

General Assembly

January Session, 2025

Offered by:

REP. GARIBAY, 60<sup>th</sup> Dist. REP. WOOD K., 29<sup>th</sup> Dist. REP. GILCHREST, 18<sup>th</sup> Dist. REP. BARRY, 31<sup>st</sup> Dist. REP. MESKERS, 150<sup>th</sup> Dist. REP. ALLIE-BRENNAN, 2<sup>nd</sup> Dist. REP. CONSTANTINE, 42<sup>nd</sup> Dist. REP. DATHAN, 142<sup>nd</sup> Dist. REP. KEITT, 134<sup>th</sup> Dist. REP. SHAKE, 120<sup>th</sup> Dist. REP. SHAKE, 120<sup>th</sup> Dist. REP. BOLINSKY, 106<sup>th</sup> Dist. REP. FORTIER, 79<sup>th</sup> Dist. Amendment

LCO No. 10453



REP. HUGHES, 135<sup>th</sup> Dist. REP. NOLAN, 39<sup>th</sup> Dist. REP. RADER, 98<sup>th</sup> Dist. REP. GAIEWSKI, 40<sup>th</sup> Dist. REP. JOHNSON, 49<sup>th</sup> Dist. REP. DELNICKI, 14<sup>th</sup> Dist. REP. MARRA T., 141<sup>st</sup> Dist. SEN. MARX, 20<sup>th</sup> Dist. SEN. HONIG, 8<sup>th</sup> Dist. SEN. GORDON, 35<sup>th</sup> Dist. SEN. LESSER, 9<sup>th</sup> Dist. SEN. CABRERA, 17<sup>th</sup> Dist.

To: Subst. House Bill No. 6870

File No. 308

Cal. No. 210

## "AN ACT ADDRESSING PATIENTS' PRESCRIPTION DRUG COSTS."

Strike everything after the enacting clause and substitute the
 following in lieu thereof:

"Section 1. (NEW) (*Effective October 1, 2025*) (a) Any pharmacy
benefits manager shall exercise good faith and fair dealing in the
performance of such pharmacy benefits manager's contractual duties to
any health carrier, as defined in section 38a-591a of the general statutes,

7 or other health benefit plan sponsor.

8 (b) Any pharmacy benefits manager shall notify the health carrier or 9 other health benefit plan sponsor, in writing, of any activity, policy or 10 practice of such pharmacy benefits manager that directly or indirectly 11 presents any conflict of interest with the duties imposed by this section.

(c) Any pharmacy benefits manager shall have an obligation of good
faith and fair dealing in performing such pharmacy benefits manager's
duties with all parties, including, but not limited to, a health carrier or
other health benefit plan sponsor with whom such pharmacy benefits
manager interacts in the performance of pharmacy benefit management
services.

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

Sec. 2. Section 38a-477cc of the general statutes is repealed and the
following is substituted in lieu thereof (*Effective January 1, 2026*):

(a) No contract for pharmacy services entered into in the state
between a health carrier, as defined in section 38a-591a, or pharmacy
benefits manager, as defined in section 38a-479aaa, and a pharmacy or
pharmacist shall:

(1) On and after January 1, 2018, contain a provision prohibiting or
penalizing, including through increased utilization review, reduced
payments or other financial disincentives, a pharmacist's disclosure to
an individual purchasing prescription medication of information
regarding:

32 (A) The cost of the prescription medication to the individual; or

(B) The availability of any therapeutically equivalent alternative
medications or alternative methods of purchasing the prescription
medication, including, but not limited to, paying a cash price, that are

less expensive than the cost of the prescription medication to theindividual; and

(2) On and after January 1, 2020, contain a provision permitting the health carrier or pharmacy benefits manager to recoup, directly or indirectly, from a pharmacy or pharmacist any portion of a claim that such health carrier or pharmacy benefits manager has paid to the pharmacy or pharmacist, unless such recoupment is permitted under section 38a-479iii or required by applicable law.

(b) (1) On and after January 1, 2018, no health carrier or pharmacy
benefits manager shall require an individual to make a payment at the
point of sale for a covered prescription medication in an amount greater
than the lesser of:

- 48 (A) The applicable copayment for such prescription medication;
- 49 (B) The allowable claim amount for the prescription medication; or

50 (C) The amount an individual would pay for the prescription 51 medication if the individual purchased the prescription medication 52 without using a health benefit plan, as defined in section 38a-591a, or 53 any other source of prescription medication benefits or discounts.

54 (2) For the purposes of this subsection, "allowable claim amount"
55 means the amount the health carrier or pharmacy benefits manager has
56 agreed to pay the pharmacy for the prescription medication.

