

General Assembly

February Session, 2024

## Raised Bill No. 5235

LCO No. **1643** 

Referred to Committee on GENERAL LAW

Introduced by: (GL)

## AN ACT CONCERNING THE DEPARTMENT OF CONSUMER PROTECTION'S RECOMMENDATIONS REGARDING CANNABIS REGULATION.

Be it enacted by the Senate and House of Representatives in General Assembly convened:

Section 1. Section 21a-240 of the 2024 supplement to the general
 statutes is repealed and the following is substituted in lieu thereof
 (*Effective from passage*):

4 The following words and phrases, as used in this chapter, shall have 5 the following meanings, unless the context otherwise requires:

6 (1) "Abuse of drugs" means the use of controlled substances solely for 7 their stimulant, depressant or hallucinogenic effect upon the higher 8 functions of the central nervous system and not as a therapeutic agent 9 prescribed in the course of medical treatment or in a program of 10 research operated under the direction of a physician or pharmacologist.

11 (2) "Administer" means the direct application of a controlled 12 substance, whether by injection, inhalation, ingestion or any other 13 means, to the body of a patient or research subject by: (A) A practitioner, or, in the practitioner's presence, by the practitioner's authorized agent; [, or] (B) the patient or research subject at the direction and in the presence of the practitioner; [,] or (C) a nurse or intern under the direction and supervision of a practitioner.

(3) "Agent" means an authorized person who acts on behalf of or at
the direction of a manufacturer, distributor, dispenser or prescribing
practitioner, but does not include a common or contract carrier, public
warehouseman [,] or employee of the carrier or warehouseman.

(4) "Amphetamine-type substances" include amphetamine, optical
isomers thereof, salts of amphetamine and its isomers, and chemical
compounds which are similar thereto in chemical structure or which are
similar thereto in physiological effect, and which show a like potential
for abuse, which are controlled substances under this chapter unless
modified.

(5) "Barbiturate-type drugs" include barbituric acid and its salts,
derivatives thereof and chemical compounds which are similar thereto
in chemical structure or [which are similar thereto in] physiological
effect, and which show a like potential for abuse, which are controlled
substances under this chapter unless modified.

(6) "Bureau" means the Bureau of Narcotics and Dangerous Drugs,United States Department of Justice, or its successor agency.

35 (7) "Cannabis-type substances" include all parts of any plant, or 36 species of the genus cannabis or any infra specific taxon thereof whether 37 growing or not; the seeds thereof; the resin extracted from any part of 38 such a plant; and every compound, manufacture, salt, derivative, 39 mixture or preparation of such plant, its seeds or resin; but shall not 40 include the mature stalks of such plant, fiber produced from such stalks, 41 oil or cake made from the seeds of such plant, any other compound, 42 manufacture, salt, derivative, mixture or preparation of such mature 43 stalks, except the resin extracted therefrom, fiber, oil or cake, the 44 sterilized seed of such plant which is incapable of germination, or hemp, 45 as defined in 7 USC 16390, as amended from time to time. Included are

46 cannabinon, cannabinol, cannabidiol and chemical compounds which
47 are similar to cannabinon, cannabinol or cannabidiol in chemical
48 structure or which are similar thereto in physiological effect, and which
49 show a like potential for abuse, which are controlled substances under
50 this chapter unless derived from hemp, as defined in section 22-61*l*.

(8) "Controlled drugs" are those drugs which contain any quantity of 51 52 a substance which has been designated as subject to the federal 53 Controlled Substances Act, or which has been designated as a 54 depressant or stimulant drug pursuant to federal food and drug laws, or which has been designated by the Commissioner of Consumer 55 56 Protection pursuant to section 21a-243, as amended by this act, as 57 having a stimulant, depressant or hallucinogenic effect upon the higher 58 functions of the central nervous system and as having a tendency to 59 promote abuse or psychological or physiological dependence, or both. Such controlled drugs are classifiable as amphetamine-type, 60 cannabis-type, 61 barbiturate-type, cocaine-type, hallucinogenic, 62 morphine-type and other stimulant and depressant drugs. Specifically 63 excluded from controlled drugs and controlled substances are alcohol, 64 nicotine and caffeine.

(9) "Controlled substance" means a drug, substance [,] or immediate
precursor in schedules I to V, inclusive, of the Connecticut controlled
substance scheduling regulations adopted pursuant to section 21a-243,
<u>as amended by this act</u>.

(10) "Counterfeit substance" means a controlled substance which, or the container or labeling of which, without authorization, bears the trademark, trade name or other identifying mark, imprint, number or device, or any likeness thereof, of a manufacturer, distributor or dispenser other than the person who in fact manufactured, distributed or dispensed the substance.

(11) "Deliver or delivery" means the actual, constructive or attempted
transfer from one person to another of a controlled substance, whether
or not there is an agency relationship.

(12) "Dentist" means a person authorized by law to practice dentistryin this state.

80 (13) "Dispense" means to deliver a controlled substance to an ultimate 81 user or research subject by or pursuant to the lawful order of a 82 practitioner, including the prescribing, administering, packaging, 83 labeling or compounding necessary to prepare the substance for the 84 delivery.

85 (14) "Dispenser" means a practitioner who dispenses.

86 (15) "Distribute" means to deliver other than by administering or87 dispensing a controlled substance.

(16) "Distributor" means a person who distributes and includes a
wholesaler who is a person supplying or distributing controlled drugs
which the person personally has not produced or prepared to hospitals,
clinics, practitioners, pharmacies, other wholesalers, manufacturers and
federal, state and municipal agencies.

93 (17) "Drug" means: (A) [substances] Substances recognized as drugs 94 in the official United States Pharmacopoeia, official Homeopathic 95 Pharmacopoeia of the United States, or official National Formulary, or 96 any supplement to any of them; (B) substances intended for use in the 97 diagnosis, cure, mitigation, treatment or prevention of disease in man 98 or animals; (C) substances, other than food, intended to affect the 99 structure or any function of the body of man or animals; and (D) 100 substances intended for use as a component of any article specified in 101 subparagraph (A), (B) or (C) of this subdivision. [It] "Drug" does not 102 include devices or their components, parts or accessories.

