	Rd SE, Tumwater, WA 98501	360-480-9	108	
Is a school district If yes, insert statem	t fiscal impact statement re ent here:	quired under	RCW 28A.305.135?	☐ Yes	⊠ No	
The public may Name Address Phone Fax	obtain a copy of the school d	istrict fiscal im	pact statement by contacting:			

1	TTY		
	Email		
	Other		
Is a cost-	benefit analysis required under RCW 34		
⊠ Ye	, , , , , , , , , , , , , , , , , , , ,	nay be obtained	by contacting:
	Name Julia Katz	DEO4 7050	
	Address PO Box 47852, Olympia, WA 98 Phone 360-236-4946	004-7002	
	Fax 360-236-2260		
	TTY		
	Email PharmacyRules@doh.wa.gov		
	Other None		
□ No	<u>'</u>		
	ry Fairness Act and Small Business Eco		Statement e (ORIA) provides support in completing this part.
	fication of exemptions:	TH AND ASSISTANCE	e (ONIA) provides support in completing this part.
		<b>be exempt</b> from	requirements of the Regulatory Fairness Act (see
	9.85 RCW). For additional information on e box for any applicable exemption(s):	exemptions, cons	ult the exemption guide published by ORIA. Please
solely to c	conform and/or comply with federal statute	or regulations. P	CW 19.85.061 because this rule making is being adopted lease cite the specific federal statute or regulation this assequences to the state if the rule is not adopted.
	alle proposal, or portions of the proposal, is 34.05.313 before filing the notice of this pro		the agency has completed the pilot rule process defined
		•	e provisions of RCW 15.65.570(2) because it was
	y a referendum.		(-) a constant
☐ This ru	le proposal, or portions of the proposal, is	exempt under Re	CW 19.85.025(3). Check all that apply:
	RCW 34.05.310 (4)(b)		RCW 34.05.310 (4)(e)
	(Internal government operations)		(Dictated by statute)
	RCW 34.05.310 (4)(c)		RCW 34.05.310 (4)(f)
	(Incorporation by reference)		(Set or adjust fees)
	RCW 34.05.310 (4)(d)		RCW 34.05.310 (4)(g)
	(Correct or clarify language)		((i) Relating to agency hearings; or (ii) process
			requirements for applying to an agency for a license or permit)
☐ This ru	lle proposal, or portions of the proposal, is	exempt under Re	CW 19.85.025(4). (Does not affect small businesses).
	le proposal, or portions of the proposal, is on of how the above exemption(s) applies t	•	
	e of exemptions: Check one.	3) Exemptions in	dentified above apply to all portions of the rule proposal.
☐ The ru proposal,		section 3.) The exide details here	exemptions identified above apply to portions of the rule (consider using this template from ORIA):
(3) Small	business economic impact statement:	Complete this se	ction if any portion is not exempt.
` ´	tion of the proposed rule is <b>not exempt</b> , do	•	re-than-minor costs (as defined by RCW 19.85.020(2))
⊠ No		or cost analysis a	and how the agency determined the proposed rule did
	of the proposed rule (\$1,669) are <u>less tha</u>	n the minor cost	threshold (\$10,305.83).
		_	,
			oufacturers and wholesalers choosing to dispense 69. Using the Governor's Office for Regulatory

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Innovation and Assistance's Minor Cost Threshold Calculator with NAICS Code Title, 424210 Drugs and Druggists' Sundries Merchant Wholesalers, the minor cost threshold is not met per RCW 19.85.020. A full SBEIS may not be required since the minor cost threshold is not met.

It was further determined that the proposed amendments to WAC 246-945-090 to add "may" and the list of dialysis devices would not affect existing cost estimates. Excerpts of the SBEIS are provided herein.

The following is a brief description of the proposed rule including the current situation/rule, followed by the history of the issue and why the proposed rule is needed. A description of the probable compliance requirements and the kinds of professional services that a small business is likely to need in order to comply with the proposed rule.

The Pharmacy Quality Assurance Commission (commission) is proposing rule amendments to increase access to dialysis devices and approved legend drugs, including dialysate, for patients undergoing kidney dialysis treatment at home by allowing manufacturers and wholesalers to dispense lawfully prescribed dialysis devices and approved legend drugs to patients' homes.

