



General Assembly

Substitute Bill No. 8

February Session, 2024



AN ACT CONCERNING DRUG AFFORDABILITY.

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

1 Section 1. (NEW) (*Effective July 1, 2024*) For the purposes of this
2 section and sections 2 to 9, inclusive, of this act, unless the context
3 otherwise requires:

4 (1) "Canadian supplier" means a manufacturer or wholesale drug
5 distributor that is licensed or permitted under applicable Canadian law
6 to manufacture or distribute prescription drugs;

7 (2) "Canadian prescription drug importation program" or "program"
8 means the Canadian prescription drug importation program
9 established by the executive director of the Office of Health Strategy, in
10 consultation with the Commissioners of Social Services, Consumer
11 Protection and Public Health, pursuant to section 2 of this act;

12 (3) "Drug" means an article that is (A) recognized in the official United
13 States Pharmacopoeia, official Homeopathic Pharmacopoeia of the
14 United States or official National Formulary, or any supplement thereto,
15 (B) intended for use in the diagnosis, cure, mitigation, treatment or
16 prevention of disease in humans, (C) not food and intended to affect the
17 structure or any function of the human body, and (D) not a device and
18 intended for use as a component of any article specified in

19 subparagraphs (A) to (C), inclusive, of this subdivision;

20 (4) "Drug Quality and Security Act" means the federal Drug Quality
21 and Security Act, 21 USC 351, et seq., as amended from time to time;

22 (5) "Food, Drug and Cosmetic Act" means the federal Food, Drug and
23 Cosmetic Act, 21 USC 301, et seq., as amended by the Drug Quality and
24 Security Act, as both may be amended from time to time;

25 (6) "Laboratory" means an environmental laboratory as defined in
26 section 19a-29a of the general statutes that is accredited as a testing
27 laboratory in accordance with International Organization for
28 Standardization (ISO) 17025 standards;

29 (7) "Laboratory testing" means a quantitative and qualitative analysis
30 of a drug consistent with the applicable provisions of the official United
31 States Pharmacopoeia;

32 (8) "Medical assistance program" means the state's Medicaid program
33 established under Title XIX of the Social Security Act, as amended from
34 time to time, and the Children's Health Insurance Program established
35 under Title XXI of the Social Security Act, as amended from time to time;

36 (9) "Participating Canadian supplier" means a Canadian supplier that
37 is exporting prescription drugs, in the manufacturer's original
38 container, to a participating wholesaler for distribution in this state
39 under the program;

40 (10) "Participating wholesaler" means a wholesaler that is (A)
41 designated by the Department of Consumer Protection to distribute
42 prescription drugs, in the manufacturer's original container, obtained
43 from a participating Canadian supplier, and (B) participating in the
44 program;

45 (11) "Track-and-trace" means the product tracing process for the
46 components of the pharmaceutical distribution supply chain as
47 described in Title II of the Drug Quality and Security Act; and

48 (12) "Wholesaler" means a wholesaler, as defined in section 21a-70 of
49 the general statutes, that has received a certificate of registration from
50 the Commissioner of Consumer Protection pursuant to said section.

51 Sec. 2. (NEW) (*Effective July 1, 2024*) (a) The executive director of the
52 Office of Health Strategy, in consultation with the Commissioners of
53 Social Services, Consumer Protection and Public Health, shall establish
54 the "Canadian prescription drug importation program".
55 Notwithstanding any provision of the general statutes, the program
56 shall provide for the importation of safe and effective prescription drugs
57 from Canada for the medical assistance program that have the highest
58 potential for cost savings in this state as determined by the executive
59 director in consultation with said commissioners.

60 (b) (1) Not later than January 1, 2025, the executive director of the
61 Office of Health Strategy shall submit a request to the federal Food and
62 Drug Administration seeking approval for the program under Section
63 804 of the federal Food, Drug and Cosmetic Act, 21 USC 384(b) to 21
64 USC 384(h), inclusive, as amended from time to time. Such request shall,
65 at a minimum:

66 (A) Describe the state's plans for operating the program;

67 (B) Demonstrate that any prescription drug that is imported and
68 distributed in this state under the program:

69 (i) Meets all applicable federal and state standards for safety and
70 effectiveness; and

71 (ii) Complies with all federal tracing procedures; and

72 (C) Disclose the costs of implementing the program.

73 (2) (A) If the federal Food and Drug Administration approves the
74 request, the executive director of the Office of Health Strategy and the
75 Commissioners of Social Services and Consumer Protection shall:

76 (i) Submit to the Commissioner of Public Health a notice disclosing

77 that the federal Food and Drug Administration approved such request;

78 (ii) Submit to the joint standing committees of the General Assembly
79 having cognizance of matters relating to appropriations and the budgets
80 of state agencies, general law, human services and public health a notice
81 disclosing that the federal Food and Drug Administration approved
82 such request; and

83 (iii) Begin operating the program in conjunction with the
84 Commissioners of Social Services, Consumer Protection and Public
85 Health not later than one hundred eighty days after the date of such
86 approval.

87 (B) The executive director of the Office of Health Strategy shall not
88 operate the program unless the federal Food and Drug Administration
89 approved the request.

90 Sec. 3. (NEW) (*Effective July 1, 2024*) Each participating wholesaler
91 may import and distribute a prescription drug in this state for use in the
92 medical assistance program from a participating Canadian supplier
93 under the program if:

94 (1) Such drug meets the United States Food and Drug
95 Administration's standards concerning drug safety, effectiveness,
96 misbranding and adulteration;

97 (2) Importing such drug would not violate federal patent laws; and

98 (3) Such drug is not:

99 (A) A controlled substance, as defined in 21 USC 802, as amended
100 from time to time;

101 (B) A biological product, as defined in 42 USC 262, as amended from
102 time to time;

103 (C) An infused drug;

104 (D) An intravenously injected drug;

105 (E) A drug that is inhaled during surgery; or

106 (F) A drug that is a parenteral drug, the importation of which is
107 determined by the federal Secretary of Health and Human Services to
108 pose a threat to the public health.

