

General Assembly

Raised Bill No. 1006

January Session, 2021

LCO No. 3680



Referred to Committee on INSURANCE AND REAL ESTATE

Introduced by: (INS)

AN ACT CONCERNING HEALTH CARE COSTS, THE CONNECTICUT HEALTH INSURANCE EXCHANGE AND HEALTH EQUITY.

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

- 1 Section 1. Section 19a-754a of the general statutes is repealed and the
- 2 following is substituted in lieu thereof (*Effective July 1, 2021*):
- 3 (a) There is established an Office of Health Strategy, which shall be
- 4 within the Department of Public Health for administrative purposes
- 5 only. The department head of said office shall be the executive director
- 6 of the Office of Health Strategy, who shall be appointed by the Governor
- 7 in accordance with the provisions of sections 4-5 to 4-8, inclusive, with
- 8 the powers and duties therein prescribed.
- 9 (b) The Office of Health Strategy shall be responsible for the 10 following:
- 11 (1) Developing and implementing a comprehensive and cohesive
- 12 health care vision for the state, including, but not limited to, a
- 13 coordinated state health care cost containment strategy;

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(2) Promoting effective health planning and the provision of quality health care in the state in a manner that ensures access for all state residents to cost-effective health care services, avoids the duplication of such services and improves the availability and financial stability of such services throughout the state;

- (3) [Directing] (A) Developing, innovating, directing and overseeing health care delivery and payment models in the state that reduce health care cost growth and improve the quality of patient care, including, but not limited to, the State Innovation Model Initiative and related successor initiatives, (B) setting an annual health care cost growth benchmark and primary care target pursuant to section 3 of this act, (C) developing and adopting health care quality benchmarks pursuant to section 8 of this act, (D) enhancing the transparency of health care entities, as defined in section 2 of this act, (E) monitoring the development of accountable care organizations and patient-centered medical homes in the state, and (F) monitoring the adoption of alternative payment methodologies in the state;
- (4) (A) Coordinating the state's health information technology initiatives, (B) seeking funding for and overseeing the planning, implementation and development of policies and procedures for the administration of the all-payer claims database program established under section 19a-775a, (C) establishing and maintaining a consumer health information Internet web site under section 19a-755b, and (D) designating an unclassified individual from the office to perform the duties of a health information technology officer as set forth in sections 17b-59f and 17b-59g;
 - (5) Directing and overseeing the Health Systems Planning Unit established under section 19a-612 and all of its duties and responsibilities as set forth in chapter 368z; and
 - (6) Convening forums and meetings with state government and external stakeholders, including, but not limited to, the Connecticut Health Insurance Exchange, to discuss health care issues designed to

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- 46 develop effective health care cost and quality strategies.
- (c) The Office of Health Strategy shall constitute a successor, in accordance with the provisions of sections 4-38d, 4-38e and 4-39, to the
- 49 functions, powers and duties of the following:
- 50 (1) The Connecticut Health Insurance Exchange, established 51 pursuant to section 38a-1081, <u>as amended by this act</u>, relating to the 52 administration of the all-payer claims database pursuant to section 19a-53 755a; and
- (2) The Office of the Lieutenant Governor, relating to the (A) development of a chronic disease plan pursuant to section 19a-6q, (B) housing, chairing and staffing of the Health Care Cabinet pursuant to section 19a-725, and (C) (i) appointment of the health information technology officer, and (ii) oversight of the duties of such health information technology officer as set forth in sections 17b-59f and 17b-59g.
- (d) Any order or regulation of the entities listed in subdivisions (1)
 and (2) of subsection (c) of this section that is in force on July 1, 2018,
 shall continue in force and effect as an order or regulation until
 amended, repealed or superseded pursuant to law.
- Sec. 2. (NEW) (*Effective July 1, 2021*) For the purposes of this section and sections 3 to 9, inclusive, of this act:
- 67 (1) "Device manufacturer" means a manufacturer that manufactures 68 a device for which annual sales in this state exceed ten million dollars;

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(2) "Drug manufacturer" means the manufacturer of a drug that is: (A) Included in information and data submitted by a health carrier pursuant to section 38a-479qqq of the general statutes; (B) studied or listed pursuant to subsection (c) or (d) of section 19a-754b of the general statutes; or (C) in a therapeutic class of drugs that the executive director determines, through public or private reports, has had a substantial impact on prescription drug expenditures, net of rebates, as a