The proposed rule is necessary to implement Substitute House Bill (SHB) 1675 (chapter 23, Laws of 2022) as well as establish important quality assurance measures for wholesalers and manufacturers dispensing approved legend drugs, and dialysis devices directly to home dialysis patients. SHB 1675 amended RCW 18.64.257 and 69.41.032 to ensure manufacturers and wholesalers may distribute approved medications and devices directly to dialysis patients.

Prior to the passage of SHB 1675, manufacturers and wholesalers would need either a pharmacy or nonresident pharmacy license to dispense directly to patients. Pharmacy licenses are issued to facilities located in Washington that dispense prescriptions to patients. Nonresident pharmacy licenses are issued to facilities located outside of Washington that dispense prescriptions to patients. Both the pharmacy and nonresident pharmacy licenses require annual renewal applications and an application fee of \$575. Current statute does not require pharmacy and nonresident pharmacy licensees dispensing legend drugs to patients to have an agreement with a pharmacist to provide consultation on shipment and delivery of prescriptions, develop a quality assurance program for shipment and delivery of prescriptions, nor maintain a record of shipment and delivery errors. WAC 246-945-016 does require pharmacy and nonresident pharmacy licensees dispensing legend drugs to patients to affix labels to prescription containers. Since the passage of SHB 1675, manufacturers and wholesalers in compliance with quality assurance measures may distribute approved medications and devices directly to patients without a pharmacy or nonresident pharmacy license.

In October 2022, the commission filed a policy statement under WSR 22-21-062 to clarify the commission's position on this subject until rulemaking can be completed. Per the policy statement, the commission will not take enforcement action against a manufacturer or wholesaler acting in compliance with the minimum requirements of SHB 1675 and WAC 246-945-090 through 246-945-093.

Small manufacturers and wholesalers that choose to distribute prescriptions to home dialysis patients must secure and utilize a pharmacist consultant. Distributing manufacturers and wholesalers must also develop and implement protocol for shipments, deliveries, and error documentation. Finally, these manufacturers and wholesalers must also provide quality assurance measures to protect medications from diversion or tampering in line with their own security policies and procedures.

At the August 2024 business meeting, the commission considered feedback from interested parties at a rule hearing on this topic and voted to approve filing a supplemental CR-102. The commission's approved revisions to the proposed rule language are applicable to small-scale manufacturers and wholesalers; however, no additional costs are anticipated as their intent is to provide clarification. The commission determined that the proposed rule needed to be further amended and required two amendments to WAC 246-945-090. The first amendment is to reinstate the term "may." The second amendment is to add a list of approved dialysis devices that manufacturers and wholesalers may sell, deliver, possess, and dispense to home dialysis patients. Amendments are also needed to WAC 246-945-091, 246-945-092, and 246-945-093 in association with the WAC 246-945-090 amendments.

SBEIS Table 1 identifies and summarizes of which businesses are required to comply with the proposed rule using the North American Industry Classification System (NAICS).

SBEIS Table 1. Summary of Businesses Required to comply to the Proposed Rule

NAICS Code (4, 5 or 6 digit)	NAICS Business Description	Number of businesses in Washington State	Minor Cost Threshold
424210	Drugs and Druggists' Sundries Merchant Wholesalers	121	\$10,305.83

The following is an Analysis of probable costs of businesses in the industry to comply to the proposed rule and includes the cost of equipment, supplies, labor, professional services, and administrative costs. The analysis considers if compliance with the proposed rule will cause businesses in the industry to lose sales or revenue.

### WAC 246-945-090 Home dialysis programs, manufacturers, and wholesalers—Legend drugs.

**Description:** The current rule allows a medicare-approved dialysis center or facility operating a medicare-approved home dialysis program to sell, deliver, or dispense directly to its home dialysis patients specified legend drugs. The proposed rule allows manufacturers and wholesalers to sell, deliver, possess, or dispense approved legend drugs used in home dialysis programs directly to patients, provided that the treatment was prescribed by a practitioner acting within the scope of their practice. Manufacturers and wholesalers that ship and deliver approved legend drugs and dialysis devices directly to patients will not be required to retain a pharmacy or nonresident pharmacy license to do so.

