S.B. NO. ⁴⁷³ S.D. 1

A BILL FOR AN ACT

RELATING TO THE PRACTICE OF PHARMACY.

BE IT ENACTED BY THE LEGISLATURE OF THE STATE OF HAWAII:

1	SECTION 1. Chapter 461, Hawaii Revised Statutes, is
2	amended by adding a new section to be appropriately designated
3	and to read as follows:
4	"§461- Distribution of dialysate drugs and devices. (a)
5	The license, registration, and permit requirements of this
6	chapter shall not apply to a drug manufacturer, wholesale
7	prescription drug distributor, or third-party logistics
8	provider, to the extent the manufacturer, wholesale distributor,
9	or third-party logistics provider is engaged in the distribution
10	of dialysate drugs or devices necessary to perform home dialysis
11	on patients with end-stage renal disease; provided that the
12	following criteria are met:
13	(1) The dialysate drugs or devices are approved by the
14	United States Food and Drug Administration, as
15	required by federal law;
16	(2) The dialysate drugs or devices are lawfully held by a

manufacturer or a manufacturer's agent that is



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1		properly registered with the board as a manufacturer
2		or wholesale distributor;
3	(3)	The dialysate drugs or devices are held and delivered
4		in their original, sealed, and labeled packaging from
5		the manufacturing facility;
6	(4)	The dialysate drugs or devices are delivered only by
7		the manufacturer or the manufacturer's agent and only
8		upon receipt of a physician's order; and
9	(5)	The manufacturer or the manufacturer's agent delivers
10		the dialysate drugs or devices directly to a:
11		(A) Patient with end stage renal disease, or the
12		patient's designee, for the patient's self-
13		administration of dialysis therapy; or
14		(B) Health care provider or institution for
15		administration or delivery of dialysis therapy to
16		a patient with end stage renal disease.
17	(b)	For the purposes of this section:
18	"Manı	ufacturer" shall have the same meaning as in section
19	328-112.	
20	<u>"Thi</u>	rd-party logistics provider" means an entity that
21	provides o	or coordinates warehousing or other logistics services

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on behalf of a pharmaceutical manufacturer, wholesale
distributor, or dispenser of a product.
"Wholesale distributor" shall have the same meaning as in
section 328-112."
SECTION 2. New statutory material is underscored.
SECTION 3. This Act shall take effect on December 31,
2050.

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Report Title:

Pharmacy; Dialysis Drugs and Supplies; Manufacturers; Wholesalers; Third-Party Logistics Providers; Exemption

Description:

Exempts drug manufacturers, wholesale prescription drug distributors, and third-party logistics providers of home dialysis drugs, supplies, and devices from the license, registration, and permit requirements for pharmacies; provided that certain conditions are met. Effective 12/31/2050. (SD1)

The summary description of legislation appearing on this page is for informational purposes only and is not legislation or evidence of legislative intent.