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76 percentage of total health care expenditures;

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- 77 (3) "Executive director" means the executive director of the office;
- 78 (4) "Health care cost growth benchmark" means the annual benchmark established pursuant to section 3 of this act;
 - (5) "Health care entity" means an accountable care organization, ambulatory surgical center, clinic, hospital or provider organization in this state, other than a health care provider contracting unit that, for a given calendar year: (A) Has a patient panel of not more than ten thousand patients; or (B) represents health care providers who collectively receive less than twenty million dollars in net patient service revenue from health carriers;
- (6) "Health care facility" has the same meaning as provided in section19a-630 of the general statutes;
- 89 (7) "Health care quality benchmark" means an annual benchmark 90 established pursuant to section 8 of this act;
- 91 (8) "Health care provider" has the same meaning as provided in 92 section 19a-17b of the general statutes;
 - (9) "Health status adjusted total medical expenses" means: (A) The total cost of care for the patient population of a provider organization with at least thirty-six thousand member months for a given calendar year, which cost (i) is calculated for such year on the basis of the allowed claims for all categories of medical expenses and all nonclaims payments for such year, including, but not limited to, cost-sharing payments, adjusted by health status and expressed on a per member, per month basis for all members in this state, (ii) is reported to the executive director for Medicaid, Medicare separately nongovernment health plans for such year, and (iii) discloses the health adjustment risk score and the version of the risk adjustment tool used to calculate such score for such provider organization for such year; and (B) the total aggregate medical expenses for all health care providers and

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- 106 provider organizations with fewer than thirty-six thousand member 107 months for a given calendar year;
- 108 (10) "Hospital outpatient department" has the same meaning as such 109 term is used in 42 CFR 413.65, as amended from time to time;
- 110 (11) "Institutional provider" means any health care provider that 111 provides skilled nursing facility services, or acute, chronic or 112 rehabilitation hospital services, in this state;
- 113 (12) "Office" means the Office of Health Strategy established under 114 section 19a-754a of the general statutes, as amended by this act;
- 115 (13) "Other entity" means a device manufacturer, drug manufacturer 116 or pharmacy benefits manager;
- 117 (14) "Payer" means a payer that, during a given calendar year, pays health care providers for health care services on behalf of, or pays 118 119 pharmacies for prescription drugs dispensed to, more than ten 120 thousand individuals in this state;
- 121 (15) "Pharmacy benefits manager" has the same meaning as provided 122 in section 38a-479000 of the general statutes;
- 123 (16) "Primary care target" means the annual target established pursuant to section 3 of this act; 124

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- (17) "Provider organization" means a group of persons, including, but not limited to, an accountable care organization, association, business trust, corporation, independent practice association, partnership, physician organization, physician-hospital organization or provider network, that is in the business of health care delivery or management in this state and represents a health care provider in contracting with a payer for payment for health care services; and
- (18) "Total health care expenditures" means the per capita sum of all health care expenditures in this state from public and private sources for a given calendar year, including: (A) All categories of medical

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- 135 expenses and all nonclaims payments to health care providers and 136 health care facilities, as included in the health status adjusted total 137 medical expenses reported, if any, by the executive director pursuant to 138 subsection (c) of section 5 of this act; (B) all patient cost-sharing 139 amounts, including, but not limited to, deductibles and copayments; (C) 140 the net cost of nongovernment health insurance; (D) prescription drug 141 expenditures net of rebates and discounts; (E) device manufacturer 142 expenditures net of rebates and discounts; and (F) any other 143 expenditures specified by the executive director.
- 144 Sec. 3. (NEW) (Effective July 1, 2021) (a) Not later than December 1, 145 2021, and annually thereafter, the executive director shall establish a 146 health care cost growth benchmark for the calendar year next 147 succeeding. Such health care cost growth benchmark shall address the 148 average growth in total health care expenditures across all payers and 149 populations in this state for such year, and the executive director shall 150 include within such health care cost growth benchmark a primary care 151 target to ensure primary care spending as a percentage of total health 152 care expenditures reaches a goal of ten per cent for the calendar year 153 beginning January 1, 2026.
- (b) In establishing each health care cost growth benchmark pursuant to subsection (a) of this section, the executive director shall, at a minimum:

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- (1) Consider any change in the consumer price index for all urban consumers in the northeast region from the preceding calendar year, and the most recent publicly available information concerning the growth rate of the gross state product;
- 161 (2) Evaluate current primary care spending as a percentage of total 162 health care expenditures; and
- 163 (3) (A) Hold an informational public hearing concerning such health 164 care cost growth benchmark:
- (i) At a time and place designated by the executive director in a notice

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166 prominently posted by the executive director on the office's Internet 167 web site;

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- (ii) In a form and manner prescribed by the executive director; and
- (iii) On the basis of the most recent report, if any, prepared by the executive director pursuant to subsection (c) of section 5 of this act, and any other information that the executive director, in the executive director's discretion, deems relevant for the purposes of such hearing.
 - (B) Notwithstanding subparagraph (A) of this subdivision, the executive director shall not be required to hold an informational public hearing concerning a health care cost growth benchmark for any calendar year beginning on or after January 1, 2023, if such health care cost growth benchmark is the same as the health care cost growth benchmark for the preceding calendar year.
 - (c) If the executive director determines, after any informational public hearing held pursuant to subdivision (3) of subsection (b) of this section, that a modification to the health care cost growth benchmark is, in the executive director's discretion, reasonably warranted, the executive director may modify such health care cost growth benchmark. The executive director need not hold an additional informational public hearing concerning such modified health care cost growth benchmark.
- (d) The executive director shall post each health care cost growth benchmark on the office's Internet web site.
- 188 (e) The executive director may enter into such contractual agreements 189 as may be necessary to carry out the purposes of this section, including, 190 but not limited to, contractual agreements with actuarial, economic and other experts and consultants to assist the executive director in 192 establishing health care cost growth benchmarks.
- 193 Sec. 4. (NEW) (Effective July 1, 2021) (a) (1) Not later than May 1, 2023, 194 and annually thereafter, the executive director shall hold an 195 informational public hearing to compare the growth in total health care

LCO No. 3680 **7** of 55 expenditures during the preceding calendar year to the health care cost growth benchmark established pursuant to section 3 of this act for such year. Such hearing shall include an examination of:

(A) The report, if any, most recently prepared by the executive director pursuant to subsection (c) of section 5 of this act;

- (B) The expenditures of health care entities and payers, including, but not limited to, health care cost trends, primary care spending as a percentage of total health care expenditures, and the factors contributing to such costs and expenditures;
- (C) Whether one category of expenditures may be offset by savings in another category of expenditures; and
- (D) Any other matters that the executive director, in the executive director's discretion, deems relevant for the purposes of this section.
 - (2) The executive director may require that any health care entity or payer that is found to be a significant contributor to health care cost growth in this state during the preceding calendar year participate in such hearing. Each such health care entity or payer that is required to participate in such hearing shall provide testimony on issues identified by the executive director, and provide additional information on actions taken to reduce such health care entity's contribution to future statewide health care costs and expenditures.
 - (b) Not later than October 1, 2023, and annually thereafter, the executive director shall prepare and submit a report, in accordance with section 11-4a of the general statutes, to the joint standing committees of the General Assembly having cognizance of matters relating to insurance and public health. Such report shall be based on the executive director's analysis of the information submitted during the most recent informational public hearing conducted pursuant to subsection (a) of this section and any other information that the executive director, in the executive director's discretion, deems relevant for the purposes of this section, and shall:

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- (1) Describe health care spending trends in this state, including, but not limited to, trends in primary care spending as a percentage of total health care expenditures, and the factors underlying such trends; and
- 230 (2) Disclose the executive director's recommendations, if any, 231 concerning strategies to increase the efficiency of this state's health care 232 system, including, but not limited to, any recommended legislation 233 concerning this state's health care system.
 - Sec. 5. (NEW) (*Effective July 1, 2021*) (a) Not later than March 1, 2023, and annually thereafter, each institutional provider, on behalf of such institutional provider and its parent organization and affiliated entities, health care provider that is not an institutional provider and provider organization in this state, shall submit to the executive director, for the preceding calendar year:
- 240 (1) Data concerning:

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- (A) The utilization of health care services provided by such provider or organization;
- (B) The charges, prices imposed and payments received by such provider or organization for such services;
- 245 (C) The costs incurred, and revenues earned, by such provider or organization in providing such services; and
- (D) Any other matter that the executive director deems relevant for the purposes of this section; and
 - (2) If such provider is a hospital, the data described in subdivision (1) of this subsection, and such additional data, information and documents designated by the executive director, including, but not limited to, charge masters, cost data, audited financial statements and merged billing and discharge data, provided such provider shall not be required to submit any data contained in a report that is filed pursuant to chapters 368aa to 368*ll*, inclusive, of the general statutes and available to the executive director.