109 Sec. 4. (NEW) (*Effective July 1, 2024*) Participating wholesalers may,
110 subject to the provisions of sections 2 to 9, inclusive, of this act, import
111 and distribute drugs in this state for use in the medical assistance
112 program from a participating Canadian supplier under the program to:

113 (1) A pharmacy or institutional pharmacy, as defined in section 20-
114 571 of the general statutes, solely for prescriptions covered under the
115 medical assistance program; and

116 (2) A laboratory registered with the Department of Public Health
117 under section 19a-29a of the general statutes to perform analytical
118 testing.

119 Sec. 5. (NEW) (*Effective July 1, 2024*) The executive director of the
120 Office of Health Strategy shall require that each participating Canadian
121 supplier and participating wholesaler (1) comply with all applicable
122 track-and-trace requirements, and shall not distribute, dispense or sell
123 outside of this state any prescription drug that is imported into this state
124 under the program, and (2) make available to the executive director all
125 track-and-trace records not later than forty-eight hours after the
126 executive director requests such records.

127 Sec. 6. (NEW) (*Effective July 1, 2024*) (a) The participating wholesaler
128 shall ensure the safety and quality of all drugs that are imported and
129 distributed in this state under the program. The participating
130 wholesaler shall:

131 (1) For each initial shipment of a drug that is imported into this state
132 by a participating wholesaler, ensure that a laboratory engaged by the
133 participating wholesaler tests a statistically valid sample size for each
134 batch of each drug in such shipment for authenticity and degradation in

135 a manner that is consistent with the Food, Drug and Cosmetic Act;

136 (2) For each shipment of a drug that is imported into this state by a
137 participating wholesaler and has been sampled and tested pursuant to
138 subdivision (1) of this subsection, ensure that a laboratory engaged by
139 the participating wholesaler tests a statistically valid sample of such
140 shipment for authenticity and degradation in a manner that is consistent
141 with the Food, Drug and Cosmetic Act;

142 (3) Certify that each drug imported into this state under the program:

143 (A) Is approved for marketing in the United States and not
144 adulterated or misbranded; and

145 (B) Meets all of the labeling requirements under 21 USC 352, as
146 amended from time to time;

147 (4) Maintain laboratory records, including, but not limited to,
148 complete data derived from all tests necessary to ensure that each drug
149 imported into this state under the program is in compliance with the
150 requirements of this section; and

151 (5) Maintain documentation demonstrating that the testing required
152 by this section was conducted at a laboratory in accordance with the
153 Food, Drug and Cosmetic Act and all other applicable federal and state
154 laws and regulations concerning laboratory qualifications.

155 (b) The participating wholesaler shall maintain all information and
156 documentation that is submitted pursuant to this section for a period of
157 not less than three years from the date of submission.

158 (c) Each participating wholesaler shall maintain all of the following
159 information for each drug that such participating wholesaler imports
160 and distributes in this state under the program, and submit such
161 information to the executive director of the Office of Health Strategy
162 upon request by the executive director:

163 (1) The name and quantity of the active ingredient of such drug;

- 164 (2) A description of the dosage form of such drug;
- 165 (3) The date on which such participating wholesaler received such
166 drug;
- 167 (4) The quantity of such drug that such participating wholesaler
168 received;
- 169 (5) The point of origin and destination of such drug;
- 170 (6) The price paid by such participating wholesaler for such drug;
- 171 (7) A report for any drug that fails laboratory testing; and
- 172 (8) Such additional information and documentation that the
173 executive director of the Office of Health Strategy deems necessary to
174 ensure the protection of the public health.

175 (d) The executive director of the Office of Health Strategy shall
176 require each participating Canadian supplier to maintain the following
177 information and documentation and, upon request by the executive
178 director, submit such information and documentation to the executive
179 director and the Commissioner of Consumer Protection for each drug
180 that such participating Canadian supplier exports into this state under
181 the program:

- 182 (1) The original source of such drug, including, but not limited to:
- 183 (A) The name of the manufacturer of such drug;
- 184 (B) The date on which such drug was manufactured; and
- 185 (C) The location where such drug was manufactured;
- 186 (2) The date on which such drug was shipped;
- 187 (3) The quantity of such drug that was shipped;
- 188 (4) The quantity of each lot of such drug originally received and the
189 source of such lot;

190 (5) The lot or control number and the batch number assigned to such
191 drug by the manufacturer; and

192 (6) Such additional information and documentation that the
193 executive director of the Office of Health Strategy, in consultation with
194 the Commissioners of Social Services, Consumer Protection and Public
195 Health, deems necessary to ensure the protection of the public health.

196 Sec. 7. (NEW) (*Effective July 1, 2024*) (a) The executive director of the
197 Office of Health Strategy shall issue a written order:

198 (1) Suspending importation and distribution of a drug under the
199 program if the executive director discovers that such distribution or
200 importation violates any provision of sections 2 to 9, inclusive, of this
201 act or any other applicable state or federal law or regulation;

202 (2) Suspending all importation and distribution of drugs by a
203 participating wholesaler under the program if the executive director
204 discovers that the participating wholesaler has violated any provision
205 of sections 2 to 9, inclusive, of this act or any other applicable state or
206 federal law or regulation;

207 (3) Suspending all importation and distribution of drugs by a
208 participating Canadian supplier under the program if the executive
209 director discovers that the participating Canadian supplier has violated
210 any provision of sections 2 to 9, inclusive, of this act or any other
211 applicable state or federal law or regulation; or

212 (4) Requiring the recall or seizure of any drug that was imported and
213 distributed under the program and has been identified as adulterated,
214 within the meaning of section 21a-105 of the general statutes, or
215 misbranded.