57 (c) On and after January 1, 2026, a pharmacy benefits manager shall
58 offer a health plan the option of being charged the same price for a
59 prescription drug that such pharmacy benefits manager pays a
60 pharmacy for such prescription drug.

61 [(c)] (d) Any provision of a contract that violates the provisions of this 62 section shall be void and unenforceable. Any general business practice 63 that violates the provisions of this section shall constitute an unfair trade 64 practice pursuant to chapter 735a. The invalidity or unenforceability of

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65 66	any contract provision under this subsection shall not affect any other provision of the contract.
67	[(d)] (e) The Insurance Commissioner may:
68	(1) Enforce the provisions of this section pursuant to chapter 697; and
69 70	(2) Upon request, audit a contract for pharmacy services for compliance with the provisions of this section.
71 72	Sec. 3. Section 38a-479ttt of the general statutes is repealed and the following is substituted in lieu thereof ( <i>Effective October 1, 2025</i> ):
<ul> <li>73</li> <li>74</li> <li>75</li> <li>76</li> <li>77</li> <li>78</li> <li>79</li> <li>80</li> <li>81</li> <li>82</li> <li>83</li> <li>84</li> <li>85</li> <li>86</li> <li>87</li> <li>88</li> </ul>	Not later than March 1, 2021, and annually thereafter, the commissioner shall prepare a report, for the immediately preceding calendar year, describing the rebate practices of health carriers. The report shall contain (1) an explanation of the manner in which health carriers accounted for rebates in calculating premiums for health care plans delivered, issued for delivery, renewed, amended or continued during such year, (2) a statement disclosing whether, and describing the manner in which, health carriers made rebates available to insureds at the point of purchase during such year, (3) any other manner in which health carriers applied rebates during such year, (4) the percentage of rebate dollars used by health carriers to reduce premiums paid by insureds during such year, (5) an evaluation of rebate practices to reduce cost-sharing for health care plans delivered, issued for delivery, renewed, amended or continued during such year, and [(4)] (6) such other information as the commissioner, in the commissioner's discretion.
88 89	discretion, deems relevant for the purposes of this section. The commissioner shall publish a copy of the report on the department's
90	Internet web site.
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91 Sec. 4. (NEW) (*Effective January 1, 2026*) The Insurance Commissioner 92 shall require any health carrier, as defined in section 38a-591a of the 93 general statutes, to report to the commissioner annually on pricing in 94 effect for the prior year and profit generated between such health carrier 95 and any pharmacy benefits manager or mail-order pharmacy doing

96 business with such health carrier, provided such information is 97 reasonably available to such health carrier and any information noted 98 by such health carrier as proprietary that is reported by such health 99 carrier to the Insurance Commissioner pursuant to the provisions of this 100 section shall be kept confidential by the Insurance Commissioner, in 101 accordance with section 38a-69a of the general statutes. 102 Sec. 5. (NEW) (*Effective from passage*) (a) There is established a task 103 force to study emergency preparedness and mitigation strategies for 104 prescription drug shortages. The task force shall identify prescription 105 drugs at risk of shortage in this state and make recommendations 106 pursuant to subsection (g) of this section. 107 (b) The task force shall consist of the following members: 108 (1) Two appointed by the speaker of the House of Representatives, 109 one of whom has expertise in prescription drug supply chains and one 110 of whom has expertise in federal law concerning prescription drug 111 shortages; 112 (2) Two appointed by the president pro tempore of the Senate, one of 113 whom represents hospitals and one of whom represents health care 114 providers who treat patients with rare diseases; 115 (3) One appointed by the majority leader of the House of 116 Representatives, who represents one of the two federally recognized 117 Indian tribes in the state; 118 (4) One appointed by the majority leader of the Senate, who 119 represents one of the two federally recognized Indian tribes in the state; 120 (5) One appointed by the minority leader of the House of 121 Representatives, who represents health insurance companies; 122 (6) One appointed by the minority leader of the Senate, who is a 123 representative of the Connecticut Health Insurance Exchange; 124 (7) The Commissioner of Health Strategy, or the commissioner's