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- (b) The executive director shall establish standards to ensure that the data, information and documents submitted to the executive director pursuant to subsection (a) of this section are submitted to the executive director in a uniform manner. Such standards shall enable the executive director to identify, on a patient-centered and health care providerspecific basis, state-wide and regional trends in the availability, cost, price and utilization of medical, surgical, diagnostic and ancillary services and prescription drugs provided by hospital outpatient departments, acute care hospitals, chronic disease hospitals, rehabilitation hospitals and other specialty hospitals, clinics, including, but not limited to, psychiatric clinics, urgent care facilities and facilities providing ambulatory care. Such standards may require hospitals to submit such data, information and documents to the executive director in an electronic form, provided such standards shall provide for a waiver of such requirement if such waiver is reasonable in the judgment of the executive director.
- (c) (1) Not later than December 1, 2022, and annually thereafter, the executive director shall prepare, to the extent practicable, and post on the office's Internet web site, a report concerning health status adjusted total medical expenses for the preceding calendar year, including, but not limited to, a breakdown of such health status adjusted total medical expenses by:
- 279 (A) Major service category;
- 280 (B) Payment methodology;
- 281 (C) Relative price;

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- 282 (D) Direct hospital inpatient cost;
- 283 (E) Indirect hospital inpatient cost;
- 284 (F) Direct hospital outpatient cost;
- 285 (G) Indirect hospital outpatient cost; and

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(H) Primary care spending as a percentage of total health care expenditures.

- (2) Notwithstanding subdivision (1) of this subsection, the executive director shall not disclose any health care provider-specific data or information unless the executive director provides at least ten days' advance written notice of such disclosure to each health care provider that would be affected by such disclosure.
 - (d) The executive director shall, at least annually, submit a request to the federal Centers for Medicare and Medicaid Services for the health status adjusted total medical expenses of provider organizations that served Medicare patients during the calendar year next preceding.
 - (e) The executive director may enter into such contractual agreements as may be necessary to carry out the purposes of this section, including, but not limited to, contractual agreements with actuarial, economic and other experts and consultants.
 - Sec. 6. (NEW) (Effective July 1, 2021) (a) (1) For each calendar year beginning on or after January 1, 2023, if the executive director determines that the average annual percentage change in total health care expenditures for the preceding calendar year exceeded the health care cost growth benchmark for such year, the executive director shall identify, not later than May first of such calendar year, each health care entity or payer that exceeded such health care cost growth benchmark for such year.
 - (2) The executive director may require any health care entity or payer that is found to be a significant contributor to health care cost growth in this state during the preceding calendar year to participate in the informational public hearing held pursuant to subsection (a) of section 4 of this act. Each such entity or payer that is required to participate in such hearing shall provide testimony on issues identified by the executive director, and provide additional information on actions taken to reduce such entity's or payer's contribution to future state-wide health care costs.

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(b) Not later than thirty days after the executive director identifies each health care entity or payer pursuant to subdivision (1) of subsection (a) of this section, the executive director shall send a notice to each such entity or payer. Such notice shall be in a form and manner prescribed by the executive director, and disclose to each such entity or payer:

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- (1) That the executive director has identified such entity or payer pursuant to subdivision (1) of subsection (a) of this section;
- 325 (2) The factual basis for the executive director's identification of such 326 entity or payer pursuant to subdivision (1) of subsection (a) of this 327 section; and
 - (3) That such entity or payer shall file a proposed performance improvement plan pursuant to subdivision (1) of subsection (e) of this section, provided such entity or payer may:
- (A) File a request for an extension of time, or a waiver, pursuant to subdivision (1) of subsection (c) of this section; and
- (B) Request a hearing pursuant to subsection (d) of this section.
- (c) (1) (A) Each health care entity or payer identified by the executive director pursuant to subdivision (1) of subsection (a) of this section may, not later than thirty days after the executive director sends a notice to such entity or payer pursuant to subsection (b) of this section, file with the executive director, in a form and manner prescribed by the executive director, a request seeking:
- 340 (i) An extension of time to file a proposed performance improvement 341 plan pursuant to subdivision (1) of subsection (e) of this section; or
- 342 (ii) A waiver from the requirement that such entity or payer file a 343 proposed performance improvement plan pursuant to subdivision (1) 344 of subsection (e) of this section.
 - (B) Each health care entity or payer that files a request pursuant to subparagraph (A) of this subdivision shall set forth in such request the

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347 reasons for such request.