216 (b) The executive director of the Office of Health Strategy shall send
217 a notice to each participating Canadian supplier and participating
218 wholesaler affected by an order issued pursuant to subsection (a) of this
219 section notifying such participating Canadian supplier or participating

220 wholesaler that:

221 (1) The executive director of the Office of Health Strategy has issued
222 such order, and provide the legal and factual basis for such order; and

223 (2) Such participating Canadian supplier or participating wholesaler
224 may request, in writing, a hearing before the executive director of the
225 Office of Health Strategy, provided such request is received by the
226 executive director not later than thirty days after the date of such notice.

227 (c) If a hearing is timely requested pursuant to subsection (b) of this
228 section, the executive director of the Office of Health Strategy shall, not
229 later than thirty days after the receipt of the request, convene the hearing
230 as a contested case in accordance with the provisions of chapter 54 of
231 the general statutes. Not later than sixty days after the receipt of such
232 request, the executive director shall issue a final decision vacating,
233 modifying or affirming the order. The participating Canadian supplier
234 or participating wholesaler aggrieved by such final decision may appeal
235 such decision in accordance with the provisions of section 4-183 of the
236 general statutes.

237 Sec. 8. (NEW) (*Effective July 1, 2024*) The executive director of the
238 Office of Health Strategy may, in consultation with the Commissioners
239 of Social Services, Consumer Protection and Public Health, adopt
240 regulations in accordance with the provisions of chapter 54 of the
241 general statutes to implement the provisions of sections 2 to 9, inclusive,
242 of this act.

243 Sec. 9. (NEW) (*Effective July 1, 2024*) Not later than one hundred eighty
244 days after the program begins, and annually thereafter, the executive
245 director of the Office of Health Strategy established under section 19a-
246 754a of the general statutes shall submit a report, in accordance with the
247 provisions of section 11-4a of the general statutes, to the joint standing
248 committees of the General Assembly having cognizance of matters
249 relating to appropriations and the budgets of state agencies, general law,
250 human services and public health. Such report shall describe the
251 operations of the program established pursuant to section 2 of this act

252 and recommendations for expanding the program to other state-funded
253 and privately funded health care programs.

254 Sec. 10. (NEW) (*Effective July 1, 2024*) (a) There is established the
255 Prescription Drug Affordability Board to advise the executive director
256 of the Office of Health Strategy on decisions regarding the affordability
257 of prescription drugs. The board shall be within the Office of Health
258 Strategy for administrative purposes only.

259 (b) The purposes of the Prescription Drug Affordability Board shall
260 be to (1) explore strategies to reduce out-of-pocket drug costs to
261 consumers while supporting innovations in biotechnology and scientific
262 discovery, (2) study the prescription drug supply chain and
263 pharmaceutical pricing strategies to identify opportunities for consumer
264 savings, (3) monitor prescription drug prices in the state, (4) promote
265 innovative strategies for the use of more affordable drugs, (5) take into
266 consideration recommendations of a stakeholder council established
267 pursuant to section 11 of this act, and (6) recommend a range of options
268 of prescription drug cost affordability tools to the executive director of
269 the Office of Health Strategy.

270 (c) The board shall consist of five members, each of whom shall have
271 an advanced degree and experience or expertise in health care
272 economics, health services research, pharmacoeconomics,
273 pharmacology or clinical medicine. At least one such member shall have
274 direct experience with consumer advocacy and health equity. The
275 members shall be appointed by and serve at the pleasure of the
276 Governor with the advice and consent of either house of the General
277 Assembly. The Governor shall make all initial appointments not later
278 than January 1, 2025. Any vacancy shall be filled for the remainder of
279 the unexpired term by the Governor.

280 (d) Each member of the board shall serve a term of three years, except
281 as to the terms of the members who are first appointed to the board.
282 Two such members shall serve an initial term of three years, two such
283 members shall serve an initial term of two years and one such member

284 shall serve an initial term of one year, to be determined by the Governor.
285 The Governor may remove any appointed member of the board for
286 malfeasance in office, failure to regularly attend meetings or any cause
287 that renders the member incapable or unfit to discharge the duties of the
288 member's office. Any such removal is not subject to review.

289 (e) The Governor shall designate one member of the board to serve as
290 the chairperson of the board. Such chairperson shall schedule the first
291 meeting of the board, which shall be held not later than February 1, 2025.

292 (f) The board shall meet not less than four times annually to carry out
293 its purposes as set forth in subsection (b) of this section. A majority of
294 the board shall constitute a quorum. The concurrence of a majority of
295 the board in any matter within its powers and duties is required for any
296 determination made by the board. Any conflict of interest involving a
297 member of the board shall be disclosed at the next board meeting after
298 the conflict is identified.

299 (g) Not later than December 31, 2025, and annually thereafter, the
300 board shall report, in accordance with the provisions of section 11-4a of
301 the general statutes, to the joint standing committees of the General
302 Assembly having cognizance of matters relating to aging, general law,
303 human services, insurance and public health. The report shall include,
304 but need not be limited to: (1) Strategies for identifying and eliminating
305 pricing or business practices that do not support or enhance innovation
306 in drug development, (2) price trends and affordability strategies for
307 any drug identified pursuant to subsection (b) or (c) of section 13 of this
308 act, (3) any recommendations the board may have for legislation needed
309 to make prescription drug products more affordable in the state while
310 supporting and enhancing innovation in drug development, (4)
311 purchasing strategies, cost effectiveness evaluations and the
312 development of new technologies and drugs that increase affordability,
313 and (5) a summary and evaluation of state prescription drug advisory
314 board activities and recommendations.