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125	designee;	
126	(8) The Commissioner of Consumer Protection, or the commissioner's	
127	designee;	
128	(9) The Commissioner of Social Services, or the commissioner's	
129	designee;	
130	(10) The Commissioner of Public Health, or the commissioner's	
131	designee;	
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132	(11) The chief executive officer of The University of Connecticut	
133	Health Center, or the chief executive officer's designee;	
134	(12) The Insurance Commissioner, or the commissioner's designee;	
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135	(13) The Commissioner of Economic and Community Development,	
136	or the commissioner's designee; and	
137	(14) Any other members as deemed necessary by the chairpersons of	
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139	(c) Any member of the task force appointed under subdivision (1),	
140	(2), (3), (4), (5) or (6) of subsection (b) of this section may be a member	
141	of the General Assembly.	
142	(d) All initial appointments to the task force shall be made not later	
143	than thirty days after the effective date of this section. Any vacancy shall	
144	be filled by the appointing authority.	
145	(e) The speaker of the House of Representatives and the president pro	
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148 140	the first meeting of the task force, which shall be held not later than sixty	
149	days after the effective date of this section.	
150	(f) The administrative staff of the joint standing committee of the	
151	General Assembly having cognizance of matters relating to general law	

152 shall serve as administrative staff of the task force.

153 (g) Not later than January 1, 2026, and annually thereafter, the task 154 force shall submit a report on its findings and recommendations to the 155 joint standing committees of the General Assembly having cognizance 156 of matters relating to general law, human services, insurance and real 157 estate and public health, in accordance with the provisions of section 11-158 4a of the general statutes, including, but not limited to, identification of 159 prescription drugs the task force determines are at risk of shortage and 160 strategies that would mitigate these shortages, including methods to 161 increase in-state production of such drugs deemed both at risk of 162 shortage and critically necessary for the provision of health care within 163 the state.

Sec. 6. (NEW) (*Effective July 1, 2025*) The Commissioner of Economic and Community Development may utilize bond proceeds pursuant to section 32-235 of the general statutes to support prescription drug production capacity in the state, provided the commissioner may give preference to financial assistance applications that incorporate recommendations by the task force established pursuant to section 5 of this act to prevent or mitigate prescription drug shortages.

171 Sec. 7. (Effective from passage) (a) Not later than July 1, 2025, the 172 chairpersons of the joint standing committee of the General Assembly 173 having cognizance of matters relating to insurance, or their designees, 174 shall convene a working group to study and make recommendations for 175 legislation related to the compensation of pharmacists licensed under 176 chapter 400j of the general statutes, who provide certain health care 177 services, including, but not limited to, vaccine administration, HIV-178 related tests, influenza-related tests and the prescribing of contraceptive 179 devices or products approved by the federal Food and Drug 180 Administration. For the purposes of this section, (1) "chain pharmacy" 181 means any community pharmacy that is publicly traded or has not less 182 than six stores located in this state, (2) "HIV-related test" and "influenza-183 related test" have the same meanings as provided in section 20-633f of 184 the general statutes, (3) "independent pharmacy" means any privately

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185	owned pharmacy that has not more than five stores located in this state,		
186	(4) "pharmacist" has the same meaning as provided in section 20-571 of		
187	the general statutes, and (5) "pharmacy benefits manager" has the same		
188	meaning as provided in section 38a-479aaa of the general statutes.		
189	(b) The working group convened pursuant to subsection (a) of this		
190	section shall consist of the following members:		
191	(1) The chairpersons of the joint standing committee of the General		
192	Assembly having cognizance of matters relating to insurance, or their		
193	designees;		
194	(2) The ranking members of the joint standing committee of the		
195	General Assembly having cognizance of matters relating to insurance,		
196	or their designees;		
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197	(3) The Insurance Commissioner, or the commissioner's designee;		
198	(4) The Commissioner of Consumer Protection, or the commissioner's		
199	designee;		
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200	(5) One pharmacist licensed under chapter 400j of the general statutes		
201	who is employed by any independent pharmacy;		
202	(6) One pharmacist licensed under chapter 400j of the general statutes		
203	who is employed by any chain pharmacy;		
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204	(7) One pharmacist licensed under chapter 400j of the general statutes		
205	who is employed by any health system pharmacy;		
206	(8) One representative of any organization representing pharmacy		
207	benefits managers;		
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208	(9) One representative of any health insurance company doing		
209	business in this state;		
210	(10) One representative of any pharmaceutical company doing		
211	business in this state;		

