

General Assembly

Substitute Bill No. 6768

January Session, 2023

AN ACT CONCERNING THE DEPARTMENT OF CONSUMER PROTECTION'S RECOMMENDATIONS REGARDING PRESCRIPTION DRUG REGULATION.

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

Section 1. (NEW) (*Effective January 1, 2024*) (a) For the purposes of this
 section:

3 (1) "Centralized dispensing practitioner" means a prescribing 4 practitioner (A) who is employed by, or affiliated with, a dispensing 5 group practice, and (B) whom the dispensing group practice designates 6 as the prescribing practitioner who is authorized to dispense legend 7 drugs and legend devices on behalf of other prescribing practitioners 8 who are employed by, or affiliated with, such dispensing group 9 practice;

10 (2) "Department" means the Department of Consumer Protection;

(3) "Dispense" has the same meaning as provided in section 20-571 ofthe general statutes;

(4) "Dispensing assistant" means an individual who is (A) registered
with the department under subdivision (1) of subsection (d) of this
section, (B) employed by a dispensing group practice, and (C)

supervised by (i) the centralized dispensing practitioner, or (ii) apharmacist employed by the dispensing group practice;

(5) "Dispensing group practice" means a group practice that (A) centralizes the dispensing of legend drugs or legend devices prescribed by prescribing practitioners who are employed by, or affiliated with, the group practice through (i) a centralized dispensing practitioner, or (ii) a pharmacist employed by the dispensing group practice, and (B) is registered with the department pursuant to subsection (b) of this section;

(6) "Group practice" has the same meaning as provided in section 19a486i of the general statutes;

27 (7) "Legend device" has the same meaning as provided in section 20-28 571 of the general statutes;

(8) "Legend drug" has the same meaning as provided in section 20-571 of the general statutes;

(9) "Pharmacist" has the same meaning as provided in section 20-571of the general statutes;

(10) "Pharmacy technician" means an individual who is registered
with the department and qualified in accordance with section 20-598a
of the general statutes;

36 (11) "Prescribing practitioner" has the same meaning as provided in
37 section 20-571 of the general statutes;

(12) "Prescription" has the same meaning as provided in section 20-635 of the general statutes;

40 (13) "Professional samples" has the same meaning as provided in41 section 20-14c of the general statutes; and

(14) "Seventy-two-hour supply" means a quantity of a legend drug orlegend device that does not exceed the dosage amount necessary for

seventy-two hours according to the directions for use of the legend drugor legend device.

46 (b) (1) No group practice may dispense legend drugs or legend 47 devices as a dispensing group practice unless such group practice 48 submits an application to, and receives a registration from, the 49 department under this subdivision. Each application submitted to the 50 department under this subdivision shall be submitted on a form, and in 51 a manner, prescribed by the department and designate a centralized 52 dispensing practitioner or a pharmacist who is employed by the group 53 practice and shall serve as the primary contact for the department, and 54 shall be accompanied by a registration fee in the amount of two hundred 55 dollars. Each registration issued pursuant to this subdivision shall be 56 valid for a period of two years, and the department may renew such 57 registration for additional two-year periods upon its receipt of a 58 complete renewal application submitted on a form, and in a manner, 59 prescribed by the department and a renewal fee of two hundred dollars.

60 (2) Except as provided in subdivision (3) of this subsection, each 61 dispensing group practice that dispenses, or proposes to dispense, in 62 this state more than a seventy-two-hour supply of any legend drug or 63 legend device shall (A) register for access to the electronic prescription 64 drug monitoring program established pursuant to subsection (j) of 65 section 21a-254 of the general statutes, and (B) comply with all reporting 66 and usage requirements for the electronic prescription drug monitoring 67 program as set forth in subsection (j) of section 21a-254 of the general 68 statutes.

(3) No dispensing group practice that dispenses, or proposes to
dispense, less than a seventy-two-hour supply of legend drugs or
legend devices shall be subject to the provisions of subdivision (2) of this
subsection if such dispensing group practice exclusively dispenses such
supply of legend drugs or legend devices as professional samples.

(c) A dispensing group practice that employs a pharmacist for thepurpose of dispensing legend drugs or legend devices shall not be

76 required to obtain a pharmacy license for the dispensing group 77 practice's premises under section 20-594 of the general statutes. The pharmacist shall report directly to a prescribing practitioner who is 78 79 employed by, or affiliated with, the dispensing group practice, and may 80 supervise dispensing assistants employed by such dispensing group 81 practice, perform in-process and final checks without obtaining any 82 additional verification from the prescribing practitioner to whom such 83 pharmacist reports and perform any component of the practice of 84 pharmacy.

85 (d) (1) No individual may act as a dispensing assistant unless such 86 individual submits an application to, and receives a registration from, 87 the department under this subdivision. Each application submitted to 88 the department under this subdivision shall be submitted on a form, and 89 in a manner, prescribed by the department, and shall be accompanied 90 by a registration fee in the amount of one hundred dollars. Each 91 registration issued pursuant to this subdivision shall be valid for a 92 period of two years, and the department may renew such registration 93 for additional two-year periods upon its receipt of a complete renewal 94 application submitted on a form, and in a manner, prescribed by the 95 department and a renewal fee of one hundred dollars.

96 (2) A dispensing assistant who is registered with the department 97 under subdivision (1) of this subsection may perform the duties of a 98 pharmacy technician, provided the dispensing assistant performs such 99 duties under the supervision of a prescribing practitioner who is 100 employed by or affiliated with, or a pharmacist who is employed by, the 101 dispensing group practice that employs such dispensing assistant. Each 102 dispensing assistant shall be subject to the same responsibilities and 103 liabilities set forth in chapter 400j of the general statutes, and any 104 regulations adopted pursuant to chapter 400j of the general statutes, 105 concerning pharmacy technicians.

(e) A prescribing practitioner who is employed by, or affiliated with,
a dispensing group practice may dispense legend drugs or legend
devices to the prescribing practitioner's patients without engaging the

services of the centralized dispensing practitioner or a pharmacist whois employed by the dispensing group practice.

(f) (1) No centralized dispensing practitioner or pharmacist employed by a dispensing group practice shall dispense a legend drug, legend device or controlled substance for, or order that a legend drug, legend device or controlled substance be dispensed to, any individual who is not being treated by a prescribing practitioner who is employed by, or affiliated with, the dispensing group practice.

(2) No dispensing group practice shall accept or dispense anyprescription from a prescribing practitioner who is not employed by, oraffiliated with, the dispensing group practice.

(3) No dispensing group practice shall exhibit within or upon the
outside of the premises occupied by such dispensing group practice, or
include in any advertisement for such dispensing group practice, (A) the
words "drug store", "pharmacy", "apothecary" or "medicine shop" or any
combination thereof, or (B) any other display, symbol or word
indicating that such dispensing group practice or premises is a
pharmacy.