(18) "Drug dependence" means a psychoactive substance dependence
on drugs as that condition is defined in the most recent edition of the
"Diagnostic and Statistical Manual of Mental Disorders" of the American
Psychiatric Association.

107 (19) "Drug-dependent person" means a person who has a

psychoactive substance dependence on drugs as that condition is
defined in the most recent edition of the "Diagnostic and Statistical
Manual of Mental Disorders" of the American Psychiatric Association.

111 (20) (A) "Drug paraphernalia" means equipment, products and 112 materials of any kind that are used, intended for use or designed for use planting, 113 cultivating, in propagating, growing, harvesting, 114 manufacturing, compounding, converting, producing, processing, 115 preparing, testing, analyzing, packaging, repackaging, storing, 116 containing or concealing, or ingesting, inhaling or otherwise 117 introducing into the human body, any controlled substance contrary to the provisions of this chapter, including, but not limited to: (i) Kits 118 119 intended for use or designed for use in planting, propagating, 120 cultivating, growing or harvesting of any species of plant that is a 121 controlled substance or from which a controlled substance can be 122 derived; (ii) kits used, intended for use or designed for use in 123 manufacturing, compounding, converting, producing, processing or 124 preparing controlled substances; (iii) isomerization devices used or 125 intended for use in increasing the potency of any species of plant that is 126 a controlled substance; (iv) testing equipment used, intended for use or 127 designed for use in identifying or analyzing the strength, effectiveness 128 or purity of controlled substances; (v) dilutents and adulterants, 129 including, but not limited to, quinine hydrochloride, mannitol, mannite, 130 dextrose and lactose used, intended for use or designed for use in 131 cutting controlled substances; (vi) separation gins and sifters used, 132 intended for use or designed for use in removing twigs and seeds from, 133 or in otherwise cleaning or refining, marijuana; (vii) capsules and other 134 containers used, intended for use or designed for use in packaging small 135 quantities of controlled substances; (viii) containers and other objects 136 used, intended for use or designed for use in storing or concealing 137 controlled substances; and (ix) objects used, intended for use or 138 designed for use in ingesting, inhaling, or otherwise introducing 139 marijuana, cocaine, hashish [,] or hashish oil into the human body, 140 including, but not limited to, wooden, acrylic, glass, stone, plastic or 141 ceramic pipes with screens, permanent screens, hashish heads or

142 punctured metal bowls; water pipes; carburetion tubes and devices; 143 smoking and carburetion masks; roach clips; miniature cocaine spoons 144 and cocaine vials; chamber pipes; carburetor pipes; electric pipes; airdriven pipes; chillums; bongs; ice pipes and chillers. "Drug 145 146 paraphernalia" does not include a product used by a manufacturer 147 licensed pursuant to this chapter for the activities permitted under the 148 license or by an individual to test any substance prior to injection, 149 inhalation or ingestion of the substance to prevent accidental overdose 150 by injection, inhalation or ingestion of the substance, provided the 151 licensed manufacturer or individual is not using the product to engage in the unlicensed manufacturing or distribution of controlled 152 153 substances. As used in this subdivision, "roach clip" means an object 154 used to hold burning material, including, but not limited to, a marijuana 155 cigarette, that has become too small or too short to be held between the 156 fingers.

(B) "Factory" means any place used for the manufacturing, mixing,
compounding, refining, processing, packaging, distributing, storing,
keeping, holding, administering or assembling illegal substances
contrary to the provisions of this chapter, or any building, rooms or
location which contains equipment or paraphernalia used for this
purpose.

(21) "Federal Controlled Substances Act, 21 USC 801 et seq." means
Public Law 91-513, the Comprehensive Drug Abuse Prevention and
Control Act of 1970.

(22) "Federal food and drug laws" means the federal Food, Drug andCosmetic Act, as amended, Title 21 USC 301 et seq.

(23) "Hallucinogenic substances" are psychodysleptic substances,
other than cannabis-type substances, which assert a confusional or
disorganizing effect upon mental processes or behavior and mimic
acute psychotic disturbances. Exemplary of such drugs are mescaline,
peyote, psilocyn and d-lysergic acid diethylamide, which are controlled
substances under this chapter unless modified.

(24) "Hospital", as used in sections 21a-243 to 21a-283, inclusive, as
amended by this act, means an institution for the care and treatment of
the sick and injured, approved by the Department of Public Health or
the Department of Mental Health and Addiction Services as proper to
be entrusted with the custody of controlled drugs and substances and
professional use of controlled drugs and substances under the direction
of a licensed practitioner.

181 (25) "Intern" means a person who holds a degree of doctor of 182 medicine or doctor of dental surgery or medicine and whose period of 183 service has been recorded with the Department of Public Health and 184 who has been accepted and is participating in training by a hospital or 185 institution in this state. Doctors meeting the foregoing requirements and 186 commonly designated as "residents" and "fellows" shall be regarded as 187 interns for purposes of this chapter.

188 (26) "Immediate precursor" means a substance which the 189 Commissioner of Consumer Protection has found to be, and by 190 regulation designates as being, the principal compound commonly used 191 or produced primarily for use, and which is an immediate chemical 192 intermediary used or likely to be used, in the manufacture of a 193 controlled substance, the control of which is necessary to prevent, curtail 194 or limit manufacture.

(27) "Laboratory" means a laboratory approved by the Department of
Consumer Protection as proper to be entrusted with the custody of
controlled substances and the use of controlled substances for scientific
and medical purposes and for purposes of instruction, research or
analysis.

(28) "Manufacture" means the production, preparation, cultivation,
growing, propagation, compounding, conversion or processing of a
controlled substance, either directly or indirectly by extraction from
substances of natural origin, or independently by means of chemical
synthesis, or by a combination of extraction and chemical synthesis, and
includes any packaging or repackaging of the substance or labeling or

206 relabeling of its container, except that this term does not include the 207 preparation or compounding of a controlled substance by an individual 208 for the individual's own use or the preparation, compounding, 209 packaging or labeling of a controlled substance: (A) By a practitioner as 210 an incident to the practitioner administering or dispensing of a 211 controlled substance in the course of such practitioner's professional 212 practice; [,] or (B) by a practitioner, or by the practitioner's authorized 213 agent under such practitioner's supervision, for the purpose of, or as an 214 incident to, research, teaching or chemical analysis and not for sale.