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- 348 (2) Not later than thirty days after a health care entity or payer files a 349 request pursuant to subdivision (1) of this subsection, the executive 350 director shall:
 - (A) Examine the reasons set forth in the request and decide, on the basis of such reasons, whether to approve or deny such request; and
- 353 (B) Send a notice, in a form and manner prescribed by the executive 354 director, to the entity or payer that filed such request disclosing, at a 355 minimum:
- 356 (i) The executive director's decision concerning such request and the reasons therefor:
 - (ii) If the executive director denies such entity's or payer's request, that such entity or payer may file a request for a hearing pursuant to subsection (d) of this section; and
 - (iii) If such entity's or payer's request is a request for an extension of time to file a proposed performance improvement plan pursuant to subdivision (1) of subsection (e) of this section and the executive director approves such request, the date by which such entity or payer shall file such proposed performance improvement plan.
 - (d) Each health care entity or payer identified by the executive director pursuant to subsection (a) of this section may, not later than thirty days after the executive director sends a notice to such entity or payer pursuant to subsection (b) of this section or subparagraph (B) of subdivision (2) of subsection (c) of this section, as applicable, file with the executive director a request for a hearing. Each hearing conducted pursuant to this subsection shall be conducted in accordance with the procedures for hearings on contested cases established in chapter 54 of the general statutes.
 - (e) (1) Each health care entity or payer identified by the executive director pursuant to subdivision (1) of subsection (a) of this section, or

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377 required by the executive director pursuant to subparagraph (C)(ii)(III) 378 of subdivision (4) of subsection (f) of this section, shall, subject to the 379 provisions of subsections (b) to (d), inclusive, of this section, file with 380 the executive director a proposed performance improvement plan. Such 381 entity or payer shall file such proposed performance improvement plan, 382 which shall include an implementation timetable, with the executive 383 director, in a form and manner prescribed by the executive director, not 384 later than whichever of the following dates first occurs:

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- (A) The date that is thirty days after the date on which the executive director sent a notice to such entity or payer pursuant to subsection (b) of this section;
- (B) The date that the executive director disclosed to such entity or payer pursuant to subparagraph (B)(iii) of subdivision (2) of subsection (c) of this section; or
- (C) The date that is thirty days after the date on which the notice of a final decision is issued following a hearing conducted pursuant to subsection (d) of this section.
- (2) (A) The executive director shall review each health care entity's and payer's proposed performance improvement plan filed pursuant to subdivision (1) of this subsection to determine whether, in the executive director's judgment, it is reasonably likely that:
- 398 (i) Such proposed performance improvement plan will address the 399 cause of such entity's or payer's excessive cost growth; and
- 400 (ii) Such entity or payer will successfully implement such proposed 401 performance improvement plan.
- (B) After the executive director reviews a proposed performance improvement plan pursuant to subparagraph (A) of this subdivision, the executive director shall:
 - (i) Approve such proposed performance improvement plan if the executive director determines, in the executive director's judgment, that

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such proposed plan satisfies the criteria established in subparagraph (A) of this subdivision; or

- (ii) Deny such proposed performance improvement plan if the executive director determines, in the executive director's judgment, that such proposed performance improvement plan does not satisfy the criteria established in subparagraph (A) of this subdivision.
- (C) (i) Not later than thirty days after the executive director approves or denies a proposed performance improvement plan pursuant to subparagraph (B) of this subdivision, the executive director shall send a notice to the health care entity or payer that filed such proposed performance improvement plan disclosing, at a minimum, that:
- 418 (I) The executive director approved such proposed performance 419 improvement plan; or
 - (II) The executive director denied such proposed performance improvement plan, the reasons for such denial and that such entity or payer shall file with the executive director such amendments as are necessary for such proposed performance improvement plan to satisfy the criteria established in subparagraph (A) of this subdivision.
 - (ii) The executive director shall post a notice on the office's Internet web site disclosing:
- 427 (I) The name of each health care entity or payer that files, and receives 428 approval for, a proposed performance improvement plan; and
- 429 (II) That such health care entity or payer is implementing such 430 performance improvement plan.
 - (D) Each health care entity or payer that receives a notice from the executive director pursuant to subparagraph (C)(i) of this subdivision notifying such entity or payer that the executive director has denied such entity's or payer's proposed performance improvement plan shall file with the executive director, in a form and manner prescribed by the executive director and not later than thirty days after the date that the