The language proposed as part of the supplemental clarifies that that a medicare-approved dialysis center or facility operating a medicare-approved home dialysis program, a manufacturer or a wholesaler may sell, deliver, or dispense directly to its home dialysis patients specified legend drugs. The proposed rule also list the dialysis devices that may be sold, delivered or dispensed directly to patients.

**Cost(s):** \$187 total probable cost per participating manufacturer or wholesaler for ninety minutes of staff time to prepare and deliver training to employees. This probable cost assumes an average health service manufacturer or wholesaler employing 200 employees has a shipping and receiving team of 10 production workers and 1 manager. Commission staff estimate that the training will require 60 minutes of the manager's time (\$62/hour) to prepare and deliver the training on patient home deliveries and 30 minutes of each production worker's time (\$25/hour) to receive the training. There is no anticipated costs associated with adding the list of devices that may be sold, delivered or dispensed. The list provides clarification but does not require the devices to be sold.

# WAC 246-945-091 Home dialysis programs, manufacturers, and wholesalers—Pharmacist consultant.

**Description:** The current rule requires home dialysis program involved in the distribution of legend drugs to have an agreement with a pharmacist with provides consultation as necessary. The proposed rule requires manufacturers and wholesalers that ship and deliver approved legend drugs and dialysis devices to patients to also establish an agreement with a pharmacist for consultation on an as needed basis. This is not currently required by statute. The shipment and delivery content of the agreement may be in addition to or stand alone to an existing pharmacist consultant agreement. The language proposed as part of the supplemental clarifies that the agreement shall include advice on both the drug and device shipment and delivery process.

**Cost(s):** It is estimated that there will be \$426 ongoing probable cost to manufacturers and wholesalers for 6 hours of a pharmacist's time (\$71/hour) for consultation annually. Commission staff estimate that approximately 6 hours, one hour every other month of a year, will be necessary for a wholesaler or manufacturer to discuss shipment and delivery protocol with a pharmacist in order to deliver and dispense dialysis devices and approved legend drugs safely to patients. The clarification provided as part of the supplemental is not anticipated to add any additional time or costs.

### WAC 246-945-092 Home dialysis programs, manufacturers, and wholesalers—Records.

**Description:** The current rule outlines what is required to be on the record of shipment and attached to the prescriber's order. The proposed rule requires manufacturers and wholesalers that ship and deliver approved legend drugs and dialysis devices directly to home patients to attach a record of shipment to each practitioner's order that includes additional information than currently required of manufacturers and wholesalers with pharmacy or nonresident pharmacy licensees by WAC 246-945-016. The record of shipment needs to include the name of the patient, strengths and quantities of drugs, manufacturers' names, date of shipment, names of people who selected, assembled and packaged the shipment, and the name of the pharmacist or designated person responsible for the shipment. The language proposed as part of the supplemental clarifies that the record shall also include information on devices if applicable.

<sup>&</sup>lt;sup>1</sup> What is compliance training, and why is it important? What is compliance training, and why is it important? (powerdms.com). (Accessed March 26, 2024)

<sup>&</sup>lt;sup>2</sup> 43.5 percent of manufacturing workers in establishments with 250 or more workers in March 2018. <u>43.5 percent of manufacturing workers in establishments with 250 or more workers in March 2018 : The Economics Daily: U.S. Bureau of Labor Statistics (bls.gov)</u>. (Accessed March 25, 2024)

<sup>&</sup>lt;sup>3</sup> The Ideal Manager to Employee Ratio: How Many Managers Do You Need? <u>The Ideal Manager to Employee Ratio: How Many Managers Do You Need? - Don Romans</u> (Accessed March 25, 2024)

<sup>&</sup>lt;sup>4</sup> Occupational Employment and Wages, May 2023. <u>Transportation, Storage, and Distribution Managers (bls.gov)</u> (Accessed March 25, 2024)

<sup>&</sup>lt;sup>5</sup> Occupational Employment and Wages, May 2023. Production Workers, All Other (bls.gov) (Accessed March 25, 2024)

<sup>&</sup>lt;sup>6</sup> Occupational Employment and Wages, May 2023 - 29-1051 Pharmacists. Pharmacists (bls.gov) (Accessed March 25, 2024)

**Cost(s):** It is estimated that a manufacturer or wholesaler will incur \$300 one-time probable cost for a printer and \$304 ongoing probable cost for toner and paper for printing records of shipment. These probable costs are based on an assumption of 10,000 shipments annually requiring printed records. The clarification provided as part of the supplemental is not anticipated to add any additional time or costs.