315 (h) Members of the board may engage in private employment, or in

316 a profession or business, subject to any applicable laws, rules and
317 regulations of the state regarding official ethics or conflict of interest. As
318 used in this subsection, (1) "conflict of interest" means (A) an association
319 of a board member, including a financial or personal association, that
320 has the potential to bias or appear to bias a board member's decisions in
321 matters related to the board, and (B) any instance in which a board
322 member, a staff member, a contractor of the division on behalf of the
323 board or an immediate family member of a board member has received
324 or could receive (i) a financial benefit of any amount derived from the
325 results or findings of a study or determination that is reached by or for
326 the board, or (ii) a financial benefit from an individual or company that
327 owns or manufactures a prescription drug, service or item that is being
328 or will be studied by the board; and (2) "financial benefit" means
329 honoraria, fees, stock or any other form of compensation, including
330 increases to the value of existing stock holdings.

331 (i) In carrying out its purposes, the board may:

332 (1) Collect and review publicly available information and
333 information available via private subscriptions regarding prescription
334 drug pricing and business practices of health carriers, health
335 maintenance organizations, managed care organizations,
336 manufacturers, wholesale distributors and pharmacy benefit managers,
337 including, but not limited to, the annual report by pharmacy benefit
338 managers required pursuant to section 38a-479ppp of the general
339 statutes;

340 (2) Identify innovative strategies that may reduce the cost of
341 prescription drugs to consumers, including importation of certain
342 prescription drugs from Canada and other foreign countries and
343 jurisdictions;

344 (3) Identify states with innovative programs to lower prescription
345 drug costs and, if approved by the board, enter into memoranda of
346 understanding with such states to aid in the collection of transparency
347 data for prescription drug products or any other information needed to

348 establish similar programs in this state; and

349 (4) Receive and accept aid or contributions from any source of money,
350 property, labor or other things of value, to be held, used and applied to
351 carry out the purposes of the board, provided acceptance of such aid or
352 contributions does not present a conflict of interest for any board
353 member or any purpose of the board.

354 Sec. 11. (NEW) (*Effective July 1, 2024*) (a) There is established a
355 Prescription Drug Affordability Stakeholder Council to advise the
356 Prescription Drug Affordability Board established pursuant to section
357 10 of this act on decisions regarding the affordability of prescription
358 drugs.

359 (b) Members of the council shall serve for three years and shall consist
360 of:

361 (1) Three appointed by the speaker of the House of Representatives,
362 who shall be (A) a representative of a state-wide health care advocacy
363 coalition, (B) a representative of a state-wide advocacy organization for
364 elderly persons, and (C) a representative of a state-wide organization
365 for diverse communities;

366 (2) Three appointed by the president pro tempore of the Senate, who
367 shall be (A) a representative of a labor union, (B) a health services
368 researcher, and (C) a consumer who has experienced barriers to
369 obtaining prescription drugs due to the cost of such drugs;

370 (3) Two appointed by the majority leader of the House of
371 Representatives, who shall be (A) a representative of physicians, and (B)
372 a representative of nurses;

373 (4) Two appointed by the minority leader of the House of
374 Representatives, who shall be (A) a representative of private insurers,
375 and (B) a representative of brand-name drug corporations;

376 (5) Two appointed by the minority leader of the Senate, who shall be
377 (A) a representative of generic drug corporations, and (B) a

378 representative of an academic institution with expertise in health care
379 costs;

380 (6) Two appointed by the Governor, who shall be (A) a representative
381 of pharmacists, and (B) a representative of pharmacy benefit managers;

382 (7) The Secretary of the Office of Policy and Management, or the
383 secretary's designee;

384 (8) The Commissioner of Social Services, or the commissioner's
385 designee;

386 (9) The Commissioner of Public Health, or the commissioner's
387 designee;

388 (10) The Insurance Commissioner, or the commissioner's designee;

389 (11) The Commissioner of Consumer Protection, or the
390 commissioner's designee;

391 (12) The executive director of the Office of Health Strategy, or the
392 executive director's designee; and

393 (13) The Healthcare Advocate, or the Healthcare Advocate's
394 designee.

395 (c) All initial appointments to the council shall be made not later than
396 November 1, 2024. Any vacancy shall be filled by the appointing
397 authority.

398 (d) The speaker of the House of Representatives and the president
399 pro tempore of the Senate shall select the chairpersons of the council
400 from among the members of the council. Such chairpersons shall
401 schedule the first meeting of the council, which shall be held not later
402 than sixty days after the effective date of this section.

403 (e) The administrative staff of the joint standing committee of the
404 General Assembly having cognizance of matters relating to insurance
405 shall serve as administrative staff of the council.

406 (f) Not later than September 1, 2025, and annually thereafter, the
407 council shall submit a report to the board, in accordance with the
408 provisions of section 11-4a of the general statutes, on its
409 recommendations concerning prescription drug prices. The council
410 shall also provide recommendations to the board at any time the board
411 requests such recommendations.

412 Sec. 12. (NEW) (*Effective July 1, 2024*) As used in this section and
413 section 13 of this act:

414 (1) "Biologic" means a drug licensed under 42 USC 262, as amended
415 from time to time;

416 (2) "Biosimilar" means a drug that is highly similar to a biologic and
417 is produced or distributed in accordance with a biologics license
418 application approved under 42 USC 262(k), as amended from time to
419 time;

420 (3) "Board" means the Prescription Drug Affordability Board
421 established pursuant to section 10 of this act;

422 (4) "Brand-name drug" means a drug that is produced or distributed
423 in accordance with an original new drug application approved under 21
424 USC 355, as amended from time to time, but does not include an
425 authorized generic drug as defined in 42 CFR 447.502, as amended from
426 time to time;

427 (5) "FDA breakthrough drug" means a drug granted expedited
428 review by the United States Food and Drug Administration under 21
429 USC 356, as amended from time to time;

430 (6) "Generic drug" means (A) a prescription drug product that is
431 marketed or distributed in accordance with an abbreviated new drug
432 application approved under 21 USC 355, as amended from time to time,
433 (B) an authorized generic drug as defined in 42 CFR 447.502, as
434 amended from time to time, or (C) a drug that entered the market before
435 calendar year 1962 that was not originally marketed under a new

436 prescription drug product application;

437 (7) "Manufacturer" means an entity that (A) engages in the
438 manufacture of a drug product, or (B) enters into a lease with another
439 manufacturer to market and distribute a prescription drug product
440 under the entity's own name and sets or changes the wholesale
441 acquisition cost of the prescription drug product it manufactures or
442 markets;

443 (8) "Orphan drug" has the same meaning as provided in 21 CFR 316.3,
444 as amended from time to time; and

445 (9) "Prescription drug product" means a brand-name drug, a generic
446 drug, a biologic or biosimilar.