127 (g) The department may refuse to issue or renew a dispensing group 128 practice registration under subsection (b) of this section or a dispensing 129 assistant registration under subsection (d) of this section, revoke, 130 suspend or place conditions on a dispensing group practice's 131 registration issued under subsection (b) of this section or a dispensing 132 assistant's registration under subsection (d) of this section, and assess a 133 civil penalty not to exceed one thousand dollars per violation if the 134 dispensing group practice or a centralized dispensing practitioner, 135 dispensing assistant or pharmacist employed by, or acting as an agent 136 on behalf of, such dispensing group practice violates any provision of 137 (1) subsections (a) to (f), inclusive, of this section, or (2) chapter 400j of 138 the general statutes, or any regulations adopted pursuant to chapter 400j 139 of the general statutes, concerning dispensing legend drugs or legend 140 devices.

Sec. 2. (NEW) (*Effective from passage*) (a) For the purposes of this section, "drug", "legend device", "pharmacist" and "prescribing practitioner" have the same meanings as provided in section 20-571 of the general statutes.

(b) A pharmacist may authorize or refill a prescription for a legend
device if such legend device is approved by the federal Food and Drug
Administration for use in combination with a drug prescribed by a
prescribing practitioner.

(c) A pharmacist who dispenses a legend device as described in
subsection (b) of this section shall identify the prescribing practitioner
who prescribed the drug that is associated with such legend device, and
shall send written notice to such prescribing practitioner, not later than
seventy-two hours after the pharmacist dispenses such legend device to
the patient, disclosing that such pharmacist dispensed such legend
device to such patient.

156 Sec. 3. (NEW) (*Effective from passage*) (a) For the purposes of this 157 section:

158 (1) "Department" means the Department of Consumer Protection;

(2) "Emergency contraceptive" means a drug, or a combination of
drugs, approved by the federal Food and Drug Administration to
prevent pregnancy as soon as possible following (A) unprotected sexual
intercourse, or (B) a known or suspected contraceptive failure;

(3) "Hormonal contraceptive" means a drug, including, but not
limited to, a hormonal contraceptive patch, an intravaginal hormonal
contraceptive or an oral hormonal contraceptive, composed of a
hormone, or a combination of hormones, approved by the federal Food
and Drug Administration to prevent pregnancy;

(4) "Legend drug" has the same meaning as provided in section 20-571 of the general statutes;

(5) "Pharmacist" has the same meaning as provided in section 20-571of the general statutes;

(6) "Pharmacy" has the same meaning as provided in section 20-571of the general statutes;

(7) "Pharmacy technician" has the same meaning as provided insection 20-571 of the general statutes; and

(8) "Prescribe" means to order, or designate a remedy or anypreparation of, a legend drug for a specific patient.

(b) A pharmacist certified in accordance with the provisions of this
section may prescribe, in good faith, an emergency contraceptive or
hormonal contraceptive to a patient subject to the following conditions:

(1) The pharmacist has completed an educational training program
that (A) concerns prescribing emergency contraceptives and hormonal
contraceptives by a pharmacist, (B) addresses appropriate medical
screening of patients, contraindications, drug interactions, treatment
strategies and modifications and when to refer patients to medical
providers, and (C) is accredited by the Accreditation Council for
Pharmacy Education;

(2) The pharmacist has reviewed the most current version of the
United States Medical Eligibility Criteria for Contraceptive Use
published by the Centers for Disease Control and Prevention, or any
successor document thereto, prior to prescribing any emergency
contraceptive or hormonal contraceptive and, if the pharmacist deviates
from the guidance provided in such document, documents the
pharmacist's rationale in deviating from such guidance in writing;

(3) Prior to dispensing an emergency contraceptive or hormonal
contraceptive and at least once per calendar year thereafter for any
returning patient, the pharmacist completes a screening document,
which the department shall make available on the department's Internet
web site, and the pharmacist, or the pharmacy that employs such

pharmacist, retains such document for at least three years, except
nothing in this subdivision shall be construed to prevent a pharmacist,
in the pharmacist's professional discretion, from issuing a prescription
for a hormonal contraceptive for a period not to exceed twelve months
or from requiring more frequent screenings;

205 (4) If the pharmacist determines that prescribing an emergency 206 contraceptive or hormonal contraceptive to a patient is clinically 207 appropriate, the pharmacist shall (A) counsel the patient about what the 208 patient should monitor and when the patient should seek additional 209 medical attention, and (B) send notice to any health care provider that 210 the patient identifies as the patient's primary care provider or, if the 211 patient does not disclose the identity of the patient's primary care 212 provider, provide to the patient any relevant documentation; and

(5) The pharmacist provides to the patient a document outlining ageappropriate health screenings that are consistent with recommendations
made by the Centers for Disease Control and Prevention.

216 (c) A pharmacy technician may, at a pharmacist's request, assist the 217 pharmacist in prescribing an emergency contraceptive or hormonal 218 contraceptive to a patient by providing screening documentation to the 219 patient, taking and recording the patient's blood pressure and 220 documenting the patient's medical history, provided the pharmacy 221 technician has completed an educational training program that satisfies 222 the requirements established in subdivision (1) of subsection (b) of this 223 section.

(d) If a pharmacist is morally or ethically opposed to issuing a
prescription for an emergency contraceptive or a hormonal
contraceptive, the pharmacist shall provide to any patient who requests
such a prescription a list of the nearest pharmacies that may provide
such a prescription to the patient.

(e) Each pharmacy shall maintain copies of all documents concerningany screening performed under this section for at least three years, and

each pharmacy shall, upon request by the department, make suchscreening documents available to the department for inspection.

(f) The Commissioner of Consumer Protection may adopt
regulations, in accordance with chapter 54 of the general statutes, to
implement the provisions of this section.

Sec. 4. (NEW) (*Effective from passage*) (a) For the purposes of this section, "drug", "pharmacist" and "pharmacy" have the same meanings as provided in section 20-571 of the general statutes.

(b) A pharmacist who is employed by a pharmacy that has been approved to dispense drugs for the termination of a pregnancy shall provide to any patient who is seeking any such drug a list of the pharmacies nearest to such patient that dispense such drug if (1) the pharmacy does not have a supply of such drug, or (2) the pharmacist is morally or ethically opposed to dispensing such drug to such patient.

(c) A pharmacist who is, or has been, licensed in another state or
jurisdiction shall not be subject to automatic reciprocal discipline in this
state for any disciplinary action taken in such other state or jurisdiction,
provided such disciplinary action was based solely on the termination
of a pregnancy under conditions which would not violate the laws of
this state.

