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A BILL FOR AN ACT

RELATING TO MENTAL HEALTH.

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

1 SECTION 1. The legislature finds that mental health 2 conditions are treated in various ways, depending on the 3 condition, and can include medication, therapy, and psychosocial 4 services. Congress, through the Breakthrough Therapies Act, and 5 the Food and Drug Administration has indicated that 3,4methylenedioxymethamphetamine, commonly known as MDMA, and 6 7 psilocybin will be rescheduled to enable therapeutic use. MDMA 8 and psilocybin have already been granted the Food and Drug 9 Administration's breakthrough therapy designation to fast-track 10 research and potential approval given efficacy in treating 11 treatment-resistant depression and post-traumatic stress 12 disorder. These treatments, while effective for certain 13 conditions and patients, do not treat all mental health 14 conditions. However, research supports the effectiveness of 15 natural and alternative medicines and therapies, such as the use 16 of MDMA, psilocybin, and other therapies, as a safe and 17 effective way to potentially treat depression, post-traumatic



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stress disorder, addiction, end-of-life psychological distress,
 and other afflictions.

The legislature further finds that the department of health should be empowered to review relevant laws, regulations, and studies each time a breakthrough therapy designation is issued to review any new treatment intended for mental health or substance abuse to prepare the State for the treatment's eventual approval by the federal Food and Drug Administration.

9 The purpose of this Act is to authorize the director of
10 health to establish a temporary breakthrough therapy
11 designations advisory council within three months of a
12 breakthrough therapy designation approval by the Food and Drug
13 Administration.

SECTION 2. Chapter 321, Hawaii Revised Statutes, is amended by adding a new section to part I to be appropriately designated and to read as follows:

17 "<u>§321-</u> Temporary breakthrough therapy designation 18 advisory council. (a) The director of health may establish a 19 temporary breakthrough therapy designation advisory council to 20 assess a breakthrough therapy designation for a mental health or 21 substance abuse treatment within three months of a breakthrough



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1	therapy d	esignation approval by the United States Food and Drug
2	Administr	ation. The advisory council is established within the
3	departmen	t of health for administrative purposes only.
4	(b)	The advisory council shall consist of the following
5	members o	r their designees:
6	(1)	The director of health, who shall serve as the
7		chairperson of the advisory council;
8	(2)	The attorney general;
9	(3)	The director of law enforcement;
10	(4)	The chairpersons of the standing committees within the
11		senate and house of representatives with primary
12		jurisdiction over health;
13	(5)	A physician who is duly licensed pursuant to chapter
14		453 or an advanced practice registered nurse who is
15		authorized to prescribe psychotropic medication and is
16		duly licensed pursuant to chapter 457; and
17	(6)	Other members as recommended by the director of
18		health, president of the senate, or speaker of the
19		house of representatives, who represent relevant
20		community, advocacy, or stakeholder interests.

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1	(c)	Members shall serve without compensation, but may be
2	reimburse	d for necessary expenses, including reasonable travel
3	expenses,	incurred in the performance of their duties.
4	(d)	The advisory council shall:
5	(1)	Examine federal and state laws, regulations,
6		administrative rules, and community practices
7		regarding the treatment of mental health or substance
8		abuse conditions for which the breakthrough therapy
9		designation applies;
10	(2)	Examine available clinical and scientific studies,
11		research, and other information relating to the safety
12		and efficacy of methods to treat mental health or
13		substance abuse conditions for which the breakthrough
14		therapy designation applies;
15	(3)	Examine requirements, specifications, and guidelines
16		for a health care professional to prescribe and
17		provide various treatments for patients who may
18		benefit; and
19	(4)	Submit a report of its findings and recommendations,
20		including any proposed legislation, to the legislature

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1	no later than one year after the advisory council is
2	convened.
3	(e) The advisory council may convene as necessary but
4	shall terminate upon the withdrawal of the breakthrough therapy
5	designation or final approval by the United States Food and Drug
6	Administration.
7	(f) As used in this section, "breakthrough therapy
8	designation" means a designation by the United States Food and
9	Drug Administration, pursuant to the Food and Drug
10	Administration Safety and Innovation Act (P.L. 112-144)."
11	SECTION 3. New statutory material is underscored.
12	SECTION 4. This Act shall take effect on June 30, 3000.



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Report Title:

Temporary Breakthrough Therapy Designation Advisory Council; DOH; Mental Health

Description:

Authorizes the director of health to establish a temporary breakthrough therapy designation advisory council within three months of a breakthrough therapy designation approval by the United States Food and Drug Administration. Effective 6/30/3000. (HD2)

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