215 (29) "Marijuana" means all parts of any plant, or species of the genus cannabis or any infra specific taxon thereof, whether growing or not; 216 217 [the seeds thereof;] the resin extracted from any part of the plant; every 218 compound, manufacture, salt, derivative, mixture [,] or preparation of 219 such plant, or its [seeds or] resin; [,] any high-THC hemp product; 220 manufactured cannabinoids; [, synthetic cannabinoids, except as 221 provided in subparagraph (E) of this subdivision;] or cannabinon, 222 cannabinol or cannabidiol and chemical compounds which are similar 223 to cannabinon, cannabinol or cannabidiol in chemical structure or which 224 are similar thereto in physiological effect, which are controlled 225 substances under this chapter, except cannabidiol derived from hemp, 226 as defined in section 22-61*l*, that is not a high-THC hemp product. 227 "Marijuana" does not include: (A) The mature stalks of such plant, fiber 228 produced from such stalks, oil or cake made from the seeds of such 229 plant, any other compound, manufacture, salt, derivative, mixture or 230 preparation of such mature stalks, except the resin extracted from such 231 mature stalks or fiber, oil or cake; (B) the sterilized seed of such plant 232 which is incapable of germination; (C) hemp, as defined in section 22-233 61*l*, (i) with a total THC concentration of not more than three-tenths per 234 cent on a dry-weight basis, and (ii) that is not a high-THC hemp product; 235 or (D) any substance approved by the federal Food and Drug 236 Administration or successor agency as a drug and reclassified in any 237 schedule of controlled substances or unscheduled by the federal Drug 238 Enforcement Administration or successor agency which is included in 239 the same schedule designated by the federal Drug Enforcement Administration or successor agency. [; or (E) synthetic cannabinoids which are controlled substances that are designated by the Commissioner of Consumer Protection, by whatever official, common, usual, chemical or trade name designation, as controlled substances and are classified in the appropriate schedule in accordance with subsections (i) and (j) of section 21a-243.]

246 (30) "Narcotic substance" means any of the following, whether 247 produced directly or indirectly by extraction from a substance of 248 vegetable origin, or independently by means of chemical synthesis, or 249 by a combination of extraction and chemical synthesis: (A) Morphine-250 type: (i) Opium or opiate, or any salt, compound, derivative [,] or 251 preparation of opium or opiate which is similar to any such substance 252 in chemical structure or which is similar to any such substance in 253 physiological effect and which shows a like potential for abuse, which 254 is a controlled substance under this chapter unless modified; (ii) any 255 salt, compound, isomer, derivative [,] or preparation of any such 256 substance which is chemically equivalent or identical to any substance 257 referred to in clause (i) of this [subdivision] subparagraph, but not 258 including the isoquinoline alkaloids of opium; (iii) opium poppy or 259 poppy straw; or (iv) (I) fentanyl or any salt, compound, derivative or 260 preparation of fentanyl which is similar to any such substance in 261 chemical structure or which is similar to any such substance in 262 physiological effect and which shows a like potential for abuse, which 263 is a controlled substance under this chapter unless modified, or (II) any 264 salt, compound, isomer, derivative or preparation of any such substance 265 which is chemically equivalent or identical to any substance referred to 266 in subclause (I) of this clause; or (B) cocaine-type, [;] coca leaves or any 267 salt, compound, derivative or preparation of coca leaves, or any salt, 268 compound, isomer, derivatives or preparation of any such substance 269 which is chemically equivalent or identical to any such substance or 270 which is similar to any such substance in physiological effect and which 271 shows a like potential for abuse, but not including decocainized coca 272 leaves or extractions of coca leaves which do not contain cocaine or 273 ecgonine.

(31) "Nurse" means a person performing nursing as defined in section20-87a.

(32) "Official written order" means an order for controlled substances
written on a form provided by the bureau for that purpose under the
federal Controlled Substances Act.

(33) "Opiate" means any substance having an addiction-forming or
addiction-sustaining liability similar to morphine or being capable of
conversion into a drug having addiction-forming or addictionsustaining liability; it does not include, unless specifically designated as
controlled under this chapter, the dextrorotatory isomer of 3-methoxyn-methylmorthinan and its salts (dextro-methorphan) but shall include
its racemic and levorotatory forms.

(34) "Opium poppy" means the plant of the species papaversomniferum l., except its seed.

288 (35) Repealed by P.A. 99-102, S. 51.

(36) "Other stimulant and depressant drugs" means controlled
substances other than amphetamine-type, barbiturate-type, cannabistype, cocaine-type, hallucinogenics and morphine-type which are found
to exert a stimulant and depressant effect upon the higher functions of
the central nervous system and which are found to have a potential for
abuse and are controlled substances under this chapter.

(37) "Person" includes any corporation, limited liability company,
association or partnership, or one or more individuals, government or
governmental subdivisions or agency, business trust, estate, trust, or
any other legal entity. Words importing the plural number may include
the singular; words importing the masculine gender may be applied to
females.

301 (38) "Pharmacist" means a person authorized by law to practice
302 pharmacy pursuant to section 20-590, 20-591, 20-592 or 20-593.

303 (39) "Pharmacy" means an establishment licensed pursuant to section

304 20-594.

(40) "Physician" means a person authorized by law to practicemedicine in this state pursuant to section 20-9.

307 (41) "Podiatrist" means a person authorized by law to practice308 podiatry in this state.

309 (42) "Poppy straw" means all parts, except the seeds, of the opium310 poppy, after mowing.

311 (43) "Practitioner" means: (A) A physician, dentist, veterinarian, 312 podiatrist, scientific investigator or other person licensed, registered or 313 otherwise permitted to distribute, dispense, conduct research with 314 respect to or to administer a controlled substance in the course of 315 professional practice or research in this state; and (B) a pharmacy, 316 hospital or other institution licensed, registered or otherwise permitted 317 to distribute, dispense, conduct research with respect to or to administer 318 a controlled substance in the course of professional practice or research 319 in this state.

320 (44) "Prescribe" means order or designate a remedy or any321 preparation containing controlled substances.

322 (45) "Prescription" means a written, oral or electronic order for any
323 controlled substance or preparation from a licensed practitioner to a
324 pharmacist for a patient.

325 (46) "Production" includes the manufacture, planting, cultivation,326 growing or harvesting of a controlled substance.