251 Sec. 5. Section 20-579 of the general statutes is repealed and the 252 following is substituted in lieu thereof (*Effective from passage*):

253 (a) The commission may refuse to authorize the issuance of a 254 temporary permit to practice pharmacy, [may] refuse to authorize the 255 issuance or renewal of a license to practice pharmacy, a license to 256 operate a pharmacy or a registration of a pharmacy intern or pharmacy 257 technician, [and may] revoke, suspend or place conditions on a license 258 or temporary permit to practice pharmacy, a license to operate a 259 pharmacy [,] or a registration of a pharmacy intern or a pharmacy 260 technician [,] and [may] assess a civil penalty [of up to] not to exceed 261 one thousand dollars per violation of any provision of this chapter, or

take other action permitted in subdivision (7) of subsection (a) of section 262 263 21a-7, if the applicant or holder of the license, temporary permit or 264 registration: (1) Has violated a statute or regulation relating to drugs, 265 devices or the practice of pharmacy of this state, any state of the United 266 States, the United States, the District of Columbia, the Commonwealth 267 of Puerto Rico, any territory or insular possession subject to the 268 jurisdiction of the United States or a foreign jurisdiction; (2) has been 269 convicted of violating any criminal statute relating to drugs, devices or 270 the practice of pharmacy of this state, any state of the United States, the 271 United States, the District of Columbia, the Commonwealth of Puerto 272 Rico, any territory or insular possession subject to the jurisdiction of the 273 United States or a foreign jurisdiction; (3) has been disciplined by, or is 274 the subject of pending disciplinary action or an unresolved complaint 275 before, the duly authorized pharmacy disciplinary agency of any state 276 of the United States, the United States, the District of Columbia, the 277 Commonwealth of Puerto Rico, any territory or insular possession 278 subject to the jurisdiction of the United States or a foreign jurisdiction; 279 (4) has been refused a license or registration or renewal of a license or 280 registration by any state of the United States, the United States, the 281 District of Columbia, the Commonwealth of Puerto Rico, any territory 282 or insular possession subject to the jurisdiction of the United States or a 283 foreign jurisdiction based on grounds that are similar to grounds on 284which Connecticut could refuse to issue or renew such a license or 285 registration; (5) has illegally possessed, diverted, sold or dispensed 286 drugs or devices; (6) abuses or excessively uses drugs, including, but not 287 limited to, alcohol; (7) has made false, misleading or deceptive 288 representations to the public or the commission; (8) has maintained 289 exclusive telephone lines to, has maintained exclusive electronic 290 communication with, or has exclusive access to computers located in 291 offices of prescribing practitioners, nursing homes, clinics, hospitals or 292 other health care facilities; (9) has substituted drugs or devices except as 293 permitted in section 20-619; (10) has accepted, for return to regular stock, 294 any drug already dispensed in good faith or delivered from a pharmacy, 295 and exposed to possible and uncontrolled contamination or 296 substitution; (11) has split fees for professional services, including, but

297 not limited to, a discount or rebate, with a prescribing practitioner or an 298 administrator or owner of a nursing home, hospital or other health care 299 facility; (12) has entered into an agreement with a prescribing 300 practitioner or an administrator or owner of a nursing home, hospital or 301 other health care facility for the compounding or dispensing of secret 302 formula or coded prescriptions; (13) has performed or been a party to a fraudulent or deceitful practice or transaction; (14) has presented to the 303 304 commission a diploma, license or certificate illegally or fraudulently 305 obtained, or obtained from a college or school of pharmacy not 306 approved by the commission; (15) has performed incompetent or 307 negligent work; (16) has falsified a continuing education document 308 submitted to the commission or department or a certificate retained in 309 accordance with the provisions of subsection (d) of section 20-600; (17) 310 has permitted a person not licensed to practice pharmacy in this state to 311 practice pharmacy in violation of section 20-605, to use a pharmacist 312 license or pharmacy display document in violation of section 20-608, or 313 to use words, displays or symbols in violation of section 20-609; (18) has 314 failed to maintain the entire pharmacy premises, its components and 315 contents in a clean, orderly and sanitary condition; (19) has failed to 316 demonstrate adherence to applicable provisions of United States 317 Pharmacopeia, Chapter 797, Pharmaceutical Compounding - Sterile 318 Preparations, as amended from time to time; or (20) has failed to demonstrate adherence to applicable provisions of United States 319 320 Pharmacopeia, Chapter 795, Pharmaceutical Compounding Nonsterile Preparations, as amended from time to time. 321

322 (b) The commission may refuse to authorize the issuance or renewal 323 of a license to operate a pharmacy, revoke, suspend or place conditions 324 on a license to operate a pharmacy and assess a civil penalty not to 325 exceed one thousand dollars per violation for any violation of this 326 chapter, or take other action permitted in subdivision (7) of subsection 327 (a) of section 21a-7, if the applicant or holder of the license: (1) Implements policies, procedures, systems or processes that result in any 328 deviation from the safe practice of pharmacy; (2) prevents or delays 329 330 patient access to prescribed drugs or other pharmacy services

331 unreasonably or without providing adequate notice and an opportunity 332 to transfer such services to avoid such delay; (3) allows pharmacy 333 conditions that inhibit the safe and competent practice of pharmacy by pharmacists or other pharmacy staff, or creates an unreasonable risk to 334 335 patient care; or (4) fails to provide adequate resources, including, but 336 not limited to, staffing, to pharmacists in such a manner as to inhibit a 337 pharmacist's ability to perform all duties required under state and 338 federal law.

339 [(b)] (c) The commission may refuse to authorize the issuance of a 340 temporary permit to practice pharmacy, [may] refuse to authorize the issuance or renewal of a license to practice pharmacy, a license to 341 342 operate a pharmacy or a registration of a pharmacy intern or pharmacy 343 technician [,] and [may] revoke, suspend or place conditions on a license 344 or temporary permit to practice pharmacy, a license to operate a 345 pharmacy, or a registration of a pharmacy intern or a pharmacy 346 technician, or take other action permitted in subdivision (7) of 347 subsection (a) of section $21a-7_{L}$ if the commission determines that the applicant or holder of the license, temporary permit or registration has 348 349 a condition, including, but not limited to, physical illness or loss of skill 350 or deterioration due to the aging process, emotional disorder or mental 351 illness, abuse or excessive use of drugs or alcohol that would interfere 352 with the practice of pharmacy, operation of a pharmacy or activities as 353 a pharmacy intern or pharmacy technician, provided the commission 354 may not, in taking action against a license, temporary permit or 355 registration holder on the basis of such a condition, violate the 356 provisions of section 46a-73 or 42 USC Section 12132 of the federal 357 Americans with Disabilities Act.