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212	(11) One faculty member of a school of pharmacy in this state;
213	(12) One representative of a small employer in this state that employs
214	less than fifty employees;
215 216	(13) One representative of a large employer in this state that employs more than one hundred employees;
217 218	(14) One representative of the Connecticut Health Insurance Exchange; and
219	(15) Any other members as deemed necessary by the chairpersons of
220	the joint standing committee of the General Assembly having
221	cognizance of matters relating to insurance.
222	(c) All initial appointments to the working group shall be made not
223 224	later than thirty days after the effective date of this section. Any vacancy shall be filled by the appointing authority.
225 226 227 228 229 230	(d) Working group members shall be appointed by the chairpersons of the joint standing committee of the General Assembly having cognizance of matters relating to insurance. The administrative staff of the joint standing committee of the General Assembly having cognizance of matters relating to insurance shall serve as administrative staff of the working group.
231	(e) Not later than February 1, 2026, the working group shall submit a
232	report on its findings and legislative recommendations to the joint
233 234	standing committee of the General Assembly having cognizance of matters relating to insurance, in accordance with the provisions of
235	section 11-4a of the general statutes. The working group shall terminate
236	on the date the working group submits such report or February 1, 2026,
237	whichever is later.
238	Sec. 8. (NEW) ( <i>Effective July 1, 2026</i> ) (a) As used in this section:
239	(1) "Enrollee" has the same meaning as provided in section 38a-478 of
240	the general statutes;

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241	(2) "Health benefit plan" has the same meaning as provided in section
242	38a-472f of the general statutes; and
243	(3) "Health carrier" has the same meaning as provided in section 38a-
244	591a of the general statutes.
245	(b) Each insurer, health care center, hospital service corporation,
246	medical service corporation, fraternal benefit society or other entity that
247	delivers, issues for delivery, renews, amends or continues an individual
248	or a group health insurance policy or health benefit plan in this state on
249	or after January 1, 2026, providing coverage of the type specified in
250	subdivisions (1), (2), (4), (11) and (12) of section 38a-469 of the general
251	statutes in this state, shall, when calculating an insured's or enrollee's
252	in-network liability for such insured's or enrollee's annual coinsurance,
253	copayment, deductible or other in-network out-of-pocket expense, give
254	credit for any out-of-pocket expense such insured or enrollee pays
255	directly to any pharmacy licensed pursuant to section 20-594 of the
256	general statutes, or health care provider licensed in this state, for any
257	prescription drug, provided (1) no claim for such prescription drug was
258	submitted to such insurer, center, corporation, society, or other entity,
259	and (2) such out-of-pocket expense paid by such insured or enrollee to
260	such pharmacy or health care provider is less than the average
261	discounted rate for such prescription drug paid to an in-network health
262	care provider pursuant to the terms of such health insurance policy or
263	health benefit plan.
264	(c) If any insured or enrollee purchases a prescription drug from any
265	out-of-network health care provider for a lower amount than the
266	average amount paid by such insured's or enrollee's health carrier to any
267	in-network health care provider for the same prescription drug, such
268	health carrier, when calculating such insured's or enrollee's liability for
269	such insured's or enrollee's in-network annual coinsurance, copayment,
270	deductible or other out-of-pocket expense, shall give credit for such
271	purchase, provided such insured or enrollee provides such health

272 carrier with proof of payment for such prescription drug in accordance

with the provisions of subsection (d) of this section. Nothing in this

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subsection shall be construed to restrict any health insurance policy or
health benefit plan from requiring a prior authorization or
precertification otherwise provided for in the insured's or enrollee's
health insurance policy or health benefit plan.

(d) Each health carrier shall (1) develop a proof of payment form and
publish such form on such health carrier's Internet web site for insureds
and enrollees to submit proof of payment for any out-of-network
prescription drug purchase pursuant to subsection (c) of this section,
and (2) annually provide to such insureds and enrollees written notice
of, and instructions for downloading or electronic submission of, such
proof of payment form.

285 (e) Upon receipt of any such proof of payment form submitted by any 286 insured or enrollee pursuant to subsection (d) of this section, each health 287 carrier shall give credit for any out-of-pocket payments that such 288 insured or enrollee paid to any out-of-network pharmacy or health care 289 provider in accordance with the provisions of subsection (c) of this 290 section, provided (1) the prescription drug purchased by such insured 291 or enrollee is included under such insured's or enrollee's health 292 insurance policy or health benefit plan, and (2) such insured or enrollee 293 purchased such prescription drug for a lower price than the average 294 amount paid by such insured or enrollee's health carrier to an in-295 network health care provider for the same prescription drug.

(f) The total amount credited toward any insured's or enrollee's annual coinsurance, copayment, deductible or other out-of-pocket expense pursuant to subsection (e) of this section shall not (1) exceed the total amount that such insured or enrollee is required to pay out-ofpocket under the terms of the health insurance policy or health benefit plan during a policy period, and (2) carry over to a new policy period.