327 (47) "Registrant" means any person licensed by this state and
328 assigned a current federal Bureau of Narcotics and Dangerous Drug
329 Registry Number as provided under the federal Controlled Substances
330 Act.

(48) "Registry number" means the alphabetical or numericaldesignation of identification assigned to a person by the federal Drug

Enforcement Administration, or other federal agency, which iscommonly known as the federal registry number.

335 (49) "Restricted drugs or substances" are the following substances 336 without limitation and for all purposes: Datura stramonium; 337 hyoscyamus niger; atropa belladonna, or the alkaloids atropine; 338 hyoscyamine; belladonnine; apatropine; or any mixture of these 339 alkaloids such as daturine, or the synthetic homatropine or any salts of 340 these alkaloids, except that any drug or preparation containing any of 341 the above-mentioned substances which is permitted by federal food and 342 drug laws to be sold or dispensed without a prescription or written 343 order shall not be a controlled substance; amyl nitrite; the following 344 volatile substances to the extent that said chemical substances or 345 compounds containing said chemical substances are sold, prescribed, 346 dispensed, compounded, possessed or controlled or delivered or 347 administered to another person with the purpose that said chemical 348 substances shall be breathed, inhaled, sniffed or drunk to induce a 349 stimulant, depressant or hallucinogenic effect upon the higher functions 350 of the central nervous system: Acetone; benzene; butyl alcohol; butyl 351 nitrate and its salts, isomers, esters, ethers or their salts; cyclohexanone; 352 dichlorodifluoromethane; ether; ethyl acetate; formaldehyde; hexane; 353 isopropanol; methanol; methyl cellosolve acetate; methyl ethyl ketone; 354 methyl isobutyl ketone; nitrous oxide; pentochlorophenol; toluene; 355 toluol; trichloroethane; trichloroethylene; 1,4 butanediol.

(50) "Sale" is any form of delivery which includes barter, exchange or
gift, or offer therefor, and each such transaction made by any person
whether as principal, proprietor, agent, servant or employee.

(51) "State", when applied to a part of the United States, includes any
state, district, commonwealth, territory or insular possession thereof,
and any area subject to the legal authority of the United States of
America.

(52) "State food, drug and cosmetic laws" means the Uniform Food,Drug and Cosmetic Act, section 21a-91 et seq.

(53) "Ultimate user" means a person who lawfully possesses a
controlled substance for the person's own use or for the use of a member
of such person's household or for administering to an animal owned by
such person or by a member of such person's household.

369 (54) "Veterinarian" means a person authorized by law to practice370 veterinary medicine in this state.

(55) "Wholesaler" means a distributor or a person who supplies
controlled substances that the person personally has not produced or
prepared to registrants.

(56) "Reasonable times" means the time or times any office, caregiving institution, pharmacy, clinic, wholesaler, manufacturer,
laboratory, warehouse, establishment, store or place of business, vehicle
or other place is open for the normal affairs or business or the practice
activities usually conducted by the registrant.

379 (57) "Unit dose drug distribution system" means a drug distribution 380 system used in a hospital or chronic and convalescent nursing home in 381 which drugs are supplied in individually labeled unit of use packages, 382 each patient's supply of drugs is exchanged between the hospital 383 pharmacy and the drug administration area or, in the case of a chronic 384 and convalescent nursing home between a pharmacy and the drug 385 administration area, at least once each twenty-four hours and each 386 patient's medication supply for this period is stored within a patient-387 specific container, all of which is conducted under the direction of a 388 pharmacist licensed in Connecticut and, in the case of a hospital, directly 389 involved in the provision and supervision of pharmaceutical services at 390 such hospital at least thirty-five hours each week.

(58) "Cocaine in a free-base form" means any substance which
contains cocaine, or any compound, isomer, derivative or preparation
thereof, in a nonsalt form.

(59) "THC" means tetrahydrocannabinol, including, but not limitedto, delta-7, delta-8-tetrahydrocannabinol, delta-9-tetrahydrocannabinol

396 and delta-10-tetrahydrocannabinol, and any material, compound, 397 mixture or preparation which contain their salts, isomers and salts of 398 isomers, whenever the existence of such salts, isomers and salts of 399 isomers is possible within the specific chemical designation, regardless 400 of the source, except: (A) Dronabinol substituted in sesame oil and 401 encapsulated in a soft gelatin capsule in a federal Food and Drug 402 Administration or successor agency approved product; [,] or (B) any 403 tetrahydrocannabinol product that has been approved by the federal 404 Food and Drug Administration or successor agency to have a medical 405 use and reclassified in any schedule of controlled substances or 406 unscheduled by the federal Drug Enforcement Administration or 407 successor agency.

(60) "Total THC" means the sum of the percentage by weight of
tetrahydrocannabinolic acid, multiplied by eight hundred seventyseven-thousandths, plus the percentage of weight of THC.

(61) "Manufactured cannabinoid" means cannabinoids [naturally
occurring from a source other than marijuana that are similar in
chemical structure or physiological effect to cannabinoids derived from
marijuana, as defined in section 21a-243, but are derived by a chemical
or biological process] created by directly converting one cannabinoid to
a different cannabinoid through: (A) Application of light or heat; or (B)
decarboxylation of naturally occurring acidic forms of cannabinoids.

418 (62) "Synthetic cannabinoid": [means] (A) Means any [material, 419 compound, mixture or preparation which contains any quantity of a 420 substance having a psychotropic response primarily by agonist activity 421 at cannabinoid-specific receptors affecting the central nervous system 422 that is produced artificially and not derived from an organic source 423 naturally containing cannabinoids, unless listed in another schedule 424 pursuant to section 21a-243] substance converted, by a chemical process, 425 to create a cannabinoid or cannabinoid-like substance that (i) has 426 structural features which allow interaction with at least one of the 427 known cannabinoid-specific receptors, or (ii) has any physiological or 428 psychotropic response on at least one cannabinoid-specific receptor; and (B) excludes any cannabinoid that is (i) naturally produced, or (ii)
 manufactured through (I) application of light or heat, or (II)
 decarboxylation of naturally occurring acidic forms of cannabinoids.