Sec. 6. Subsection (d) of section 20-613 of the general statutes is repealed and the following is substituted in lieu thereof (*Effective from passage*):

361 (d) Nothing in sections 20-570 to 20-630, inclusive, shall prevent a
362 prescribing practitioner from dispensing the prescribing practitioner's
363 own prescriptions to the prescribing practitioner's own patients when

364 authorized within the scope of the prescribing practitioner's own 365 practice, [and] when done in compliance with sections 20-14c to 20-14g, 366 inclusive, and, if the prescribing practitioner is compounding, performing compounding in adherence to all applicable provisions of 367 368 Pharmacopeia, 797, United States Chapter Pharmaceutical 369 Compounding - Sterile Preparations, or Chapter 795, Pharmaceutical 370 Compounding - Nonsterile Preparations, as both may be amended from time to time. 371

Sec. 7. Section 20-617a of the general statutes is repealed and the following is substituted in lieu thereof (*Effective from passage*):

374 (a) For purposes of this section, "flavoring agent" means an additive 375 used in food or drugs when such additive [:] (1) [Is] is used in 376 accordance with good manufacturing practice principles and in the 377 minimum quantity required to produce its intended effect, (2) consists 378 of one or more ingredients generally recognized as safe in food and 379 drugs, has been previously sanctioned for use in food and drugs by the 380 state or the federal government, meets United States Pharmacopeia 381 standards or is an additive permitted for direct addition to food for 382 human consumption pursuant to 21 CFR 172, (3) is inert and produces 383 no effect other than the instillation or modification of flavor, and (4) is 384 not greater than five per cent of the total weight of the product.

(b) A flavoring agent may be added to a prescription product by [:]
(1) [A] <u>a</u> pharmacist upon the request of the prescribing practitioner,
patient for whom the prescription is ordered or such patient's agent, or
(2) a pharmacist acting on behalf of a hospital, as defined in section 19a490.

390 (c) The addition of a flavoring agent in accordance with subsections
391 (a) and (b) of this section shall be exempt from the requirements
392 established in subsections (a) to (m), inclusive, of section 20-633b, as
393 amended by this act, any regulations adopted pursuant to subsection (o)
394 of section 20-633b, as amended by this act, and United States
395 Pharmacopeia, Chapter 795, Pharmaceutical Compounding –

396 <u>Nonsterile Preparations, and Chapter 800, Hazardous Drugs, as both</u>
 397 <u>may be amended from time to time.</u>

398 Sec. 8. Section 20-623 of the general statutes is repealed and the 399 following is substituted in lieu thereof (*Effective from passage*):

400 (a) No nonlegend drug may be sold at retail except at a pharmacy, 401 [or] at a store or in a vending machine that is owned and operated by a business that has obtained from the commission or the department a 402 403 permit to sell nonlegend drugs pursuant to section 20-624. Nonlegend drugs may be sold in a vending machine, which vending machine shall 404 be owned and operated by a business that has obtained from the 405 406 department a permit for each vending machine in which such business 407 offers nonlegend drugs for sale. If an applicant seeks to locate two or more vending machines selling nonlegend drugs at a single premises, 408 409 only one permit to sell nonlegend drugs shall be required. Any person who is not licensed as a pharmacy and wishes to sell nonlegend drugs 410 411 in a vending machine shall apply to the department, in a form and 412 manner prescribed by the commissioner, in order to obtain a permit to 413 sell nonlegend drugs. Nonlegend drugs shall be labeled and packaged 414 in accordance with state and federal law.

415 (b) (1) A vending machine offering nonlegend drugs may also offer 416 nonlegend devices or test strips intended for use by an individual to test 417 for a particular substance prior to injection, inhalation or ingestion of the substance to prevent accidental overdose by injection, inhalation or 418 ingestion of such substance. Each vending machine offering nonlegend 419 drugs or nonlegend devices shall be individually registered with the 420 421 department, and each application to register a vending machine offering nonlegend drugs or nonlegend devices shall designate an individual 422 who shall be responsible for properly maintaining such vending 423 424 machine.

425 (2) Each person who registers a vending machine pursuant to
426 subdivision (1) of this subsection, and the individual designated as the
427 individual responsible for properly maintaining the registered vending

machine, shall ensure that such vending machine (A) maintains the 428 429 proper temperature and humidity for each nonlegend drug offered in such vending machine as required by the original manufacturer of such 430 nonlegend drug, (B) only contains nonlegend drugs and nonlegend 431 432 devices that remain in the original containers provided by the 433 manufacturers of such nonlegend drugs or nonlegend devices, (C) only 434 offers nonlegend drugs and nonlegend devices that are unexpired and unadulterated, (D) only offers nonlegend drugs and nonlegend devices 435 that are not subject to a recall, provided any nonlegend drug or 436 437 nonlegend device that is the subject of a recall shall be promptly 438 removed from such vending machine, (E) only contains nonlegend drugs and nonlegend devices, sundries and other nonperishable items, 439 (F) has a clear and conspicuous written statement attached to such 440 vending machine disclosing the name, address and toll-free telephone 441 442 number of the owner and operator of such vending machine, (G) has a 443 clear and conspicuous written statement attached to such vending machine advising a consumer to check the expiration date of a 444 nonlegend drug or nonlegend device contained in such vending 445 machine before the consumer uses such nonlegend drug or nonlegend 446 447 device, (H) has attached to such vending machine, in a size and prominent location visible to consumers, a written notice stating "Drug 448 449 tampering or expired product? Notify the Department of Consumer 450 Protection, Drug Control Division, by calling (telephone number of the 451 toll-free telephone line established by the department pursuant to 452 section 21a-2)", (I) does not offer any nonlegend drug or nonlegend device that requires age verification, is subject to any quantity limit or is 453 subject to any sales restriction under state or federal law, and (J) does 454 455 not contain any package of a nonlegend drug that contains more than a five-day supply of the nonlegend drug as determined according to the 456 usage directions provided by the manufacturer of such nonlegend drug. 457 [(b)] (c) Any person who violates any provision of this section shall 458 459 be fined not [less than one hundred dollars nor more than five hundred

460 dollars] more than one thousand dollars per violation.

461 Sec. 9. Section 20-633b of the general statutes is repealed and the 462 following is substituted in lieu thereof (*Effective from passage*):

463 (a) As used in this section:

(1) "Medical order" means a written, oral or electronic order by a
prescribing practitioner, as defined in section 20-14c, for a drug to be
dispensed by a pharmacy for administration to a patient;

467 (2) "Sterile compounding pharmacy" means a pharmacy, as defined
468 in section 20-571, a nonresident pharmacy registered pursuant to section
469 20-627, that dispenses or compounds sterile pharmaceuticals;

(3) "Sterile pharmaceutical" means any dosage form of a drug,
including, but not limited to, parenterals, injectables, surgical irrigants
and ophthalmics devoid of viable microorganisms; and

(4) "USP chapters" means chapters 797, 800 and 825 of the United
States Pharmacopeia that pertain to compounding sterile
pharmaceuticals and their referenced companion documents, as
amended from time to time.

477 (b) (1) If an applicant for a new pharmacy license pursuant to section 478 20-594 intends to compound sterile pharmaceuticals, the applicant shall 479 file an addendum to its pharmacy license application to include sterile 480 pharmaceutical compounding. The Department of Consumer 481 Protection shall inspect the proposed pharmacy premises of the 482 applicant and the applicant shall not compound sterile pharmaceuticals 483 until it receives notice that the addendum application has been 484 approved by the department and the Commission of Pharmacy.

