THE SENATE THIRTIETH LEGISLATURE, 2019 STATE OF HAWAII

541 S.D. 1 S.B. NO.

A BILL FOR AN ACT

RELATING TO MEDICAL CANNABIS PRODUCTS.

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

1 SECTION 1. The legislature finds that the list of medical 2 cannabis products that may be manufactured and distributed 3 pursuant to section 329D-10, Hawaii Revised Statutes, omits viable products. The legislature notes that the medical 4 5 cannabis legislative oversight working group, established 6 pursuant to Act 230, Session Laws of Hawaii 2016, recommends 7 amending section 329D-10 to refer to "transdermal devices", 8 instead of "transdermal patches", as excluding non-patch devices 9 that deliver through the dermis was unintentional.

10 The purpose of this Act is to further ensure access to 11 medical cannabis for qualifying patients, by updating references 12 in the medical cannabis dispensary laws from transdermal patches 13 to transdermal devices and simplifying the list of manufactured 14 cannabis products that may be manufactured and distributed by 15 dispensaries to section 329-10.

SB541 HD1 HMS 2019-3115

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1	SECT:	ION 2. Section 329D-1, Hawaii Revised Statutes, is		
2	amended by amending the definition of "manufactured cannabis			
3	product" to read as follows:			
4	""Manufactured cannabis product" means any [capsule,			
5	lozenge, oil or oil extract, tincture, ointment or skin lotion,			
6	pill, transdermal patch, or pre-filled and sealed container used			
7	to aerosolize and deliver cannabis orally, such as an inhaler or			
8	nebulizer, that has been manufactured using cannabis, or any			
9	other products] product as specified by the department pursuant			
10	to section [329D-10(a)(9).] <u>329D-10.</u> "			
11	SECTION 3. Section 329D-10, Hawaii Revised Statutes, is			
12	amended by amending subsection (a) to read as follows:			
13	"(a)	The types of medical cannabis products that may be		
14	manufactured and distributed pursuant to this chapter shall be			
15	limited to:			
16	(1)	Capsules;		
17	(2)	Lozenges;		
18	(3)	Pills;		
19	(4)	Oils and oil extracts;		
20	(5)	Tinctures;		
21	(6)	Ointments and skin lotions;		



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1 (7) Transdermal [patches;] devices as approved by the 2 department;

3 Pre-filled and sealed containers used to aerosolize (8) and deliver cannabis orally, such as with an inhaler 4 5 or nebulizer; provided that containers need not be 6 manufactured by the licensed dispensary but shall be 7 filled with cannabis, cannabis oils, or cannabis 8 extracts manufactured by the licensed dispensary; 9 shall not contain nicotine, tobacco-related products, or any other non-cannabis derived products; and shall 10 11 be designed to be used with devices used to provide 12 safe pulmonary administration of manufactured cannabis 13 products;

- 14 (9) Devices that provide safe pulmonary administration;15 provided that:
- 16 (A) The heating element of the device, if any, is
 17 made of inert materials such as glass, ceramic,
 18 or stainless steel, and not of plastic or rubber;
 19 (B) The device is distributed solely for use with
 20 single-use, pre-filled, tamper-resistant, sealed



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1		containers that do not contain nicotine or other
2		tobacco products;
3	(C)	The device is used to aerosolize and deliver
4		cannabis by inhalation, such as an inhaler,
5		medical-grade nebulizer, or other similar medical
6		grade volitization device;
7	(D)	There is a temperature control on the device that
8		is regulated to prevent the combustion of
9		cannabis oil; and
10	(E)	The device need not be manufactured by the
11		licensed dispensary; and
12	(10) Othe	r products as specified by the department."
13	SECTION 4	. Statutory material to be repealed is bracketed
14	and stricken.	New statutory material is underscored.
15	SECTION 5	. This Act shall take effect on July 1, 2050.





Report Title:

Medical Cannabis; Manufactured Cannabis Products; Transdermal Devices

Description:

Simplifies the list of manufactured cannabis products that may be manufactured and distributed by dispensaries. Updates transdermal patches to department-approved transdermal devices in section 329D-10, Hawaii Revised Statutes, thereby including non-patch devices that deliver through the dermis. (SB541 HD1)

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