432 (63) "High-THC hemp product" means a manufacturer hemp 433 product, as defined in section 22-61*l*, that has, or is advertised, labeled 434 or offered for sale as having, total THC that exceeds: (A) [for] For a hemp 435 edible, hemp topical or hemp transdermal patch (i) one milligram on a 436 per-serving basis, or (ii) five milligrams on a per-container basis; [,] (B) 437 for a hemp tincture, including, but not limited to, oil intended for 438 ingestion by swallowing, buccal administration or sublingual 439 absorption, (i) one milligram on a per-serving basis, or (ii) twenty-five 440 milligrams on a per-container basis; [,] (C) for a hemp concentrate or 441 extract, including, but not limited to, a vape oil, wax or shatter, twenty-442 five milligrams on a per-container basis; [,] or (D) for a manufacturer 443 hemp product not described in subparagraph (A), (B) or (C) of this 444 subdivision, (i) one milligram on a per-serving basis, (ii) five milligrams 445 on a per-container basis, or (iii) three-tenths per cent on a dry-weight 446 basis for cannabis flower or cannabis trim.

447 Sec. 2. Subsection (j) of section 21a-243 of the general statutes is 448 repealed and the following is substituted in lieu thereof (*Effective from* 449 *passage*):

(j) Notwithstanding the provisions of subsection (c) of this section, the Commissioner of Consumer Protection shall designate the following substances, by whatever official, common, usual, chemical or trade name designation, as controlled substances in schedule I of the controlled substances scheduling regulations:

- 455 (1) Mephedrone (4-methylmethcathinone); [and]
- 456 (2) Synthetic cannabinoids; and
- 457 [(2)] (3) MDPV (3,4-methyenedioxypyrovalerone).
- 458 Sec. 3. Subparagraph (N) of subdivision (5) of subsection (b) of section

459 21a-421j of the 2024 supplement to the general statutes is repealed and460 the following is substituted in lieu thereof (*Effective from passage*):

(N) A requirement that, prior to being sold and transferred to a
consumer, qualifying patient or caregiver, cannabis packaging be
clearly labeled, whether printed directly on such packaging or affixed
by way of a separate label, other than an extended content label, with:

(i) A unique identifier generated by a cannabis analytic tracking
system maintained by the department and used to track cannabis under
the policies and procedures issued, and final regulations adopted, by
the commissioner pursuant to this section; and

(ii) The following information concerning the cannabis contained in
such packaging, which shall be in legible English, black lettering, Times
New Roman font, flat regular typeface, on a contrasting background
and in uniform size of not less than one-tenth of one inch, based on a
capital letter "K", which information shall also be available on the
Internet web site of the cannabis establishment that sells and transfers
such cannabis:

(I) The name of such cannabis, as registered with the department
under the policies and procedures issued, and final regulations adopted,
by the commissioner pursuant to this section.

(II) The expiration date, which shall not account for any refrigerationafter such cannabis is sold and transferred to the consumer, qualifyingpatient or caregiver.

(III) The net weight or volume, expressed in metric and imperialunits.

(IV) The standardized serving size, expressed in customary units, andthe number of servings included in such packaging, if applicable.

486 (V) Directions for use and storage.

487 (VI) Each active ingredient comprising at least one per cent of such

488 cannabis, including cannabinoids, isomers, esters, ethers and salts and 489 salts of isomers, esters and ethers, and all quantities thereof expressed 490 in metric units and as a percentage of volume. 491 (VII) A list of all known allergens, as identified by the federal Food 492 and Drug Administration, contained in such cannabis, or the denotation 493 "no known FDA identified allergens" if such cannabis does not contain 494 any allergen identified by the federal Food and Drug Administration. 495 (VIII) The following warning statement within, and outlined by, a red 496 box: 497 "This product is not FDA-approved, may be intoxicating, cause long-498 term physical and mental health problems, and have delayed side 499 effects. It is illegal to operate a vehicle or machinery under the influence 500 of cannabis. Keep away from children." 501 (IX) At least one of the following warning statements, rotated 502 quarterly on an alternating basis: 503 "Warning: Frequent and prolonged use of cannabis can contribute to 504 mental health problems over time, including anxiety, depression, 505 stunted brain development and impaired memory." 506 "Warning: Consumption while pregnant or breastfeeding may be harmful." 507 508 "Warning: Cannabis has intoxicating effects and may be habit-509 forming and addictive." 510 "Warning: Consuming more than the recommended amount may 511 result in adverse effects requiring medical attention.". 512 (X) All information necessary to comply with labeling requirements 513 imposed under the laws of this state [or] and federal law, including, but 514 not limited to, sections 21a-91 to 21a-120, inclusive, and 21a-151 to 21a-515 159, inclusive, the Federal Food, Drug and Cosmetic Act, 21 USC 301 et 516 seq., as amended from time to time, and the federal Fair Packaging and

517 Labeling Act, 15 USC 1451 et seq., as amended from time to time, for518 similar products that do not contain cannabis.

(XI) Such additional warning labels for certain cannabis products as
the commissioner may require and post on the department's Internet
web site.

522 Sec. 4. Section 21a-421aa of the general statutes is repealed and the 523 following is substituted in lieu thereof (*Effective from passage*):

524 (a) No cannabis retailer or hybrid retailer shall accept payment or 525 other form of compensation directly or indirectly from a cultivator, 526 micro-cultivator, producer, food and beverage manufacturer, product 527 manufacturer or product packager to carry a cannabis product or for 528 placement or promotion of such product in a retailer or hybrid retailer's 529 establishment or through other promotional initiatives. No retailer or 530 hybrid retailer shall enter into a contract with a cultivator, micro-531 cultivator, producer, food and beverage manufacturer, product 532 manufacturer or product packager that requires or permits preferential 533 treatment, exclusivity or near exclusivity or limits a retailer or hybrid 534 retailer from purchasing from other cultivators, micro-cultivators, 535 producers, food and beverage manufacturers or product manufacturers 536 in any way.

(b) No cannabis establishment shall produce, manufacture or sellcannabis that is intended for use or consumption by animals.