(2) If an existing pharmacy licensed pursuant to section 20-594
intends to compound sterile pharmaceuticals for the first time on or
after July 1, 2014, such pharmacy shall file an addendum application to
its application on file with the department to include sterile
pharmaceutical compounding. The Department of Consumer
Protection shall inspect the pharmacy premises and the pharmacy shall

491 not compound sterile pharmaceuticals until it receives notice that such
492 addendum application has been approved by the department and the
493 Commission of Pharmacy.

494 (3) If an applicant for a nonresident pharmacy registration intends to 495 compound sterile pharmaceuticals for sale or delivery in this state, the 496 applicant shall file an addendum to its application to include sterile 497 pharmaceutical compounding. The applicant shall provide the 498 department with written proof it has passed inspection by the 499 appropriate state agency in the state where such nonresident pharmacy 500 is located. Such pharmacy shall not compound sterile pharmaceuticals 501 for sale or delivery in this state until it receives notice that the addendum 502 application has been approved by the department and the Commission 503 of Pharmacy.

504 (4) If a nonresident pharmacy registered pursuant to section 20-627 505 intends to compound sterile pharmaceuticals for sale or delivery in this 506 state for the first time on or after July 1, 2014, the nonresident pharmacy shall file an addendum to its application to include sterile 507 pharmaceutical compounding. The nonresident pharmacy shall provide 508 509 the department with written proof it has passed inspection by the 510 appropriate state agency in the state where such nonresident pharmacy 511 is located. Such pharmacy shall not compound sterile pharmaceuticals 512 until it receives notice that the addendum application has been 513 approved by the department and the Commission of Pharmacy.

(c) A sterile compounding pharmacy shall comply with the USP
chapters. A sterile compounding pharmacy shall also comply with all
applicable federal and state statutes and regulations.

(d) An institutional pharmacy within a facility licensed pursuant to
section 19a-490 that compounds sterile pharmaceuticals shall comply
with the USP chapters, and shall also comply with all applicable federal
and state statutes and regulations. Such institutional pharmacy may
request from the Commissioner of Consumer Protection an extension of
time, not to exceed six months, to comply, for state enforcement

523 purposes, with any amendments to USP chapters, for good cause 524 shown. The commissioner may grant an extension for a length of time 525 not to exceed six months. Nothing in this section shall prevent such 526 institutional pharmacy from requesting a subsequent extension of time 527 or shall prevent the commissioner from granting such extension.

(e) (1) A sterile compounding pharmacy may only provide patientspecific sterile pharmaceuticals to patients, practitioners of medicine,
osteopathy, podiatry, dentistry or veterinary medicine, or to an acute
care or long-term care hospital or health care facility licensed by the
Department of Public Health.

533 sterile compounding pharmacy (2)If а provides sterile 534 pharmaceuticals without a patient-specific prescription or medical 535 order, the sterile compounding pharmacy shall also obtain a certificate 536 of registration from the Department of Consumer Protection pursuant 537 to section 21a-70 and any required federal license or registration. A 538 sterile compounding pharmacy may prepare and maintain on-site 539 inventory of sterile pharmaceuticals no greater than a thirty-day supply, 540 calculated from the completion of compounding, which thirty-day 541 period shall include the period required for third-party analytical 542 testing, to be performed in accordance with the USP chapters.

543 (f) (1) If a sterile compounding pharmacy plans to remodel any area 544 utilized for the compounding of sterile pharmaceuticals or adjacent 545 space, relocate any space utilized for the compounding of sterile 546 pharmaceuticals or upgrade or conduct a nonemergency repair to the 547 heating, ventilation, air conditioning or primary or secondary 548 engineering controls for any space utilized for the compounding of 549 sterile pharmaceuticals, the sterile compounding pharmacy shall notify 550 the Department of Consumer Protection, in writing, not later than forty-551 five days prior to commencing such remodel, relocation, upgrade or 552 repair. Such written notification shall include a plan for such remodel, 553 relocation, upgrade or repair and such plan shall be subject to 554 department review and approval. If a sterile compounding pharmacy 555 makes an emergency repair, the sterile compounding pharmacy shall

notify the department of such emergency repair, in writing, not laterthan twenty-four hours after such repair is commenced.

(2) If the USP chapters require sterile recertification after such remodel, relocation, upgrade or repair, the sterile compounding pharmacy shall provide a copy of its sterile recertification to the Department of Consumer Protection not later than five days after the sterile recertification approval. The recertification shall only be performed by an independent licensed environmental monitoring entity.

(g) A sterile compounding pharmacy shall report, in writing, to the
Department of Consumer Protection any known violation or
noncompliance with viable and nonviable environmental sampling
testing, as defined in the USP chapters, not later than the end of the next
business day after discovering such violation or noncompliance.

570 (h) (1) If a sterile compounding pharmacy initiates a recall of sterile 571 pharmaceuticals that were dispensed pursuant to a patient-specific 572 prescription or medical order, the sterile compounding pharmacy shall 573 notify each patient or patient care giver, the prescribing practitioner and 574 the Department of Consumer Protection of such recall not later than 575 twenty-four hours after such recall was initiated.

576 (2) If a sterile compounding pharmacy initiates a recall of sterile 577 pharmaceuticals that were not dispensed pursuant to a patient-specific 578 prescription or a medical order, the sterile compounding pharmacy 579 shall notify [:] (A) [Each] each purchaser of such sterile pharmaceuticals, 580 to the extent such sterile compounding pharmacy possesses contact 581 information for each such purchaser, (B) the Department of Consumer 582 Protection, and (C) the federal Food and Drug Administration of such 583 recall not later than the end of the next business day after such recall 584 was initiated.

585 (i) Each sterile compounding pharmacy and each institutional 586 pharmacy within a facility licensed pursuant to section 19a-490 shall prepare and maintain a policy and procedure manual. The policy andprocedure manual shall comply with the USP chapters.

(j) Each sterile compounding pharmacy shall report to the
Department of Consumer Protection any administrative or legal action
commenced against it by any state or federal regulatory agency or
accreditation entity not later than five business days after receiving
notice of the commencement of such action.

594 (k) Notwithstanding the provisions of subdivisions (3) and (4) of 595 subsection (b) of this section, a sterile compounding pharmacy that is a 596 nonresident pharmacy shall provide the Department of Consumer 597 Protection proof that it has passed an inspection in such nonresident 598 pharmacy's home state, based on the USP chapters. Such nonresident 599 pharmacy shall submit to the Department of Consumer Protection a 600 copy of the most recent inspection report with its initial nonresident 601 pharmacy application and shall submit to the department a copy of its 602 most recent inspection report every two years thereafter. If the state in 603 which the nonresident pharmacy is located does not conduct 604 inspections based on standards required in the USP chapters, such 605 nonresident pharmacy shall provide satisfactory proof to the 606 department that it is in compliance with the standards required in the 607 USP chapters.