447 Sec. 13. (NEW) (*Effective July 1, 2024*) (a) To the extent practicable, the
448 Prescription Drug Affordability Board established pursuant to section
449 10 of this act may assess pricing information for prescription drug
450 products by: (1) Entering into a memorandum of understanding with
451 another state to which a manufacturer reports pricing information, (2)
452 assessing spending for the drug in the state, (3) utilizing data and
453 findings, including consumer affordability strategies, developed by
454 another state's board, (4) utilizing data and findings, including cost
455 containment strategies, developed by any other state or federal entity,
456 (5) utilizing the maximum fair price for a prescription drug for persons
457 eligible for Medicare established pursuant to the federal Inflation
458 Reduction Act of 2022, P.L. No. 117-169, as amended from time to time,
459 and (6) assessing any other available pricing information.

460 (b) On and after July 1, 2025, the board shall identify prescription
461 drug products that, as adjusted annually for inflation in accordance with
462 the consumer price index for all urban consumers published by the
463 United States Department of Labor, Bureau of Labor Statistics, are:

464 (1) Brand-name drugs that have a launch wholesale acquisition cost
465 of thirty thousand dollars or more per year or course of treatment;

466 (2) Brand-name drugs that have a wholesale acquisition cost increase
467 of three thousand dollars or more in any twelve-month period;

468 (3) Biosimilars that have a launch wholesale acquisition cost that is
469 not at least fifteen per cent lower than the referenced brand biologic at
470 the time the biosimilars are launched; and

471 (4) Generic drugs that have:

472 (A) A wholesale acquisition cost of one hundred dollars or more for
473 (i) a thirty-day supply lasting a patient for a period of thirty consecutive
474 days based on the recommended dosage approved for labeling by the
475 United States Food and Drug Administration, (ii) a supply lasting a
476 patient for fewer than thirty days based on the recommended dosage
477 approved for labeling by the United States Food and Drug
478 Administration, or (iii) one unit of the drug if the labeling approved by
479 the United States Food and Drug Administration does not recommend
480 a finite dosage; and

481 (B) A wholesale acquisition cost that increased by two hundred per
482 cent or more during the immediately preceding twelve-month period,
483 as determined by the difference between the resulting wholesale
484 acquisition cost and the average of the wholesale acquisition cost
485 reported over the immediately preceding twelve months.

486 (c) On and after July 1, 2025, the board shall identify any other
487 prescription drug products or pricing practices that may create
488 affordability challenges for the health care system in the state or
489 patients, including, but not limited to, drugs needed to address
490 significant public health priorities.

491 (d) After identifying prescription drug products as required by
492 subsections (b) and (c) of this section, the board may conduct, within
493 available appropriations, a review for any identified prescription drug
494 product or pricing practice if, after (1) seeking input from relevant
495 stakeholders, and (2) considering the average patient cost share of the
496 prescription drug product, the board determines such review is in the

497 interest of consumers.

498 (e) In conducting a review of prescription drugs, the board shall
499 examine any document and research related to the pricing of the
500 prescription drug product, including, but not limited to, (1) net average
501 price in the state, (2) market competition and context, (3) projected
502 revenue to the manufacturer, (4) the estimated value or cost
503 effectiveness, (5) whether and how the prescription drug product
504 represents an innovative therapy or is likely to improve health or health
505 outcomes for the target consumer, and (6) any rebates, discounts, patient
506 access programs or other cost mitigation strategies relevant to the
507 prescription drug product. As part of its review, the board may also
508 examine the costs or potential costs of FDA breakthrough and orphan
509 drugs.

510 (f) The board shall determine whether use of the prescription drug
511 product, consistent with the labeling approved by the federal Food and
512 Drug Administration or standard medical practice, has led or will lead
513 to affordability challenges for the health care system in the state or high
514 out-of-pocket costs for patients. In determining whether a prescription
515 drug product has led or will lead to an affordability challenge, the board
516 may consider the following factors:

517 (1) The wholesale acquisition cost for the prescription drug product
518 sold in the state;

519 (2) The average monetary price concession, discount or rebate
520 provided or expected to be provided to health plans in the state as
521 reported by manufacturers and health plans, expressed as a percentage
522 of the wholesale acquisition cost for the prescription drug product
523 under review;

524 (3) The total amount of the price concession, discount or rebate the
525 manufacturer provides to each pharmacy benefits manager operating in
526 the state for the prescription drug product under review, as reported by
527 manufacturers and pharmacy benefits managers, expressed as a
528 percentage of the wholesale acquisition costs;

529 (4) The price at which therapeutic alternatives have been sold in the
530 state;

531 (5) The average monetary concession, discount or rebate the
532 manufacturer provides or is expected to provide to health plan payors
533 and pharmacy benefits managers in the state for therapeutic
534 alternatives;

535 (6) The costs to health plans based on patient access consistent with
536 United States Food and Drug Administration labeled indications and
537 recognized standard medical practice;

538 (7) The impact on patient access resulting from the cost of the
539 prescription drug product relative to health plan benefit design;

540 (8) The current or expected dollar value of drug-specific patient
541 access programs that are supported by the manufacturer;

542 (9) The relative financial impacts to health, medical or social services
543 costs as may be quantified and compared to baseline effects of existing
544 therapeutic alternatives;

545 (10) The average patient copayment or other cost sharing for the
546 prescription drug product in the state;

547 (11) Any information a manufacturer chooses to provide; and

548 (12) Any other factors as determined by the board.