539 (c) A retailer or hybrid retailer shall not knowingly sell to a consumer 540 more than one ounce of cannabis or the equivalent amount of cannabis 541 products or combination of cannabis and cannabis products, as set forth 542 in subsection (i) of section 21a-279a, per day, except that a hybrid retailer 543 or dispensary facility may sell up to five ounces of cannabis or the 544 equivalent amount of cannabis products or combination of cannabis and 545 cannabis products to a qualifying patient or caregiver per day. 546 Notwithstanding the requirements of sections 4-168 to 4-172, inclusive, 547 to avoid cannabis supply shortages or address a public health and safety 548 concern, the commissioner may set temporary lower per-transaction

549 limits, which shall be published on the department's Internet web site.
550 Such limits shall become ineffective upon the commissioner's
551 determination that a supply shortage or public health and safety
552 concern no longer exists.

553 (d) No cannabis establishment, except a producer, cultivator or 554 micro-cultivator, may acquire or possess a live cannabis plant.

(e) No person issued a license or registration pursuant to RERACA
shall (1) assign or transfer such license or registration without the
commissioner's prior approval, or (2) sell, transfer or transport cannabis
to, or obtain cannabis from, a location outside of this state if such activity
would be in violation of federal law.

560 (f) Synthetic cannabinoids, as defined in section 21a-240, as amended
561 by this act, are prohibited in cannabis, and no synthetic cannabinoid
562 may be sold at any cannabis establishment.

563 Sec. 5. Subsection (a) of section 21a-421dd of the general statutes is 564 repealed and the following is substituted in lieu thereof (*Effective from* 565 *passage*):

566 (a) No member of the Social Equity Council and no employee of the 567 Social Equity Council or department who carries out the licensing, 568 inspection, investigation, enforcement or policy decisions authorized by 569 [RERACA] this chapter, and any regulations enacted pursuant thereto, 570 may, directly or indirectly, have any management or financial interest 571 in the cultivation, manufacture, sale, transportation, delivery or testing 572 of cannabis in this state, nor receive any commission or profit from nor 573 have any financial interest in purchases or sales made by [persons] 574 cannabis establishments that are licensed pursuant to this chapter and 575 authorized to make such purchases or sales pursuant to [RERACA] such 576 license. No provision of this section shall prevent any such member or 577 employee from purchasing and keeping in his or her possession, for his 578 or her personal use or the use of such member's or employee's family or 579 guests, any cannabis which may be purchased or kept by any person by 580 virtue of [RERACA] this chapter.

581 Sec. 6. Section 22-61m of the 2024 supplement to the general statutes
582 is repealed and the following is substituted in lieu thereof (*Effective from passage*):

(a) No person shall manufacture in the state without a license tomanufacture issued by the Commissioner of Consumer Protection.

(b) Each applicant for a manufacturer license shall submit anapplication on a form and in a manner prescribed by the Commissionerof Consumer Protection.

589 (c) The following fees shall apply for a license to manufacture:

590 (1) A nonrefundable license application fee of seventy-five dollars;591 and

592 (2) A nonrefundable licensing fee of three hundred seventy-five593 dollars for a license to manufacture hemp.

(d) A license to manufacture issued by the Commissioner of
Consumer Protection pursuant to this section shall expire triennially on
June thirtieth. Such licenses shall not be transferable.

(e) In accordance with a hearing held pursuant to chapter 54, the
Commissioner of Consumer Protection may deny, suspend or revoke a
manufacturer license, issue fines of not more than two thousand five
hundred dollars per violation and place conditions upon a
manufacturer licensee who violates the provisions of this section and
any regulation adopted pursuant to this section.

(f) (1) Any individual who manufactures in this state without
obtaining a license pursuant to this section or who manufactures in this
state after such entity's license is suspended or revoked shall be fined
two hundred fifty dollars in accordance with the provisions of section
51-164n.

608 (2) Any entity who manufactures in this state without obtaining a 609 license pursuant to this section, or who manufactures in this state after having a license suspended, shall be fined not more than two thousand
five hundred dollars per violation after a hearing conducted in
accordance with the provisions of chapter 54.

(g) Nothing in this chapter or any regulations adopted pursuant to
this chapter shall be construed to apply to persons licensed pursuant to
section 21a-408i nor to require persons licensed pursuant to said section
to obtain a license pursuant to this chapter.

(h) The Commissioner of Consumer Protection may inspect and shall
have access to the buildings, equipment, supplies, vehicles, records, real
property and other information of any manufacturer applicant or
licensee that the commissioner deems necessary to carry out the
commissioner's duties pursuant to this section.

622 (i) (1) Each manufacturer shall follow the protocol in this subsection 623 for disposing of cannabis in the event that any hemp or hemp product 624 is deemed to exceed the prescribed THC concentration, as determined 625 by the Commissioner of Consumer Protection, or a manufacturer 626 licensee in possession of hemp or hemp products who desires to dispose 627 of obsolete, misbranded, excess or otherwise undesired product. Each 628 manufacturer licensee shall be responsible for all costs of disposal of 629 hemp samples and any hemp produced by such licensee that violates the provisions of this section or any regulation adopted pursuant to this 630 631 section. Any cannabis that exceeds the prescribed THC concentration 632 allowable in hemp or hemp products shall be immediately embargoed 633 by such manufacturer and clearly labeled as adulterated by such 634 licensee and such licensee shall immediately notify both the Department 635 of Consumer Protection and the Department of Agriculture, in writing, 636 of such adulterated product. Such adulterated product shall be 637 destroyed and disposed of by the following method, as determined by 638 the Commissioner of Consumer Protection:

(A) Surrender, without compensation, of such hemp or hemp product
to the Commissioner of Consumer Protection who shall be responsible
for the destruction and disposal of such adulterated product; or

(B) By disposal in a manner prescribed by the Commissioner ofConsumer Protection.

(2) Notwithstanding the provisions of subdivision (1) of this
subsection, upon written request of a manufacturer, the Commissioner
of Consumer Protection may permit such manufacturer to combine
different batches of raw hemp plant material to achieve a THC
concentration of 0.3 per cent on a dry weight basis, in lieu of embargo
or destruction.

(j) The manufacturer or manufacturer's authorized designee
disposing of the hemp or hemp products shall maintain and make
available to the Commissioner of Consumer Protection a record of each
such disposal or destruction of product indicating:

(1) The date, time and location of disposal or destruction;

655 (2) The manner of disposal or destruction;

(3) The batch or lot information and quantity of hemp or hempproduct disposed of or destroyed; and

(4) The signatures of the persons disposing of the hemp or hemp
products, the authorized representative of the Commissioner of
Consumer Protection and any other persons present during the
disposal.