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- executive director sends such notice to such entity or payer, such amendments as are necessary for such proposed performance improvement plan to satisfy the criteria established in subparagraph (A)
- 440 of this subdivision.

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- (f) (1) Each health care entity or payer that receives a notice from the executive director pursuant to subparagraph (C)(i) of subdivision (2) of subsection (e) of this section notifying such entity or payer that the executive director has approved such entity's or payer's proposed performance improvement plan:
- (A) Shall immediately make good faith efforts to implement such performance improvement plan; and
 - (B) May amend such plan at any time during the implementation timetable included in such performance improvement plan, provided the executive director approves such amendment.
 - (2) The office may provide such assistance to each health care entity or payer that the executive director, in the executive director's discretion, deems necessary and appropriate to ensure that such entity or payer successfully implements such entity's or payer's performance improvement plan.
 - (3) Each health care entity or payer shall be subject to such additional reporting requirements that the executive director, in the executive director's discretion, deems necessary to ensure that such entity or payer successfully implements such entity's or payer's performance improvement plan.
 - (4) (A) Each health care entity or payer that files, and receives approval for, a performance improvement plan pursuant to this section shall, not later than thirty days after the last date specified in the implementation timetable included in such performance improvement plan, submit to the executive director, in a form and manner prescribed by the executive director, a report regarding the outcome of such entity's or payer's implementation of such performance improvement plan.

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- (B) If the executive director determines, on the basis of the report submitted by a health care entity or payer pursuant to subparagraph (A) of this subdivision, that such entity or payer successfully implemented such entity's or payer's performance improvement plan, the executive director shall:
- 473 (i) Send a notice to such entity or payer, in a form and manner 474 prescribed by the executive director, disclosing such determination; and
- (ii) Remove from the office's Internet web site the notice concerning such entity or payer that the executive director posted on such Internet web site pursuant to subparagraph (C)(ii) of subdivision (2) of subsection (e) of this section.
 - (C) If the executive director determines, on the basis of the report submitted by a health care entity or payer pursuant to subparagraph (A) of this subdivision, that such entity or payer failed to successfully implement such entity's or payer's performance improvement plan, the executive director shall:
- (i) Send a notice to such entity or payer, in a form and manner prescribed by the executive director, disclosing such determination and any action taken by the executive director pursuant to subparagraph (C)(ii) of this subdivision; and
- 488 (ii) In the executive director's discretion:

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- 489 (I) Extend the implementation timetable included in such 490 performance improvement plan;
 - (II) Require such entity or payer to file with the executive director, in a form and manner prescribed by the executive director, such amendments to such performance improvement plan as are, in the executive director's judgment, necessary to ensure that such entity or payer successfully implements such performance improvement plan;
 - (III) Require such entity or payer to file a new proposed performance improvement plan pursuant to subdivision (1) of subsection (e) of this

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section; or

- (IV) Waive or delay the requirement that such entity or payer file any future proposed performance improvement plan until the executive director determines, in the executive director's discretion, that such entity or payer has successfully implemented its current performance improvement plan.
- (g) The executive director shall keep confidential all nonpublic clinical, financial, operational or strategic documents and information filed with, or submitted to, the executive director pursuant to this section. The executive director shall not disclose any such document or information to any person without the consent of the health care entity or payer that filed such document or information with, or submitted such document or information to, the executive director pursuant to this section, except in summary form as part of an evaluative report if the executive director determines that such disclosure should be made in the public interest after taking into account any privacy, trade secret or anti-competitive considerations. Notwithstanding any provision of the general statutes, no document or information filed with, or submitted to, the executive director pursuant to this section shall be deemed to be a public record or subject to disclosure under the Freedom of Information Act, as defined in section 1-200 of the general statutes.
- Sec. 7. (NEW) (Effective July 1, 2021) (a) (1) For each calendar year beginning on or after January 1, 2023, if the executive director determines that the average annual percentage change in total health care expenditures for the preceding calendar year exceeded the health care cost growth benchmark for such year, the executive director shall identify each other entity that significantly contributed to exceeding such benchmark. Each identification shall be based on:
- (A) The report, if any, prepared by the executive director pursuant to subsection (c) of section 5 of this act for such calendar year;
- (B) The report filed pursuant to section 38a-479ppp of the general statutes for such calendar year;

