
A BILL FOR AN ACT

RELATING TO HEALTH.

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

SECTION 1. Chapter 329, Hawaii Revised Statutes, is amended by adding a new section to be appropriately designated and to read as follows:

"§329- Food and Drug Administration-approved drugs; cannabidiol. (a) Upon approval by the federal Food and Drug Administration of one or more prescription drugs containing cannabidiol, the following activities shall be lawful in the State:

(1) The clinically appropriate prescription for a patient of a Food and Drug Administration-approved prescription drug containing cannabidiol by a health care provider licensed to prescribe medications in this State and acting within the health care provider's authorized scope of practice;

(2) The dispensing, pursuant to a valid prescription, of a Food and Drug Administration-approved prescription drug containing cannabidiol to a patient or a



patient's authorized representative by a pharmacist or
another health care provider licensed to dispense
medications in this State and acting within the health
care provider's authorized scope of practice;

(3) The possession and transportation of a Food and Drug
Administration-approved prescription drug containing
cannabidiol by a patient to whom a valid prescription
was issued or by the patient's authorized
representative;

(4) The possession and transportation of a Food and Drug
Administration-approved prescription drug containing
cannabidiol by a licensed pharmacy or wholesaler to
facilitate the appropriate dispensing and use of the
drug; and

(5) The use of a Food and Drug Administration-approved
prescription drug containing cannabidiol by a patient
to whom a valid prescription was issued; provided that
the patient uses the drug only for legitimate medical
purposes in conformity with instructions from the
prescriber and dispenser.



1 (b) Upon approval by the Food and Drug Administration of
2 one or more prescription drugs containing cannabidiol, the
3 department shall amend its rules to conform to the requirements
4 of subsection (a).

5 (c) Nothing in this section shall be construed to amend,
6 alter, or otherwise restrict access to medical cannabis,
7 recreational marijuana, or both, as authorized under state law."

8 SECTION 2. New statutory material is underscored.

9 SECTION 3. This Act shall take effect upon its approval.

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Report Title:

Relating to Health. Cannabidiol; Prescription Drugs; Food and Drug Administration

Description:

Specifies certain activities that shall become lawful, upon approval by the federal Food and Drug Administration of one or more prescription drugs containing cannabidiol.

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