662 (k) Any hemp intended to be manufactured by a manufacturer into a 663 manufacturer hemp product shall be tested by an independent testing 664 laboratory located in this state. A manufacturer licensee shall make 665 available samples, in an amount and type determined by the Commissioner of Consumer Protection, of hemp for an independent 666 667 testing laboratory employee to select random samples. The independent 668 testing laboratory shall test each sample in accordance with the 669 laboratory testing standards established in policies, procedures and 670 regulations adopted by the commissioner pursuant to section 21a-421j, 671 as amended by this act.

672 (1) Once a batch of hemp, intended to be sold as a manufacturer hemp 673 product, has been homogenized for sample testing and eventual 674 packaging and sale, until the independent testing laboratory provides 675 the results from its tests and analysis, the manufacturer shall segregate 676 and withhold from use the entire batch of hemp that is intended for use 677 as a manufacturer hemp product, except the samples that have been 678 removed by the independent testing laboratory for testing. During this 679 period of segregation, the manufacturer licensee shall maintain the 680 hemp batch in a secure, cool and dry location, as prescribed by the 681 Commissioner of Consumer Protection, so as to prevent the hemp from 682 becoming adulterated. Such manufacturer shall not manufacture or sell 683 a manufacturer hemp product prior to the time that the independent 684 testing laboratory completes testing and analysis and provides such 685 results, in writing, to the manufacturer licensee who initiated such 686 testing.

(m) An independent testing laboratory shall immediately return or
dispose of any hemp or manufacturer hemp product upon the
completion of any testing, use or research. If an independent testing
laboratory disposes of hemp or manufacturer hemp products, the
laboratory shall dispose of such hemp in the following manner, as
determined by the Commissioner of Consumer Protection:

(1) By surrender, without compensation, of such hemp or
manufacturer hemp product to the Commissioner of Consumer
Protection who shall be responsible for the destruction and disposal of
such hemp or hemp product; or

697 (2) By disposal in a manner prescribed by the Commissioner of698 Consumer Protection.

(n) If a sample does not pass the microbiological, mycotoxin, heavy
metal or pesticide chemical residue test, based on the laboratory testing
standards established in policies, procedures and regulations adopted
by the Commissioner of Consumer Protection pursuant to section 21a421j, as amended by this act, the manufacturer licensee who sent such

704 batch for testing shall:

705 (1) Retest and reanalyze the hemp from which the sample was taken 706 by having an employee from the same laboratory randomly select 707 another sample from the same hemp batch. If the sample used to retest 708 or reanalyze such hemp yields satisfactory results for all testing 709 required under this section, an employee from a different laboratory 710 shall randomly select a different sample from the same hemp batch for 711 testing. If both samples yield satisfactory results for all testing required 712 under this section, the hemp batch from which the samples were taken 713 shall be released for manufacturing, processing and sale;

714 (2) If a remediation plan sufficient to ensure public health and safety 715 is submitted to and approved by the commissioner, remediate the hemp 716 batch from which the sample was taken and have a laboratory employee 717 randomly select a sample from such remediated hemp batch for testing. 718 If such randomly selected sample yields satisfactory results for any 719 testing required under this section, an employee from a different 720 laboratory shall randomly select a different sample from the same hemp 721 batch for testing. If both samples yield satisfactory results for all testing 722 required under this section, the hemp batch from which the samples 723 were taken may be released for manufacturing, processing or sale; or

(3) If the manufacturer does not retest or remediate, or if any
subsequent laboratory testing does not yield satisfactory results for any
testing required under this section, dispose of the entire batch from
which the sample was taken in accordance with procedures established
by the Commissioner of Consumer Protection pursuant to subdivision
(1) of subsection (i) of this section.

(o) If a sample passes the microbiological, mycotoxin, heavy metal
and pesticide chemical residue test, the independent testing laboratory
shall release the entire batch for manufacturing, processing or sale.

(p) The independent testing laboratory shall file with the Department
of Consumer Protection an electronic copy of each laboratory test result
for any batch that does not pass the microbiological, mycotoxin, heavy

736 metal or pesticide chemical residue test, at the same time that it 737 transmits such results to the manufacturer licensee who requested such 738 testing. Each independent testing laboratory shall maintain the test 739 results of each tested batch for a period of three years and shall make 740 such results available to the Department of Consumer Protection upon 741 request.

(q) Manufacturers shall maintain records required by the federal act,
this section, any regulation adopted pursuant to this section and the
policies, procedures and regulations adopted by the Commissioner of
Consumer Protection pursuant to section 21a-421j, as amended by this
act. Each manufacturer shall make such records available to the
Department of Consumer Protection immediately upon request and in
electronic format, if available.

749 (r) The Commissioner of Consumer Protection may adopt 750 regulations, in accordance with the provisions of chapter 54, to 751 implement the provisions of this section including, but not limited to, 752 establishing sampling and testing procedures to ensure compliance 753 with this section, prescribing storage and disposal procedures for hemp, 754 marijuana and manufacturer hemp products that fail to pass 755 Department of Consumer Protection prescribed independent testing 756 laboratory testing standards and establishing advertising and labeling 757 requirements for manufacturer hemp products.

(s) Any claim of health impacts, medical effects or physical or mental
benefits shall be prohibited on any advertising for, labeling of or
marketing of manufacturer hemp products regardless of whether such
manufacturer hemp products were manufactured in this state or
another jurisdiction. Any violation of this subsection shall be deemed an
unfair or deceptive trade practice under subsection (a) of section 42110b.

(t) Not later than February 1, 2020, the Commissioners of Agriculture
and Consumer Protection shall submit a report, in accordance with
section 11-4a, to the joint standing committee of the general assembly

having cognizance of matters relating to the environment on the status
of the pilot program, the development of the state plan and any
regulations for such pilot program or state plan. Such report shall also
include any legislative recommendations, including, but not limited to,
any recommendations for requiring the registration of any
manufacturer hemp product offered for sale in this state.