(l) A practitioner, as specified in subdivision (1) of subsection (e) of
this section, a hospital or a health care facility that receives sterile
pharmaceuticals shall report any errors related to such dispensing or
any suspected adulterated sterile pharmaceuticals to the Department of
Consumer Protection.

(m) (1) For purposes of this subsection, a "designated pharmacist"
means a pharmacist responsible for overseeing the compounding of
sterile pharmaceuticals and the application of the USP chapters, as said
chapters pertain to sterile compounding.

617 (2) Any pharmacy licensed pursuant to section 20-594 or institutional

618 pharmacy licensed pursuant to section 19a-490 that provides sterile619 pharmaceuticals shall notify the department of its designated620 pharmacist.

(3) The designated pharmacist shall be responsible for providing
proof he or she has completed a program approved by the commissioner
that demonstrates the competence necessary for the compounding of
sterile pharmaceuticals, in compliance with all applicable federal and
state statutes and regulations.

626 (4) The designated pharmacist shall immediately notify the627 department whenever he or she ceases such designation.

(5) Nothing in this section shall prevent a designated pharmacistfrom being the pharmacy manager.

(n) Notwithstanding the provisions of this section, the addition of a
flavoring agent in accordance with subsections (a) and (b) of section 20632 617a, as amended by this act, shall be exempt from the requirements of
633 United States Pharmacopeia, Chapter 795, Pharmaceutical
634 Compounding – Nonsterile Preparations, and Chapter 800, Hazardous
635 Drugs, as both may be amended from time to time.

[(n)] (o) The Commissioner of Consumer Protection may adopt
regulations, in accordance with chapter 54, to implement the provisions
of <u>subsections (a) to (n), inclusive, of</u> this section.

639 Sec. 10. Subdivision (6) of section 21a-92 of the general statutes is
640 repealed and the following is substituted in lieu thereof (*Effective from passage*):

(6) "Device", except when used in subdivision (15) of this section and
in [subsection (i)] <u>subdivision (9)</u> of section 21a-93, <u>as amended by this</u>
<u>act</u>, subdivision (6) of subsection (a) of section 21a-102, subsection (c) of
section 21a-106 and subsection (c) of section 21a-112, means
instruments, apparatus and contrivances, including their components,
parts and accessories, intended (A) for use in the diagnosis, cure,

648 mitigation, treatment or prevention of disease in humans or other 649 animals, or (B) to affect the structure or any function of the body of 650 humans or other animals;

651 Sec. 11. Section 21a-93 of the general statutes is repealed and the 652 following is substituted in lieu thereof (*Effective from passage*):

653 The following acts and the causing thereof shall be prohibited: [(a)] 654 (1) The sale in intrastate commerce of any food, drug, device or cosmetic 655 that is adulterated or misbranded; [(b)] (2) the adulteration or 656 misbranding of any food, drug, device or cosmetic in intrastate 657 commerce; [(c)] (3) the receipt in intrastate commerce of any food, drug, 658 device or cosmetic that is adulterated or misbranded, and the sale 659 thereof in such commerce for pay or otherwise; [(d)] (4) the introduction 660 or delivery for introduction into intrastate commerce of [(1)] (A) any 661 food in violation of section 21a-103 or [(2)] (B) any new drug in violation 662 of section 21a-110; [(e)] (5) the dissemination within this state, in any 663 manner or by any means or through any medium, of any false 664 advertisement; [(f)] (6) the refusal to permit [(1)] (A) entry and the taking of a sample or specimen or the making of an investigation as authorized 665 666 by section 21a-116, or [(2)] (B) access to or copying of any record as authorized by section 21a-117; [(g)] (7) the refusal to permit entry or 667 668 inspection as authorized by section 21a-118; [(h)] (8) the giving of a 669 guaranty or undertaking in intrastate commerce, referred to in 670 subsection (c) of section 21a-95, as amended by this act, that is false; [(i)] 671 (9) the forging, counterfeiting, simulating or falsely representing, or, 672 without proper authority, using, any mark, stamp, tag, label or other 673 identification device authorized or required by regulations 674 promulgated under the provisions of this chapter or of the federal act; [(i)] (10) the alteration, mutilation, destruction, obliteration or removal 675 676 of the whole or any part of the labeling of a food, drug, device or 677 cosmetic, or the doing of any other act with respect to a food, drug, 678 device or cosmetic, or the labeling or advertisement thereof, which results in a violation of this chapter; [(k)] (11) the using in interstate 679 680 commerce, in the labeling or advertisement of any drug, of any

681 representation or suggestion that an application with respect to such 682 drug is effective under Section 355 of the federal act or under section 683 21a-110, or that such drug complies with the provisions of either such 684 section; [(1)] (12) the violation of any provision of section 21a-108; [(m)] 685 (13) in the case of a prescription drug distributed or offered for sale in 686 this state, the failure of the manufacturer, packer or distributor thereof 687 to maintain for transmittal, or to transmit, to any practitioner licensed 688 by applicable state law to administer such drug who makes written 689 request for information as to such drug, true and correct copies of all 690 printed matter which is required to be included in any package in which 691 that drug is distributed or sold, or such other printed matter as is 692 approved by the commissioner or under the federal act. Nothing in this 693 [subsection] subdivision shall be construed to exempt any person from any labeling requirement imposed by or under other provisions of this 694 695 chapter unless specifically exempted under the federal act, as effective 696 on April 26, 1974; [(n)] (14) the using by any person to his own 697 advantage, or revealing, other than to the commissioner or his duly 698 authorized agents or to the courts when relevant in any judicial 699 proceeding under this chapter, of any information acquired under 700 authority of this chapter concerning any method, process, substance or 701 any other subject which as a trade secret is entitled to protection; [(0) (1)]702 (15) (A) placing or causing to be placed upon any drug or device or upon 703 the container of any drug or device, with intent to defraud, the 704 trademark, trade name or other identifying mark, imprint or device of 705 another or any likeness thereof; or [(2)] (B) selling, dispensing, disposing 706 of or causing to be sold, dispensed or disposed of or concealing or 707 keeping in possession, control or custody, with intent to sell, dispense 708 or dispose of, any drug, device or any container thereof transported, 709 received or held for transportation in commerce, with knowledge that 710 the trademark, trade name or other identifying mark, imprint or device 711 of another or any likeness thereof has been placed thereon in a manner 712 prohibited by [subdivision (1) hereof] subparagraph (A) of this 713 subdivision; or [(3)] (C) making, selling, disposing of or causing to be 714 made, sold or disposed of or keeping in possession, control or custody, 715 or concealing, with intent to defraud, any punch, die, plate, stone or

716 other thing designed to print, imprint or reproduce the trademark, trade 717 name or other identifying mark, imprint or device of another or any 718 likeness thereof upon any drug, device or container thereof; (16) failing 719 to demonstrate adherence to applicable provisions of United States 720 Pharmacopeia, Chapter 797, Pharmaceutical Compounding - Sterile 721 Preparations, as amended from time to time, concerning compounding 722 or preparation of sterile drugs; or (17) failing to demonstrate adherence 723 to applicable provisions of United States Pharmacopeia, Chapter 795, 724 Pharmaceutical Compounding - Nonsterile Preparations, as amended 725 from time to time, concerning compounding or preparation of 726 nonsterile drugs.