### WAC 246-945-093 Home dialysis programs, manufacturers, and wholesalers—Quality assurance.

**Description:** The current rule requires home dialysis programs involved in the distribution of legend drugs to develop a quality assurance program for drug distribution and to maintain records of drug distribution errors and other problems, including loss due to damage or theft. The proposed rule will require manufacturers and wholesalers who ship and deliver approved legend drugs and dialysis devices directly to patients to develop quality assurance programs for shipment and delivery and maintain a record of shipment and delivery errors. The current statute does not require manufacturers and wholesalers to maintain quality assurance programs for shipment and delivery nor records of shipment and delivery errors. The shipment and delivery quality assurance plan and error record may be supplemental to an existing quality assurance program. The language proposed as part of the supplemental clarifies that the quality assurance program requirements apply to devices if applicable.

Cost(s): It is estimated that a manufacturer or wholesaler will incur \$328 one-time probable cost for three hours of a production manager's time (\$62/hour) and two hours of a pharmacist consultant's time (\$71/hour) to fulfill the quality assurance program requirements. 10,11 In addition, there are \$124 of estimated ongoing probable cost for two hours of a production manager's time (\$62/hour) to maintain a record of shipment and delivery errors. The clarification provided as part of the supplemental is not anticipated to add any additional time or costs.

## Summary of all Cost(s)

#### SBEIS Table 2. Summary of probable cost(s)

WAC Section and Title	Probable Cost(s)
WAC 246-945-090 Home dialysis programs, manufacturers, and wholesalers—Legend drugs	\$187 one-time for employee training
WAC 246-945-091 Home dialysis programs, manufacturers, and wholesalers—Pharmacist consultant	\$426 ongoing for pharmacist consultations
WAC 246-945-092 Home dialysis programs, manufacturers, and wholesalers—Records	\$300 one-time for a printer for records of shipment \$304 ongoing for toner and paper for records of shipment
WAC 246-945-093 Home dialysis programs, manufacturers, and wholesalers—Quality assurance	\$328 one-time for quality assurance program development \$124 ongoing for quality assurance program improvement
Total	\$1,669.00

Analysis on if the proposed rule may impose more than minor costs for businesses in the industry. Includes a summary of how the costs were calculated.

The costs of the proposed rule (\$1,669) are <u>less than</u> the minor cost threshold (\$10,305.83).

#### Summary of how the costs were calculated

The probable costs were calculated for participating manufacturers and wholesalers to comply with the proposed rule. Probable costs affiliated with compliance primarily pertain to staff time. Average staff wages in Washington state were sourced from data produced by the U.S. Bureau of Labor and Statistics. Additional resources were used to estimate employee quantities. Commission staff, including a Pharmacist Consultant, determined the estimated time requirements.

$\square$ Yes Calculations show the rule proposal likely imposes more-than-minor cost to businesses and a small busin	ess
economic impact statement is required. Insert the required small business economic impact statement here:	

<sup>&</sup>lt;sup>7</sup> Staples. Staples<sup>®</sup> Official Online Store (Accessed April 22, 2024)

<sup>8</sup> National ESRD Census Data. National ESRD Census Data (esrdnetworks.org) (Accessed April 9, 2024)

<sup>&</sup>lt;sup>9</sup> Manufacturing and Quality Assurance: A Comprehensive Guide. <u>Manufaturing Quality Assurance: A Comprehensive Guide</u> (<u>cashflowinventory.com</u>) (Accessed March 25, 2024)

<sup>&</sup>lt;sup>10</sup> See footnote 4

<sup>&</sup>lt;sup>11</sup> See footnote 8

The public may obtain a copy of the small business economic impact statement or the detailed cost calculations by contacting:

Name Julia Katz

Address PO Box 47852, Olympia, WA 98504-7852

Phone 360-236-4946 Fax 360-236-2260

TTY 711

Email PharmacyRules@doh.wa.gov

Other None

Date: November 22, 2024

Name: Hawkins DeFrance, PharmD

Title: Pharmacy Quality Assurance Commission Chair

Signature:

AMENDATORY SECTION (Amending WSR 20-12-072, filed 6/1/20, effective 7/1/20)