Sec. 9. (*Effective July 1, 2025*) For the purposes of this section and sections 10 to 18, inclusive, of this act, unless the context otherwise requires:

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305	(1) "Canadian supplier" means a manufacturer or wholesale drug		
306	distributor that is licensed or permitted under applicable Canadian law		
307	to manufacture or distribute prescription drugs;		
308	(2) "Canadian prescription drug importation program" or "program"		
309	means a program under which the state would seek federal approval to		
310	import prescription drugs from Canada that have the highest potential		
311	for cost savings in the state;		
312	(3) "Department" means the Department of Consumer Protection;		
313	(4) "Drug" means an article that is (A) recognized in the official United		
314	States Pharmacopoeia, official Homeopathic Pharmacopoeia of the		
315	United States or official National Formulary, or any supplement thereto,		
316	(B) intended for use in the diagnosis, cure, mitigation, treatment or		
317	prevention of disease in humans, (C) not food and intended to affect the		
318	structure or any function of the human body, and (D) not a device and		
319	intended for use as a component of any article specified in		
320	subparagraphs (A) to (C), inclusive, of this subdivision;		
321	(5) "Drug Quality and Security Act" means the federal Drug Quality		
322	and Security Act, 21 USC 351, et seq., as amended from time to time;		
323	(6) "Food, Drug and Cosmetic Act" means the federal Food, Drug and		
324	Cosmetic Act, 21 USC 301, et seq., as amended by the Drug Quality and		
325	Security Act, as both may be amended from time to time;		
326	(7) "Qualifying laboratory" has the same meaning as provided in 21		
327	CFR 251.2;		
328	(8) "Laboratory testing" means a quantitative and qualitative analysis		
329	of a drug consistent with the applicable provisions of the official United		
330	States Pharmacopoeia;		
331	(9) "Participating Canadian supplier" means a Canadian supplier that		
332	is exporting prescription drugs, in the manufacturer's original		
333	container, to a participating wholesaler for distribution in this state		

334 under the program;

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

(11) "Recall" means a person's removal or correction of a marketed
product that the department determines is in violation of this section,
but "recall" does not include a market withdrawal or a stock recovery,
as such terms are defined in 21 CFR 7.3;

344 (12) "Relabeler" has the same meaning as provided in 21 CFR 207.1;

345 (13) "Repacker" has the same meaning as provided in 21 CFR 207.1;

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

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

352 Sec. 10. (Effective July 1, 2025) The Commissioner of Consumer 353 Protection shall hire, within available resources, a consultant to study 354 the feasibility of establishing a Canadian prescription drug importation 355 program to reduce prescription drug costs in the state. Not later than six 356 months after the date of final execution of a consultant contract with the 357 Department of Consumer Protection, the consultant shall recommend to 358 the commissioner whether it is more likely than not that a prescription 359 drug importation program is feasible and will result in cost savings to 360 the state. If the consultant determines such program is not likely to 361 result in a significant cost savings, the consultant shall provide a written 362 justification for such determination and may commence a feasibility 363 review of Canadian prescription drug price benchmarking; and develop

364 policy recommendations for implementing an upper payment limit for 365 prescription drugs in the state based on the Canadian price 366 benchmarking. Not later than October 1, 2027, the commissioner shall 367 file a report, in accordance with the provisions of section 11-4a of the 368 general statutes, with the joint standing committees of the General 369 Assembly having cognizance of matters relating to appropriations and 370 the budgets of state agencies, general law and human services and the 371 Office of Policy and Management on the results of the feasibility study.

372 Sec. 11. (*Effective October 1, 2027*) (a) If after completion of the study 373 described in section 10 of this act, the Commissioner of Consumer 374 Protection, in consultation with the Secretary of the Office of Policy and 375 Management, determines a Canadian prescription drug importation 376 program is feasible, the Commissioner of Consumer Protection may 377 submit a request to the federal Food and Drug Administration seeking 378 approval for the program under Section 804 of the federal Food, Drug 379 and Cosmetic Act, 21 USC 384(b) to 21 USC 384(h), inclusive, as 380 amended from time to time. If submitted, such request shall, at a 381 minimum:

(1) Describe the state's plans for operating the program and describe
any opportunities to coordinate or operate the program in coordination
with other states;

(2) Demonstrate that any prescription drug that is imported anddistributed in this state under the program would:

(A) Meet all applicable federal and state standards for safety andeffectiveness; and

- 389 (B) Comply with all federal tracing procedures; and
- 390 (3) State the estimated costs of implementing the program.