Sec. 12. Subsection (c) of section 21a-95 of the general statutes is
repealed and the following is substituted in lieu thereof (*Effective from passage*):

730 (c) No person shall be subject to the penalties of subsection (a) of this 731 section for having violated [subsection (a)] subdivision (1) or [(c)] (3) of 732 section 21a-93, as amended by this act, if he establishes a guaranty or 733 undertaking signed by and containing the name and address of the 734 person residing in this state from whom he received the article in good 735 faith, to the effect that such article is not adulterated or misbranded 736 within the meaning of this chapter. In such guaranty this chapter shall 737 be designated by title.

Sec. 13. Subsection (b) of section 21a-97 of the general statutes is
repealed and the following is substituted in lieu thereof (*Effective from passage*):

(b) Before any violation of this chapter, except for any violation of subdivision [(l)] (12) of section 21a-93, as amended by this act, is reported by the commissioner to any such attorney for the institution of a criminal proceeding, the person against whom such proceeding is contemplated shall be given appropriate notice and an opportunity to present his views to the commissioner, either orally or in writing, with regard to such contemplated proceeding.

748 749	Sec. 14. Section 21a-286 of the general statutes is repealed and the following is substituted in lieu thereof (<i>Effective from passage</i>):		
750	(a) For <u>the</u> purposes of this section:		
751 752	(1) "Commissioner" means the Commissioner of Consumer Protection;		
753	(2) "Department" means the Department of Consumer Protection;		
754 755 756	(3) "Host agency" means a community health organization, emergency medical service provider, government agency, law enforcement agency or local or regional board of education;		
757 758	[(1)] (<u>4)</u> "Opioid antagonist" [shall have] <u>has</u> the <u>same</u> meaning set forth in section 17a-714a; [.]		
759 760	[(2)] (<u>5)</u> "Prescribing practitioner" [shall have] <u>has</u> the <u>same</u> meaning set forth in section 20-14c; [.]		
761 762	[(3)] (<u>6)</u> "Pharmacist" [shall have] <u>has</u> the <u>same</u> meaning set forth in section 20-609a; [.]		
763	(7) "Secure box" means a container that (A) is securely affixed in a		
764	public location, (B) can be accessed by individuals for public use, (C) is		
765	temperature controlled or stored in an environment with temperature		
766	controls, (D) is tamper-resistant, (E) is equipped with an alarm capable		
767	of detecting and transmitting a signal when accessed by individuals,		
768	and (F) is equipped with an alarm capable of alerting first responders		
769	when accessed by individuals, unless equipping the container with such		
770	an alarm is commercially impracticable;		
771	(8) "Secured machine" means a device that (A) restricts access to		
772	individuals participating in a syringe services program by utilizing a		
773	designated access number, personalized magnetic strip card or any		
774	other technology to identify such individuals for the purpose of		
775	providing access, and (B) is registered with the department in a form		

776 and manner prescribed by the commissioner; and

(9) "Syringe services program" means a program that is (A)
established or authorized pursuant to section 19a-124, and (B) approved
by the department under section 21a-65.

780 (b) A prescribing practitioner, or a pharmacist who is certified to 781 prescribe [naloxone] an opioid antagonist pursuant to section 20-633c, 782 may enter into an agreement with a [law enforcement agency, 783 emergency medical service provider, government agency, community 784 health organization or local or regional board of education] host agency 785 related to the distribution and administration of an opioid antagonist 786 for the reversal of an opioid overdose. The prescribing practitioner or 787 pharmacist shall provide training to persons who will distribute or 788 administer the opioid antagonist pursuant to the terms of the 789 agreement. Persons other than the prescribing practitioner or 790 pharmacist shall receive training in the distribution or administration of 791 opioid antagonists prior to distributing or administering an opioid 792 antagonist. The agreement shall address the storage, handling, labeling, 793 recalls and recordkeeping of opioid antagonists by the [law enforcement 794 agency, emergency medical service provider, government agency, 795 community health organization or local or regional board of education 796 which] host agency that is party to the agreement.

797 (c) (1) A prescribing practitioner, or a pharmacist who is certified to 798 prescribe an opioid antagonist pursuant to section 20-633c, may enter into an agreement with a host agency to provide an intranasally or orally 799 800 administered opioid antagonist, or permit a host agency to install on the 801 host agency's premises a secure box containing an intranasally or orally 802 administered opioid antagonist. The agreement shall address the 803 environmental controls necessary to store such opioid antagonist, 804 establish procedures for replenishment of such opioid antagonist, 805 establish a process for monitoring the expiration dates of such opioid 806 antagonist and disposing of any expired opioid antagonist, and require 807 that signs be posted disclosing the presence of such opioid antagonist, 808 and usage directions for such opioid antagonist, in the language or

languages spoken in the community in which the secure box is installed. 809 810 The secure box shall not contain an amount of the opioid antagonist that 811 is greater than the amount necessary to serve the community in which 812 such secure box is installed. If the host agency is unable to maintain the 813 secure box, or the supplies necessary to maintain the secure box are 814 unavailable, such host agency shall remove such secure box, and all 815 signs required under this subdivision concerning such secure box, as soon as practicable but in no event later than five days after such host 816 817 agency discovers that such host agency is unable to maintain such secure box or the supplies necessary to maintain such secure box. 818

819 (2) A prescribing practitioner, or a pharmacist who is certified to 820 prescribe an opioid antagonist pursuant to section 20-633c, may enter 821 into an agreement with a host agency to operate a vending machine for 822 the purpose of distributing an opioid antagonist for nasal administration. The vending machine shall be in a location that 823 824 maintains a temperature that is at all times consistent with the manufacturer's package insert for the opioid antagonist, or have the 825 ability to maintain an environment, independent of the external 826 827 environment, that is appropriate for the opioid antagonist based on such 828 package insert. The following shall be clearly and conspicuously displayed on the outside of the vending machine, adjacent to the 829 830 vending machine or upon distribution of an opioid antagonist contained 831 in such vending machine: (A) Information concerning the signs and symptoms of an overdose; (B) instructions for the use of the opioid 832 833 antagonist; (C) information about the services that are offered in this state to treat opioid use disorder; and (D) an Internet web site address 834 835 that contains, or a quick response code that directs an individual to an 836 Internet web site that contains, information concerning the signs and 837 symptoms of an overdose, overdose response and instructions for the 838 use of the opioid antagonist.

