STATE OF VERMONT  
CANNABIS CONTROL BOARD

EMERGENCY RULE: SYNTHETIC and HEMP-DERIVED CANNABINOIDS

The Board shall have the authority to regulate synthetic cannabinoids and hemp-derived cannabinoids, including delta-8 and delta-10 tetrahydrocannabinol.

7 V.S.A. § 862a.

Emergency Rule 1:  Prohibition

The production, manufacture, marketing, transfer, and sale of hemp-derived intoxicating cannabinoids and synthetic cannabinoids are hereby prohibited, except as set out in Emergency Rule 3. Prohibited cannabinoids include:

(a) All isomers, variants, analogs, and mimetics of delta-9 tetrahydrocannabinol, including delta-8 and delta-10 tetrahydrocannabinol, created by chemical manipulation of any part or derivative of the plant Cannabis sativa L., regardless of the delta-9 tetrahydrocannabinol concentration level of the source plant or plants; and

(b) delta-9 tetrahydrocannabinol that has been chemically or mechanically concentrated or otherwise derived from hemp and then sprayed, infused, or otherwise artificially introduced onto or into any product, including hemp or hemp products, so as to impart intoxicating properties mimicking those of cannabis and cannabis products.

Emergency Rule 2:  Presumptions

A consumable product that is not cannabis or a cannabis product is presumptively prohibited regardless of the delta-9 tetrahydrocannabinol concentration of any plant from which the product is sourced, if the product, in the form offered to consumers:

(a) contains total tetrahydrocannabinol in a concentration exceeding 0.3 percent on a dry weight basis; or

(b) contains more than 1.5 mg tetrahydrocannabinol per serving, where “serving” is the amount reasonably ingested by a typical consumer in a single instance; or

(c) contains more than 10 mg total tetrahydrocannabinol per package, unless the ratio of cannabidiol to tetrahydrocannabinol is at least 20:1; or

(d) has the dominant market appeal of mimicking the intoxicating effects of tetrahydrocannabinol.

Emergency Rule 3:  Exceptions

Rule 1 shall not apply to:

(a) a product duly evaluated, registered, and regulated by the Board as a cannabis product;

(b) an otherwise-prohibited cannabinoid-containing product that has been specifically authorized by the Board for sale at a licensed medical dispensary based upon a finding, pursuant to 7 V.S.A. § 971(b)(6), that the product is appropriate for use by a patient; or

(c) manufactured pharmaceutical drugs approved by the United States Food & Drug Administration for therapeutic use upon the prescription of a medical provider, to include Epidiolex, Marinol, Syndros, Cesamet, and Sativex.

Effective: April 24, 2023