WAC 246-945-090 Home dialysis programs, manufacturers, and wholesalers—Legend drugs and dialysis devices. Pursuant to RCW 18.64.257 and 69.41.032, a medicare-approved dialysis center ((er)), a facility operating a medicare-approved home dialysis program ((may)), a manufacturer, or a wholesaler may sell, deliver, possess, or dispense directly to its home dialysis patients, in case((er)) or full shelf ((er)) lots, and if prescribed by a ((er)) practitioner, the following:

(1) Legend drugs:

 $\overline{(((1)))}$ ) (a) Sterile heparin, 1000 u/mL, in vials;

 $((\frac{2}{2}))$  (b) Sterile potassium chloride, 2 mEq/mL, for injection;

(((3))) Commercially available dialysate; and

- ((4)))  $\overline{(d)}$  Sterile sodium chloride, 0.9%, for injection in containers of not less than 150 mL.
  - (2) Dialysis devices:
- (a) Class II medical devices that are manufactured and marketed in compliance with the Federal Food, Drug, and Cosmetic Act and indicated for acute and chronic dialysis therapy in the home; and
  - (b) Related supplies and accessories of the dialysis device.

AMENDATORY SECTION (Amending WSR 20-12-072, filed 6/1/20, effective 7/1/20)

WAC 246-945-091 Home dialysis programs, manufacturers, and wholesalers—Pharmacist consultant. ((Home dialysis programs involved in the distribution of legend drugs as)) A medicare-approved dialysis center, a facility operating a medicare-approved home dialysis program, a manufacturer, or a wholesaler who sells, delivers, possesses, or dispenses dialysis devices and legend drugs directly to its home dialysis patients permitted by RCW 18.64.257 and 69.41.032(( $\tau$ )) shall have an agreement with a pharmacist which provides for consultation as necessary. This agreement shall include advice on the drug ((distribution)) and device shipment and delivery process to home dialysis patients and on the location used for storage and ((distribution)) shipment of the authorized drugs and devices, which shall be reasonably separated from other activities and shall be secure.

AMENDATORY SECTION (Amending WSR 20-12-072, filed 6/1/20, effective 7/1/20)

WAC 246-945-092 Home dialysis programs, manufacturers, and wholesalers—Records. (1) A medicare-approved dialysis center, a facility operating a medicare-approved home dialysis program, a manufacturer, or a wholesaler who sells, delivers, possesses, or dispenses dialysis devices and legend drugs directly to its home dialysis patients permitted by RCW 18.64.257 and 69.41.032 shall attach a record

[ 1 ] OTS-5459.2

of shipment ((shall be attached)) to the ((prescriber's)) practitioner's order ((and)). The record of shipment shall include:

- (a) The name of the patient;
- (b) Strengths and quantities of drugs, if applicable;
- (c) <u>Device name</u>, if applicable;
- (d) The name of the drug manufacturer((s' names)), if applicable;
- ((<del>(d)</del>)) <u>(e) The name of the device manufacturer, if applicable;</u>
- (f) Date of shipment;
- $((\frac{(e)}{(e)}))$  (g) Names of persons who selected, assembled and packaged for shipment; and
- $((\frac{f}{f}))$  The name of the pharmacist or designated individual responsible for the  $((\frac{distribution}{f}))$  shipment.
- (2) Prescription <u>records</u>, and drug ((<del>distribution</del>)) <u>and device</u> <u>shipment</u> records shall be maintained in accordance with WAC 246-945-020.

AMENDATORY SECTION (Amending WSR 20-12-072, filed 6/1/20, effective 7/1/20)

WAC 246-945-093 Home dialysis programs, manufacturers, and wholesalers—Quality assurance. ((Home dialysis programs involved in the distribution of legend drugs as)) A medicare-approved dialysis center, a facility operating a medicare-approved home dialysis program, a manufacturer, or a wholesaler who sells, delivers, possesses, or dispenses dialysis devices and legend drugs directly to its home dialysis patients permitted by RCW 18.64.257 and 69.41.032(( $\tau$ )) shall develop a quality assurance program for drug ((distribution)) and device shipment and delivery, and shall maintain records of drug ((distribution)) and device shipment and delivery errors and other problems, including loss due to damage or theft.

[ 2 ]

OTS-5459.2