(b) If the federal Food and Drug Administration approves therequest, the Commissioner of Consumer Protection shall:

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393	(1) Submit to the Secretary of the Office of Policy and Management,	
394	and the Commissioners of Social Services and Health Strategy, a notice	
395	disclosing that the federal Food and Drug Administration approved	
396	such request; and	
397	(2) Submit to the joint standing committees of the General Assembly	
398	having cognizance of matters relating to appropriations and the budgets	
399	of state agencies, general law, human services and public health a notice	
400	disclosing that the federal Food and Drug Administration approved	
401	such request.	
402	(c) The Commissioner of Consumer Protection shall not operate the	
403	program unless the federal Food and Drug Administration approves the	
404	request. Notwithstanding the foregoing, the department may expend	
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406	implementation.	
407	Sec. 12. (Effective October 1, 2027) If the Canadian prescription drug	
408	importation program is established, each participating wholesaler may	
409	import and distribute a prescription drug in this state from a	
410	participating Canadian supplier under the program if:	
411	(1) Such drug meets the federal Food and Drug Administration's	
412	standards concerning drug safety, effectiveness, misbranding and	
413	adulteration;	
414	(2) Importing such drug would not violate federal patent laws; and	
415	(3) Such drug is not:	
416	(A) A controlled substance, as defined in 21 USC 802, as amended	
417	from time to time;	
418	(B) A biological product, as defined in 42 USC 262, as amended from	
419	time to time;	
420	(C) An infused drug;	

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421	(D) An intravenously injected drug;	
422	(E) A drug that is inhaled during surgery; or	
423 424 425	(F) A drug that is a parenteral drug, the importation of which is determined by the federal Secretary of Health and Human Services to pose a threat to the public health.	
426 427 428 429 430	Sec. 13. ( <i>Effective October 1, 2027</i> ) If a Canadian prescription drug importation program is established, participating wholesalers may, subject to the provisions of sections 9 to 12, inclusive, and sections 14 to 18, inclusive, of this act, import and distribute drugs in this state from a participating Canadian supplier under the program to:	
431 432	(1) A pharmacy or institutional pharmacy, as defined in section 20- 571 of the general statutes; and	
433	(2) A qualifying laboratory.	
434 435 436 437 438 439 440 441 442	Sec. 14. ( <i>Effective October 1, 2027</i> ) If a Canadian prescription drug importation program is established, the Commissioner of Consumer Protection shall require that each participating Canadian supplier and participating wholesaler (1) comply with all applicable track-and-trace requirements, and shall not distribute, dispense or sell outside of this state any prescription drug that is imported into this state under the program, and (2) make available to the commissioner all track-and-trace records not later than forty-eight hours after the commissioner requests such records.	
443 444 445 446 447	Sec. 15. ( <i>Effective October 1, 2027</i> ) (a) A participating wholesaler in any approved Canadian prescription drug importation program shall ensure the safety and quality of all drugs that may be imported and distributed in this state under the program. The participating wholesaler shall, if such program is established:	

448 (1) For each initial shipment of a drug that is imported into this state449 by a participating wholesaler, ensure that a qualifying laboratory

sHB 6870 Amendment 450 engaged by the participating wholesaler tests a statistically valid sample 451 size for each batch of each drug in such shipment for authenticity and 452 degradation in a manner that is consistent with the Food, Drug and 453 Cosmetic Act: 454 (2) For each shipment of a drug that is imported into this state by a 455 participating wholesaler and has been sampled and tested pursuant to 456 subdivision (1) of this subsection, ensure that a qualifying laboratory 457 engaged by the participating wholesaler tests a statistically valid sample 458 of such shipment for authenticity and degradation in a manner that is

459 consistent with the Food, Drug and Cosmetic Act;

(3) Only import drugs into this state that are (A) approved for
marketing in the United States, (B) not adulterated or misbranded, and
(C) meet all of the labeling requirements under 21 USC 352, as amended
from time to time;

(4) Maintain qualifying laboratory records, including, but not limited
to, complete data derived from all tests necessary to ensure that each
drug imported into this state under any approved Canadian
prescription drug importation program is in compliance with the
requirements of this section; and

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

(b) The participating wholesaler shall maintain all information and
documentation pursuant to this section for a period of not less than three
years from the date of submission of such information and
documentation to the participating wholesaler by a qualifying
laboratory.