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- 530 (C) The information and data reported to the office pursuant to 531 section 19a-754b of the general statutes for such calendar year;
- 532 (D) Information obtained from the all-payer claims database 533 established under section 19a-755a of the general statutes; and
- 534 (E) Any other information that the executive director, in the executive director's discretion, deems relevant for the purposes of this section.

- (2) The executive director shall account for costs, net of rebates and discounts, when identifying other entities pursuant to this section.
- (b) The executive director may require that any other entity that is found to be a significant contributor to health care cost growth in this state during the preceding calendar year participate in the informational public hearing held pursuant to subsection (a) of section 4 of this act. Each such other entity that is required to participate in such hearing shall provide testimony on issues identified by the executive director, and provide additional information on actions taken to reduce such other entity's contribution to future state-wide health care costs. If such other entity is a drug manufacturer, and the executive director requires that such drug manufacturer participate in such hearing with respect to a specific drug or class of drugs, such hearing may, to the extent possible, include representatives from at least one brand-name manufacturer, one generic manufacturer and one innovator company that is less than ten years old.
- Sec. 8. (NEW) (*Effective July 1, 2021*) (a) (1) For each calendar year beginning on or after January 1, 2023, the executive director shall develop and adopt annual health care quality benchmarks for health care entities and payers that:
- (A) Enable health care entities and payers to report to the executive director a standard set of information concerning health care quality for such year; and
- (B) Include measures concerning clinical health outcomes,

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- overutilization, underutilization and safety measures.
- 561 (2) In developing annual health care quality benchmarks pursuant to subdivision (1) of this subsection, the executive director shall:
- 563 (A) Consider:

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- (i) Nationally recognized quality measures that are recommended by medical groups or provider organizations concerning appropriate quality measures for such groups' or organizations' specialties; and
- 567 (ii) Measures, including, but not limited to, newly developed 568 measures, that:
- 569 (I) Concern health outcomes, overutilization, underutilization and patient safety; and
- 571 (II) Meet standards of patient-centeredness and ensure consideration 572 of important differences in preferences and clinical characteristics 573 within patient subpopulations;
 - (B) Provide stakeholders with an opportunity to engage with the executive director in developing such benchmarks; and
 - (C) Ensure that the processes the executive director uses to develop, and any research that the executive director relies upon in developing, such benchmarks is transparent.
- (b) Not later than October 1, 2022, and annually thereafter, the executive director shall, prior to adopting health care quality benchmarks pursuant to subdivision (1) of subsection (a) of this section for the calendar year next succeeding, hold an informational public hearing concerning the quality measures the executive director proposes to adopt as health care quality benchmarks for the calendar year next succeeding.
- 586 (c) Not later than November 1, 2022, and annually thereafter, the executive director shall send a notice to each health care entity, payer

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and other entity disclosing the health care quality benchmarks that the executive director has adopted for the calendar year next succeeding.

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- Sec. 9. (NEW) (*Effective July 1, 2021*) The executive director may adopt regulations, in accordance with chapter 54 of the general statutes, to implement the provisions of sections 2 to 8, inclusive, of this act.
- Sec. 10. (NEW) (*Effective July 1, 2021*) For the purposes of this section and sections 11 to 15, inclusive, of this act unless the context otherwise requires:
- 596 (1) "Drug" means an article that is (A) recognized in the official United 597 States Pharmacopoeia, official Homeopathic Pharmacopoeia of the 598 United States or official National Formulary, or any supplement thereto, 599 (B) intended for use in the diagnosis, cure, mitigation, treatment or 600 prevention of disease in humans, (C) not food and intended to affect the structure or any function of the human body, and (D) not a device and 601 602 intended for use as a component of any other article specified in 603 subparagraphs (A) to (C), inclusive, of this subdivision;
- 604 (2) "Drug Quality and Security Act" means the federal Drug Quality 605 and Security Act, 21 USC 351, et seq., as amended from time to time;
- (3) "Food, Drug and Cosmetic Act" means the Federal Food, Drug and
 Cosmetic Act, 21 USC 301, et seq., as amended by the Drug Quality and
 Security Act, as both may be amended from time to time;
- (4) "Laboratory testing" means a quantitative and qualitative analysis
 of a prescription drug consistent with the official United States
 Pharmacopoeia;
- (5) "Legend drug" means a drug that (A) any applicable federal or state law requires to be (i) dispensed pursuant to a prescription, or (ii) used by a prescribing practitioner, or (B) applicable federal law requires to bear the following legend: "RX ONLY" IN ACCORDANCE WITH GUIDELINES ESTABLISHED IN THE FEDERAL FOOD, DRUG AND COSMETIC ACT;