549 (g) If the board finds that the spending on a prescription drug
550 product reviewed under this section has led or will lead to an
551 affordability challenge, the board shall recommend an upper payment
552 limit to the executive director of the Office of Health Strategy and the
553 Insurance Commissioner after considering: (1) The cost of administering
554 the drug, (2) the cost of delivering the drug to patients, and (3) other
555 relevant administrative costs related to the drug. In its
556 recommendations, the board may utilize (A) upper payment limits set
557 by similar boards in other states, provided the board finds that the other

558 entity's price justification process is at least as rigorous as the process set
559 forth in state law, (B) upper payment limits set by any other state or
560 federal entity, provided the board finds that the other entity's price
561 justification process is at least as rigorous as the process set forth in state
562 law, and (C) the Medicare maximum fair price for a prescription drug.

563 Sec. 14. (NEW) (*Effective July 1, 2025*) (a) As used in this section and
564 section 15 of this act, (1) "ERISA plan" means an employee welfare
565 benefit plan subject to the Employee Retirement Income Security Act of
566 1974, as amended from time to time; (2) "health benefit plan" has the
567 same meaning as provided in section 38a-472f of the general statutes; (3)
568 "state entity" means any state agency, or any individual employed by or
569 acting on the state's behalf that purchases a prescription drug for an
570 individual with health insurance paid for by the state, including health
571 insurance offered by local, state, or federal agencies or through
572 organizations licensed in the state; and (4) "participating ERISA plan"
573 means an ERISA plan that elects to participate in the requirements of
574 this section.

575 (b) It shall be a violation of this section for a state entity or health
576 benefit plan or participating ERISA plan to purchase drugs with an
577 established upper payment limit to be dispensed or delivered to a
578 consumer in the state, whether directly or through a distributor, for a
579 cost higher than the upper payment limit as determined in subsection
580 (g) of section 13 of this act. Contracts entered into by a state entity, health
581 benefit plan or participating ERISA plan and a third party for the
582 purchase of prescription drugs shall expressly provide that rates paid
583 for drugs may not exceed the upper payment limit.

584 (c) It shall be a violation of this section for a retail pharmacy licensed
585 in this state to purchase for sale or distribution to a person whose health
586 care is provided by a state entity or health benefit plan or participating
587 ERISA plan a drug for a cost that exceeds the upper payment limit as
588 determined in subsection (g) of section 13 of this act.

589 Sec. 15. (NEW) (*Effective July 1, 2025*) Any savings generated by a state

590 entity, health benefit plan, or participating ERISA plan that are
591 attributable to the implementation of an upper payment limit
592 established by the Prescription Drug Affordability Board shall be used
593 to reduce health care costs to consumers, prioritizing the reduction of
594 out-of-pocket costs for prescription drugs. Not later than April 1, 2026,
595 and annually thereafter, each state entity, health benefit plan and
596 participating ERISA plan shall submit to the board and to the executive
597 director of the Office of Health Strategy a report describing the savings
598 achieved as a result of implementing upper payment limits and how
599 those savings were used to reduce health care costs to consumers. Not
600 later than July 1, 2026, and annually thereafter, the executive director, in
601 accordance with the provisions of section 11-4a of the general statutes,
602 shall file a report with the joint standing committees of the General
603 Assembly having cognizance of matters relating to appropriations and
604 the budgets of state agencies, general law, human services, insurance
605 and public health. The report shall include savings achieved and the
606 executive director's recommendations concerning additional savings
607 that may be achieved.

608 Sec. 16. (NEW) (*Effective July 1, 2025*) (a) As used in this section,
609 "manufacturer" means an entity that (1) engages in the manufacture of
610 a drug product, or (2) enters into a lease with another manufacturer to
611 market and distribute a prescription drug product under the entity's
612 own name and sets or changes the wholesale acquisition cost of the
613 prescription drug product it manufactures or markets. Any
614 manufacturer that intends to withdraw from sale or distribution within
615 the state a prescription drug for which the Prescription Drug
616 Affordability Board has established an upper payment limit shall
617 provide a notice of withdrawal in writing at least six months before the
618 date of the intended withdrawal of such prescription drug to the board,
619 the Insurance Commissioner, the Attorney General and any entity in the
620 state with which the manufacturer has a contract for the sale or
621 distribution of the drug.

622 (b) The board shall assess a penalty not to exceed five hundred
623 thousand dollars if the board determines that a manufacturer failed to

624 provide the notice required by subsection (a) of this section before
625 withdrawing from sale or distribution within the state a prescription
626 drug for which the board has established an upper payment limit as
627 determined in subsection (g) of section 13 of this act.

628 (c) A representative of a manufacturer that reasonably foresees an
629 impending shortage of a prescription drug it sells or distributes in the
630 state shall notify the board not later than thirty days after determining
631 that a shortage of a prescription drug is imminent.

632 Sec. 17. (NEW) (*Effective January 1, 2025*) (a) As used in this section:

633 (1) "Health benefit plan" has the same meaning as provided in section
634 38a-472f of the general statutes;

635 (2) "Insulin" means an insulin product, including, but not limited to,
636 an insulin pen or vial, that is licensed under 42 USC 262(a) or 42 USC
637 262(k), as amended from time to time;

638 (3) "Eligible insulin" means an insulin product for which at least two
639 licenses have been issued and continues to be marketed pursuant to
640 such licensure;

641 (4) "Net cost" means the cost of an insulin product taking into account
642 rebates or discounts for that specific product, excluding (A) rebates or
643 discounts required by state or federal law, including Medicaid,
644 Medicare and section 340B of the Public Health Service Act, 42 USC
645 256b, as amended from time to time, and (B) rebates or discounts related
646 to portfolio agreements that relate to purchase of multiple insulin
647 products or other drugs;

648 (5) "State entity" means any state agency, or any individual employed
649 by or acting on behalf of the state, that purchases a prescription drug for
650 an individual with health insurance paid for by the state, including
651 health insurance offered by local, state, or federal agencies or through
652 organizations licensed in the state; and

653 (6) "Wholesale acquisition cost" means the price of a medication set

654 by a pharmaceutical manufacturer in the United States when selling to
655 a wholesaler.