479 (c) Each participating wholesaler shall maintain all of the following480 information for each drug that such participating wholesaler imports

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481 482 483	and distributes in this state under the program, and submit such information to the Commissioner of Consumer Protection upon request by the commissioner:
484	(1) The name and quantity of the active ingredient of such drug;
485	(2) A description of the dosage form of such drug;
486 487	(3) The date on which such participating wholesaler received such drug;
488 489	(4) The quantity of such drug that such participating wholesaler received;
490	(5) The point of origin and destination of such drug;
491	(6) The price paid by such participating wholesaler for such drug;
492 493	(7) A report regarding any drug that fails qualifying laboratory testing; and
494 495 496	(8) Such additional information and documentation that the commissioner deems necessary to ensure the protection of the public health.
497 498 499 500 501 502 503	(d) The Commissioner of Consumer Protection shall require each participating Canadian supplier in any approved Canadian prescription drug importation program to maintain the following information and documentation and, upon request by the commissioner, submit such information and documentation to the commissioner for each drug that such participating Canadian supplier exports into this state under the program:
504	(1) The original source of such drug, including, but not limited to:
505	(A) The name of the manufacturer of such drug;
506	(B) The date on which such drug was manufactured; and

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507	(C) The location where such drug was manufactured;
508	(2) The date on which such drug was shipped;
509	(3) The quantity of such drug that was shipped;
510 511	(4) The quantity of each lot of such drug originally received and the source of such lot;
512 513	(5) The lot or control number and the batch number assigned to such drug by the manufacturer; and
514 515 516	(6) Such additional information and documentation that the Commissioner of Consumer Protection deems necessary to ensure the protection of the public health.
517	Sec. 16. (Effective October 1, 2027) (a) If the Commissioner of Consumer
518	Protection determines that public health, safety or welfare requires
519	emergency action, the commissioner may order a participating
520	Canadian supplier, participating wholesaler, relabeler, repacker and
521	qualifying laboratory to cease and desist from actions specified in the
522	order that create the need for such emergency action pending
523	administrative proceedings. Such cease and desist order shall be (1) in
524	writing; (2) signed by the Commissioner of Consumer Protection; and
525	(3) effective upon delivery to the respondent. An administrative
526	proceeding conducted in accordance with chapter 54 of the general
527	statutes shall be promptly instituted following a cease and desist order.
528	The commissioner may impose a civil penalty, in an amount not to
529	exceed ten thousand dollars, after a hearing conducted pursuant to
530	chapter 54 of the general statutes.
531	(b) The commissioner may require the recall, embargo or destruction,
<b>E</b> 22	mumulant to partian 21. Of of the period statutes of any drug that uses

(b) The commissioner may require the recall, embargo or destruction,
pursuant to section 21a-96 of the general statutes, of any drug that was
imported and distributed under the program and has been identified as
adulterated, within the meaning of section 21a-105 of the general
statutes, or misbranded.

(c) In the event of a cease and desist, recall, embargo or destruction
order, the person adversely impacted by such order shall provide
written notice to all other businesses participating in the program,
informing them of the order.

Sec. 17. (*Effective October 1, 2027*) If a Canadian prescription drug
importation program is established, the Commissioner of Consumer
Protection may adopt regulations in accordance with the provisions of
chapter 54 of the general statutes to implement the provisions of sections
9 to 16, inclusive, and section 18 of this act.

545 Sec. 18. (Effective October 1, 2027) Not later than one hundred eighty 546 days after the first importation of any Canadian prescription drug under 547 the importation program begins, and biannually thereafter, the 548 Commissioner of Consumer Protection shall submit a report, in 549 accordance with the provisions of section 11-4a of the general statutes, 550 to the joint standing committees of the General Assembly having 551 cognizance of matters relating to appropriations and the budgets of state 552 agencies, general law, human services and public health. Such report 553 shall describe (1) the operation of the program, if established, and (2) 554 any violation of sections 9 to 17, inclusive, of this act that resulted in any 555 action taken by the commissioner pursuant to section 16 of this act and 556 the status of the investigation into such violation.

557 Sec. 19. (NEW) (Effective July 1, 2025) (a) As used in this section and sections 20 and 21 of this act, "drug purchasing agency" means the 558 559 Judicial Branch and the Department of Mental Health and Addiction Services, Children and Families, Developmental Services or Public 560 561 Health. Except as provided in subsection (c) of this section, the 562 Department of Administrative Services shall negotiate bulk prices for 563 prescription drugs on behalf of drug purchasing agencies with the goal 564 of purchasing such drugs at lower prices than the prices of such drugs 565 purchased by a single drug purchasing agency.

(b) In purchasing drugs at bulk prices pursuant to this section, a drugpurchasing agency may enter into a compact with officials in other

states to increase the state's purchasing power in negotiations withpharmaceutical companies.

(c) A drug purchasing agency may negotiate its own drug prices
upon demonstrating to the Commissioner of Administrative Services
that it is able to purchase such drugs at a cheaper price than the state's
bulk pricing agreements, or if operational conditions require. Such
demonstration shall be made in writing.