774 (u) (1) Any person who sells manufacturer hemp products shall not 775 be required to be licensed, provided such person only engages in: (A) 776 The retail or wholesale sale of manufacturer hemp products in which no 777 further manufacturing of hemp occurs, provided such manufacturer 778 hemp products are acquired from a person authorized to manufacture 779 the manufacturer hemp products under the laws of this state or another 780 state, territory or possession of the United States or another sovereign 781 entity; (B) the acquisition of manufacturer hemp products for the sole 782 purpose of product distribution for resale; and (C) the retail sale of 783 manufacturer hemp products that is authorized under federal or state 784 law.

(2) The Commissioner of Consumer Protection or Commissioner of
Revenue Services may, pursuant to section 4-182, summarily suspend
any credential the Department of Consumer Protection or Department
of Revenue Services issued to any person who sells manufacturer hemp
products in violation of subdivision (1) of this subsection or subsections
(v) to (y), inclusive, of this section.

(v) No manufacturer hemp product offered for sale in this state, or to
a consumer in this state, shall contain any synthetic cannabinoid, as
defined in section 21a-240, as amended by this act.

(w) No manufacturer hemp product offered for sale in this state, or
to a consumer in this state, shall be packaged, presented or advertised
in a manner that is likely to mislead a consumer by incorporating any
statement, brand, design, representation, picture, illustration or other
depiction that: (1) Bears a reasonable resemblance to trademarked or
characteristic packaging of (A) cannabis offered for sale (i) in this state

800 by a cannabis establishment licensed in this state, or (ii) on tribal land 801 by a tribal-credentialed cannabis entity, or (B) a commercially available product other than a cannabis product, as defined in section 21a-420; or 802 803 (2) implies that the manufacturer hemp product (A) is a cannabis 804 product, as defined in section 21a-420, (B) contains a total THC 805 concentration greater than three-tenths per cent on a dry-weight basis, 806 or (C) is a high-THC hemp product, as defined in section 21a-240, as 807 amended by this act.

(x) No manufacturer hemp product that is a food, beverage, oil or
other product intended for human ingestion shall be distributed or sold
in this state unless such product is contained within a package, or a label
is affixed to such package, that includes:

812 (1) A scannable barcode, Internet web site address or quick response
813 code that is linked to the certificate of analysis of the final form product
814 batch by an independent testing laboratory and discloses:

815 (A) The name of such product;

816 (B) The name, address and telephone number of such product's817 manufacturer, packer and distributor, as applicable;

818 (C) The batch number, which shall match the batch number on such819 package or label; and

(D) The concentration of cannabinoids present in such product,
including, but not limited to, total THC and any cannabinoids or active
ingredients comprising at least one per cent of such product;

- 823 (2) The expiration or best by date for such product, if applicable;
- 824 (3) A clear and conspicuous statement disclosing that:

(A) Children, or those who are pregnant or breastfeeding, should
avoid using such product prior to consulting with a health care
professional concerning such product's safety;

(B) Products containing cannabinoids should be kept out of reach ofchildren; and

(C) The federal Food and Drug Administration has not evaluatedsuch product for safety or efficacy; and

(4) If such product is intended to be inhaled, a clear and conspicuous
warning statement disclosing that smoking or vaporizing is hazardous
to human health.

(y) No manufacturer hemp product that is a topical, soap or cosmetic,
as defined in section 21a-92, shall be distributed or sold in this state
unless such product is contained within a package, or a label is affixed
to such package, that includes:

(1) A scannable barcode, Internet web site address or quick response
code that is linked to the certificate of analysis of the final form extract
or final form product batch by an independent testing laboratory and
discloses:

843 (A) The name of such product;

(B) The name, address and telephone number of such product'smanufacturer, packer and distributor, as applicable;

846 (C) The batch number, which shall match the batch number on such847 package or label; and

848 (D) The concentration of cannabinoids present in such batch, 849 including, but not limited to, total THC and any marketed cannabinoids;

- 850 (2) The expiration or best by date for such product, if applicable; and
- 851 (3) A clear and conspicuous statement disclosing the following:

852 "THE FDA HAS NOT EVALUATED THIS PRODUCT FOR SAFETY853 OR EFFICACY.".

854 (z) Any violation of subsections (u) to (y), inclusive, of this section

shall be deemed an unfair or deceptive trade practice under subsection(a) of section 42-110b.

(aa) Not later than October 31, 2023, the Department of Emergency
Services and Public Protection shall, in consultation with the
Department of Consumer Protection, publish a training bulletin to
inform local law enforcement agencies and officers regarding the
investigation and enforcement standards concerning cannabis and highTHC hemp products.

(bb) Notwithstanding any provision of the general statutes: (1) CBD
that is found in manufacturer hemp products shall not be considered a
controlled substance, as defined in section 21a-240, as amended by this
act, or legend drug, as defined in section 20-571; and (2) CBD derived
from hemp and contained in manufacturer hemp products shall not be
considered a controlled substance or adulterant.

869 (cc) Nothing in this section shall be construed to prohibit shipment or

870 transportation through this state of any hemp that is lawfully produced

sections:		
Section 1	from passage	21a-240
Sec. 2	from passage	21a-243(j)
Sec. 3	from passage	21a-421j(b)(5)(N)
Sec. 4	from passage	21a-421aa
Sec. 5	from passage	21a-421dd(a)
Sec. 6	from passage	22-61m

This act shall take effect as follows and shall amend the following

871 <u>under federal law.</u>

## Statement of Purpose:

To: (1) Redefine "marijuana", "manufactured cannabinoid" and "synthetic cannabinoid" for the purposes of various statutes concerning cannabis regulation; (2) require the Commissioner of Consumer Protection to designate any synthetic cannabinoid as a schedule I controlled substance; (3) provide that (A) cannabis shall not contain any synthetic cannabinoid, and (B) no cannabis establishment may sell any synthetic cannabinoid; (4) modify certain provisions that prohibit members of the Social Equity Council and certain employees of the council or the Department of Consumer Protection from holding certain interests in, or receiving certain moneys from, certain cannabis establishments or cannabis transactions; (5) specify that hemp that is lawfully produced under federal law may be transported or shipped through this state; and (6) make minor, technical and conforming changes to various statutes concerning cannabis regulation.

[Proposed deletions are enclosed in brackets. Proposed additions are indicated by underline, except that when the entire text of a bill or resolution or a section of a bill or resolution is new, it is not underlined.]