656 (b) A state entity and health benefit plan shall, except as otherwise
657 required in any collective bargaining agreement affecting the state
658 employee health plan established pursuant to section 5-259 of the
659 general statutes, make available in a preferred tier with no copayment
660 or out-of-pocket cost an eligible insulin product at the lowest wholesale
661 acquisition cost to a beneficiary. Notwithstanding the provisions of this
662 section, if a state entity or health plan determines that another eligible
663 insulin product has a lower net cost than the lowest wholesale
664 acquisition cost, such entity or health plan may offer that product with
665 no out-of-pocket payment to a beneficiary of such state entity or health
666 benefit plan. Nothing in this section shall prevent such entity or health
667 benefit plan from covering more than one eligible insulin product in a
668 preferred tier with no copayment or out-of-pocket cost to a beneficiary
669 of such entity or health benefit plan.

670 Sec. 18. Section 38a-492d of the general statutes is amended by adding
671 subsection (e) as follows (*Effective January 1, 2025*):

672 (NEW) (e) Notwithstanding the provisions of subsection (c) of this
673 section, on and after January 1, 2025, any policy described in subsection
674 (b) of this section shall make available in a preferred tier with no
675 copayment or out-of-pocket cost an eligible insulin product at the lowest
676 wholesale acquisition cost in accordance with section 17 of this act.

677 Sec. 19. Section 38a-518d of the general statutes is amended by adding
678 subsection (e) as follows (*Effective January 1, 2025*):

679 (NEW) (e) Notwithstanding the provisions of subsection (c) of this
680 section, on and after January 1, 2025, any policy described in subsection
681 (b) of this section shall make available in a preferred tier with no
682 copayment or out-of-pocket cost an eligible insulin product at the lowest
683 wholesale acquisition cost in accordance with section 17 of this act.

684 Sec. 20. (NEW) (*Effective July 1, 2024*) (a) As used in this section:

685 (1) "Eligible drug" means an injectable drug product approved under
686 Section 505(j) or 505(b)(2) of the federal Food, Drug and Cosmetic Act,
687 as amended from time to time, that is on the drug shortage list, or has
688 been on such list during the prior five-year period, established under
689 Section 506E of the federal Food, Drug and Cosmetic Act, 21 USC 356e,
690 as amended from time to time, or which has otherwise been identified
691 as being at risk of shortage;

692 (2) "Drug purchasing agency" means the Departments of Correction,
693 Social Services and Mental Health and Addiction Services;

694 (3) "Long-term purchase contract" means an agreement of at least two
695 years' duration that defines price and volume commitments; and

696 (4) "Hospital" means a hospital licensed pursuant to chapter 368v of
697 the general statutes.

698 (b) Any hospital or drug purchasing agency shall have a drug
699 shortage prevention strategy covering at least forty eligible drugs,
700 corresponding to at least one-third of the hospital's or agency's expected
701 utilization of each eligible drug. The hospital or agency shall ensure that
702 any long-term purchase contract for prescription drugs requires the
703 entity contracting with the hospital or agency to:

704 (1) Hold physical reserve inventory in order to buffer supply
705 disruption or demand surge equal to two quarters of contract volume,
706 unless the drug is in shortage or otherwise subject to a supply
707 disruption;

708 (2) Have a competent quality control unit and have in place processes
709 to evaluate supplier quality;

710 (3) Have a process to ensure that critical quality attributes have been
711 met and documentation of good manufacturing practices is complete;
712 and

713 (4) Participate, in accordance with federal law, in the program
714 administered under Section 340B of the Public Health Service Act, 42

715 USC 256b, as amended from time to time.

716 (c) Not later than January 1, 2025, and annually thereafter, a hospital
717 shall file a report with the Commissioner of Public Health documenting
718 compliance with the provisions of this section. Not later than February
719 1, 2025, and annually thereafter, the Commissioners of Correction,
720 Mental Health and Addiction Services, Social Services and Public
721 Health shall each file separate reports on compliance of hospitals, drug
722 purchasing agencies and their contractors, as applicable, with the
723 executive director of the Office of Health Strategy.

724 (d) The executive director of the Office of Health Strategy shall, not
725 later than April 1, 2025, and annually thereafter, file a comprehensive
726 report, in accordance with the provisions of section 11-4a of the general
727 statutes, on compliance of hospitals, drug purchasing agencies and their
728 contractors with the provisions of this section with the joint standing
729 committees of the General Assembly having cognizance of matters
730 relating to the judiciary, general law, human services and public health.

731 Sec. 21. (NEW) (*Effective from passage*) As used in this section and
732 section 22 of this act:

733 (1) "340B drug" means a drug that (A) is a covered outpatient drug
734 within the meaning of 42 USC 256b; (B) has been subject to any offer for
735 reduced prices by a manufacturer under 42 USC 256b(a)(1); and (C) is
736 purchased by a covered entity. "340B drug" includes a drug that would
737 have been purchased but for the restriction or limitation described in
738 subsection (a) of section 22 of this act;

739 (2) "Biologic" has the same meaning as provided in section 21a-70d of
740 the general statutes;

741 (3) "Covered entity" has the same meaning as provided in Section
742 340B of the Public Health Service Act, 42 USC 256b, as amended from
743 time to time;

744 (4) "Manufacturer" has the same meaning as provided in section 21a-

745 70 of the general statutes, except that such definition shall include
746 manufacturers of biologics;

747 (5) "Package" has the same meaning as provided in 21 USC
748 360eee(11)(A);

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

751 (7) "Third-party logistics provider" has the same meaning as
752 provided in section 20-571 of the general statutes; and

753 (8) "Wholesaler" or "distributor" has the same meaning as provided
754 in section 21a-70 of the general statutes.