575 (d) Not later than February 1, 2026, the Commissioner of 576 Administrative Services, or the commissioner's designee, shall file a 577 report, in accordance with the provisions of section 11-4a of the general 578 statutes, with the joint standing committees of the General Assembly 579 having cognizance of matters relating to appropriations and the budgets 580 of state agencies, general law, human services and public health on any 581 savings realized from bulk purchases of prescription drugs pursuant to 582 subsection (a) of this section.

583 Sec. 20. (NEW) (Effective July 1, 2025) (a) As used in this section, (1) 584 "maximum fair prices" means the prices negotiated by the Centers for 585 Medicare and Medicaid Services for certain prescription drugs under 586 the Inflation Reduction Act, P.L. 117-69, and (2) "drug purchasing 587 agency" has the same meaning as provided in section 19 of this act. A 588 drug purchasing agency may incorporate as a guiding price in the 589 negotiations with a pharmaceutical drug manufacturer, the maximum 590 fair prices in any negotiation with a pharmaceutical drug manufacturer 591 to supply prescription drugs for health care programs subsidized by the 592 state.

(b) In purchasing drugs at bulk prices pursuant to section 19 of this act or maximum fair prices pursuant to this section, a drug purchasing agency may enter into a compact with officials in other states to increase the state's purchasing power in negotiations with pharmaceutical companies. A drug purchasing agency shall consider recommendations of the council established pursuant to section 21 of this act in any negotiations for prescription drugs pursuant to this section or section 19 600 of this act.

601 Sec. 21. (NEW) (*Effective October 1, 2025*) (a) There is established an 602 Advisory Council on Pharmaceutical Procurement to advise the 603 Commissioner of Administrative Services and drug purchasing 604 agencies on prescription drug negotiations pursuant to sections 19 and 605 20 of this act.

(b) The council shall consist of five members appointed by the
Governor, one of which shall be designated by the Governor to serve as
chairperson of the council. The council shall include members who have
expertise in health policy, health care economics or clinical medicine.

(c) All initial appointments to the council shall be made not later than
thirty days after the effective date of this section. Any vacancy shall be
filled by the appointing authority.

(d) The chairperson shall schedule the first meeting of the council,which shall be held not later than sixty days after the effective date ofthis section.

(e) No member of the council may (1) have a direct ownership or
investment interest in a pharmaceutical company, (2) be employed by
such company or participate in the management of such company, or
(3) receive or have the right to receive, directly or indirectly,
remuneration under a compensation arrangement with such company.

(f) Not later than January 1, 2026, and annually thereafter, the council
shall submit a report on its findings and recommendations to the
Commissioner of Administrative Services and the joint standing
committees of the General Assembly having cognizance of matters
relating to general law, human services and public health, in accordance
with the provisions of section 11-4a of the general statutes.

627 Sec. 22. (NEW) (*Effective from passage*) (a) Not later than thirty days 628 after the effective date of this section, the Commissioner of Social 629 Services shall petition the Secretary of the Department of Health and

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630	Human Services pursuant to 28 USC 1498, as amended from time to	
631	time, to authorize generic, lower cost forms of glucagon-like peptide	
632	(GLP-1) prescription drugs approved by the federal Food and Drug	
633	Administration to treat obesity or diabetes.	
634	(b) Upon approval of such petition, the commissioner may enter into	
635	a contract with any manufacturer of generic forms of such drugs	
636	approved by the federal Food and Drug Administration to supply such	
637	drugs to the state for use by HUSKY Health program members. The	
638	commissioner may enter into a consortium with officials in other states	
639	in contracting with such manufacturer for such drugs."	

This act shall take effect as follows and shall amend the following sections:		
Section 1	October 1, 2025	New section
Sec. 2	January 1, 2026	38a-477cc
Sec. 3	October 1, 2025	38a-479ttt
Sec. 4	January 1, 2026	New section
Sec. 5	from passage	New section
Sec. 6	July 1, 2025	New section
Sec. 7	from passage	New section
Sec. 8	July 1, 2026	New section
Sec. 9	July 1, 2025	New section
Sec. 10	July 1, 2025	New section
Sec. 11	October 1, 2027	New section
Sec. 12	October 1, 2027	New section
Sec. 13	October 1, 2027	New section
Sec. 14	October 1, 2027	New section
Sec. 15	October 1, 2027	New section
Sec. 16	October 1, 2027	New section
Sec. 17	October 1, 2027	New section
Sec. 18	October 1, 2027	New section
Sec. 19	July 1, 2025	New section
Sec. 20	July 1, 2025	New section
Sec. 21	October 1, 2025	New section
Sec. 22	from passage	New section