839 (3) Nothing in subdivision (1) or (2) of this subsection shall be
 840 construed to prohibit placement of an opioid antagonist in a container
 841 that also includes an automated external defibrillator or any other

842 product used to treat a medical emergency.

843 (d) A prescribing practitioner, or a pharmacist who is certified to 844 prescribe an opioid antagonist pursuant to section 20-633c, may enter 845 into an agreement with a syringe services program to permit the syringe 846 services program to include an opioid antagonist in such syringe 847 services program's secured machine. The agreement shall address the 848 environmental controls necessary to store such opioid antagonist, 849 establish procedures for replenishment of such opioid antagonist, 850 establish a process for monitoring the expiration dates of such opioid antagonist and disposing of any expired opioid antagonist, and require 851 852 that signs be posted disclosing the presence of such opioid antagonist, 853 and usage directions for such opioid antagonist, in the language or 854 languages spoken in the community in which such secured machine is 855 installed.

(e) Nothing in this section shall be construed to prevent a secured
machine from distributing a test strip intended for use by an individual
prior to injection, inhalation or ingestion of a particular substance to
prevent accidental overdose by injection, inhalation or ingestion of such
substance.

861 [(c)] (f) A prescribing practitioner or pharmacist who enters into an 862 agreement pursuant to subsection (b), (c) or (d) of this section shall not 863 be liable for damages in a civil action or subject to administrative or 864 criminal prosecution for the administration or dispensing of an opioid 865 antagonist by [such law enforcement agency, emergency medical service provider, government agency, community health organization 866 867 or local or regional board of education] the host agency who is a party 868 to such agreement.

[(d)] (g) The Commissioner of Consumer Protection may adopt
regulations, in accordance with the provisions of chapter 54, to
implement the provisions of this section.

872 Sec. 15. Section 21a-408c of the general statutes is repealed and the

following is substituted in lieu thereof (*Effective from passage*):

874 (a) A physician, physician assistant or advanced practice registered 875 nurse may issue a written certification to a qualifying patient that 876 authorizes the palliative use of marijuana by the qualifying patient. Such 877 written certification shall be in the form prescribed by the Department 878 of Consumer Protection and shall include a statement signed and dated 879 by the qualifying patient's physician, physician assistant or advanced 880 practice registered nurse stating that, in such physician's, physician 881 assistant's or advanced practice registered nurse's professional opinion, 882 the qualifying patient has a debilitating medical condition and the 883 potential benefits of the palliative use of marijuana would likely 884 outweigh the health risks of such use to the qualifying patient.

885 (b) Any written certification for the palliative use of marijuana issued 886 by a physician, physician assistant or advanced practice registered 887 nurse under subsection (a) of this section shall be valid for a period not 888 to exceed one year from the date such written certification is signed and 889 dated by the physician, physician assistant or advanced practice 890 registered nurse. Not later than ten calendar days after the expiration of 891 such period, or at any time before the expiration of such period should 892 the qualifying patient no longer wish to possess marijuana for palliative 893 use, the qualifying patient or the caregiver shall destroy all usable 894 marijuana possessed by the qualifying patient and the caregiver for 895 palliative use.

896 (c) A physician, physician assistant or advanced practice registered 897 nurse shall not be subject to arrest or prosecution, penalized in any 898 manner, including, but not limited to, being subject to any civil penalty, 899 or denied any right or privilege, including, but not limited to, being 900 subject to any disciplinary action by the Connecticut Medical Examining 901 Board, the Connecticut State Board of Examiners for Nursing or other 902 professional licensing board, for providing a written certification for the 903 palliative use of marijuana under subdivision (1) of subsection (a) of 904 section 21a-408a if:

905 (1) The physician, physician assistant or advanced practice registered
906 nurse has diagnosed the qualifying patient as having a debilitating
907 medical condition;

(2) The physician, physician assistant or advanced practice registered
nurse has explained the potential risks and benefits of the palliative use
of marijuana to the qualifying patient and, if the qualifying patient lacks
legal capacity, to a parent, guardian or person having legal custody of
the qualifying patient;

(3) The written certification issued by the physician, physician assistant or advanced practice registered nurse is based upon the physician's, physician assistant's or advanced practice registered nurse's professional opinion after having completed a medically reasonable assessment of the qualifying patient's medical history and current medical condition made in the course of a bona fide health care professional-patient relationship; and

(4) The physician, physician assistant or advanced practice registered
nurse has no financial interest in a cannabis establishment, except for
retailers and delivery services, as such terms are defined in section 21a420.

924 (d) A physician assistant or nurse shall not be subject to arrest or 925 prosecution, penalized in any manner, including, but not limited to, 926 being subject to any civil penalty, or denied any right or privilege, 927 including, but not limited to, being subject to any disciplinary action by 928 the Connecticut Medical Examining Board, Board of Examiners for 929 Nursing or other professional licensing board, for administering 930 marijuana to a qualifying patient or research program subject in a 931 hospital or health care facility licensed by the Department of Public 932 Health.

(e) Notwithstanding the provisions of this section, sections 21a-408 to
21a-408b, inclusive, and sections 21a-408d to 21a-408o, inclusive, a
physician assistant or an advanced practice registered nurse shall not

936 issue a written certification to a qualifying patient when the qualifying937 patient's debilitating medical condition is glaucoma.

- (f) Notwithstanding any provision of the general statutes or any
 regulation of Connecticut state agencies concerning the certification of
 qualifying patients through telehealth services, a physician, physician
 assistant or advanced practice registered nurse may issue a written
 certification to a qualifying patient and provide any follow-up care
 utilizing telehealth services, provided all other requirements for issuing
 such written certification to the qualifying patient, including, but not
- 945 <u>limited to, all recordkeeping requirements, are satisfied.</u>

This act shall take effect as follows and shall amend the following sections:			
Section 1	January 1, 2024	New section	
Sec. 2	from passage	New section	
Sec. 3	from passage	New section	
Sec. 4	from passage	New section	
Sec. 5	from passage	20-579	
Sec. 6	from passage	20-613(d)	
Sec. 7	from passage	20-617a	
Sec. 8	from passage	20-623	
Sec. 9	from passage	20-633b	
Sec. 10	from passage	21a-92(6)	
Sec. 11	from passage	21a-93	
Sec. 12	from passage	21a-95(c)	
Sec. 13	from passage	21a-97(b)	
Sec. 14	from passage	21a-286	
Sec. 15	from passage	21a-408c	

Statement of Legislative Commissioners:

In Section 1(g), "a" was added before "dispensing assistant registration" for clarity; in Section 8(a), "that is owned and operated by a business" was added after "vending machine" for accuracy and internal consistency; and Section 8(b)(2)(D) was redrafted for clarity.

GL Joint Favorable Subst.