755 Sec. 22. (NEW) (*Effective from passage*) (a) A manufacturer, third-party
756 logistics provider, wholesaler or distributor, or an agent or affiliate of
757 such manufacturer, third-party logistics provider, wholesaler or
758 distributor, shall not, either directly or indirectly:

759 (1) Deny, restrict, prohibit, discriminate against or otherwise limit the
760 acquisition of a 340B drug by, or delivery of a 340B drug to, a pharmacy
761 that is under contract with, or otherwise authorized by, a covered entity
762 to receive 340B drugs on behalf of the covered entity unless such receipt
763 is prohibited by the United States Department of Health and Human
764 Services; or

765 (2) Require a covered entity, or a pharmacy that is under contract
766 with a covered entity, to submit any claims or utilization data as a
767 condition for allowing the acquisition of a 340B drug by, or delivery of
768 a 340B drug to, a covered entity, or a pharmacy that is under contract
769 with a covered entity, unless the claims or utilization data sharing is
770 required by the United States Department of Health and Human
771 Services.

772 (b) (1) On and after July 1, 2024, if the executive director of the Office
773 of Health Strategy receives information and has a reasonable belief, after
774 evaluating such information, that any manufacturer, third-party

775 logistics provider, wholesaler or distributor, or an agent or affiliate of
776 such manufacturer, third-party logistics provider, wholesaler or
777 distributor, has acted in violation of any provision of this section, or rule
778 or regulation adopted thereunder, such manufacturer, third-party
779 logistics provider, wholesaler or distributor, or an agent or affiliate of
780 such manufacturer, third-party logistics provider, wholesaler or
781 distributor, shall be subject to a civil penalty of up to fifty thousand
782 dollars. The executive director may issue a notice of violation and civil
783 penalty by first-class mail or personal service. Such notice shall include:
784 (A) A reference to the section of the general statutes, rule or section of
785 the regulations of Connecticut state agencies believed or alleged to have
786 been violated; (B) a short and plain language statement of the matters
787 asserted or charged; (C) a description of the activity to cease; (D) a
788 statement of the amount of the civil penalty or penalties that may be
789 imposed; (E) a statement concerning the right to a hearing; and (F) a
790 statement that such manufacturer, third-party logistics provider,
791 wholesaler or distributor, or an agent or affiliate of such manufacturer,
792 third-party logistics provider, wholesaler or distributor, may, not later
793 than ten business days after receipt of such notice, make a request for a
794 hearing on the matters asserted.

795 (2) The manufacturer, third-party logistics provider, wholesaler or
796 distributor, or an agent or affiliate of such manufacturer, third-party
797 logistics provider, wholesaler or distributor, to whom such notice is
798 provided pursuant to subparagraph (A) of subdivision (1) of this
799 subsection may, not later than ten business days after receipt of such
800 notice, make written application to the Office of Health Strategy to
801 request a hearing to demonstrate that such violation did not occur. The
802 failure to make a timely request for a hearing shall result in the issuance
803 of a cease and desist order or imposition of a civil penalty by the office.
804 All hearings held under this subsection shall be conducted in
805 accordance with the provisions of chapter 54 of the general statutes.

806 (3) Following any hearing before the Office of Health Strategy
807 pursuant to subdivision (2) of this subsection, if the office finds, by a
808 preponderance of the evidence, that any manufacturer, third-party

809 logistics provider, wholesaler or distributor, or an agent or affiliate of
810 such manufacturer, third-party logistics provider, wholesaler or
811 distributor, violated or is violating any provision of this subsection, any
812 rule or regulation adopted thereunder or any order issued by the office,
813 the office shall issue a final cease and desist order in addition to any civil
814 penalty the office imposes.

815 (c) Nothing in this section shall be construed or applied to be in
816 conflict with or less restrictive than:

817 (1) Applicable federal law and related regulations, including 21 USC
818 355-1, as amended from time to time; or

819 (2) Other laws of this state to the extent such laws are compatible with
820 applicable federal law.

| | | |
|---|------------------------|-------------|
| This act shall take effect as follows and shall amend the following sections: | | |
| Section 1 | <i>July 1, 2024</i> | New section |
| Sec. 2 | <i>July 1, 2024</i> | New section |
| Sec. 3 | <i>July 1, 2024</i> | New section |
| Sec. 4 | <i>July 1, 2024</i> | New section |
| Sec. 5 | <i>July 1, 2024</i> | New section |
| Sec. 6 | <i>July 1, 2024</i> | New section |
| Sec. 7 | <i>July 1, 2024</i> | New section |
| Sec. 8 | <i>July 1, 2024</i> | New section |
| Sec. 9 | <i>July 1, 2024</i> | New section |
| Sec. 10 | <i>July 1, 2024</i> | New section |
| Sec. 11 | <i>July 1, 2024</i> | New section |
| Sec. 12 | <i>July 1, 2024</i> | New section |
| Sec. 13 | <i>July 1, 2024</i> | New section |
| Sec. 14 | <i>July 1, 2025</i> | New section |
| Sec. 15 | <i>July 1, 2025</i> | New section |
| Sec. 16 | <i>July 1, 2025</i> | New section |
| Sec. 17 | <i>January 1, 2025</i> | New section |
| Sec. 18 | <i>January 1, 2025</i> | 38a-492d(e) |
| Sec. 19 | <i>January 1, 2025</i> | 38a-518d(e) |
| Sec. 20 | <i>July 1, 2024</i> | New section |

| | | |
|---------|---------------------|-------------|
| Sec. 21 | <i>from passage</i> | New section |
| Sec. 22 | <i>from passage</i> | New section |

HS *Joint Favorable Subst.*

JUD *Joint Favorable*