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618 (6) "Participating Canadian supplier" means a manufacturer or 619 wholesale drug distributor that is (A) licensed or permitted under 620 applicable Canadian law to manufacture or distribute prescription drugs, (B) exporting legend drugs, in the manufacturer's original 622 container, to a participating wholesaler for distribution in this state 623 under the program, and (C) properly registered, if such Canadian 624 supplier is required to be registered, with the United States Food and 625 Drug Administration, or any successor agency;

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- (7) "Participating wholesaler" means a wholesaler, as defined in section 21a-70 of the general statutes, that (A) has received a certificate of registration from the Commissioner of Consumer Protection pursuant to said section, and (B) is designated by the commissioner to participate in the program;
- 631 (8) "Prescription" means a lawful verbal, written or electronic order 632 by a prescribing practitioner for a drug for a specific patient;
- 633 (9) "Program" means the Canadian legend drug importation program 634 established by the Commissioner of Consumer Protection pursuant to 635 section 11 of this act:
- 636 (10) "Qualified laboratory" means a laboratory that is (A) adequately 637 equipped and staffed to properly perform laboratory testing on legend 638 drugs, and (B) accredited to International Organization for 639 Standardization (ISO) 17025; and
- 640 (11) "Track-and-trace" means the product tracing process for the 641 components of the pharmaceutical distribution supply chain, as 642 described in Title II of the Drug Quality and Security Act.
- 643 Sec. 11. (NEW) (Effective July 1, 2021) (a) The Commissioner of 644 Consumer Protection shall establish a program to be known as the 645 "Canadian legend drug importation program". Under such program, 646 the commissioner shall, notwithstanding any provision of the general 647 statutes:

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648 649	from Canada that have the highest potential for cost savings in this state;
650	and
651	(2) Designate one or more participating wholesalers to distribute
652	legend drugs in this state:
653	(A) In the manufacturer's original container;
654	(B) From a participating Canadian supplier; and
655	(C) To a pharmacy or institutional pharmacy, as both terms are
656	defined in section 20-571 of the general statutes, or a qualified
657	laboratory.
658	(b) (1) Not later than July 1, 2022, the Commissioner of Consumer
659	Protection shall submit a request to the federal Secretary of Health and
660	Human Services seeking approval for the program under 21 USC 384,
661	as amended from time to time. Such request shall, at a minimum:
662	(A) Describe the commissioner's plans for operating the program;
663	(B) Demonstrate that the legend drugs that will be imported and
664	distributed in this state under the program shall:
665	(i) Meet all applicable federal and state standards for safety and
666	effectiveness; and
667	(ii) Comply with all federal tracing procedures; and
668	(C) Disclose the costs of implementing the program.
669	(2) (A) If the federal Secretary of Health and Human Services
670	approves the commissioner's request, the commissioner shall:
671	(i) Submit to the Commissioner of Public Health a notice disclosing
672	that the federal Secretary of Health and Human Services has approved
673	such request;

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675 having cognizance of matters relating to appropriations, general law,

(ii) Submit to the joint standing committees of the General Assembly

- 676 human services and public health a notice disclosing that the federal
- 677 Secretary of Health and Human Services has approved such request;
- 678 and

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- (iii) Begin operating the program not later than one hundred eighty days after the date of such approval.
- 681 (B) Except as otherwise provided in this subsection, the 682 Commissioner of Consumer Protection shall not operate the program
- unless the federal Secretary of Health and Human Services approves the
- 684 commissioner's request.
- Sec. 12. (NEW) (Effective July 1, 2021) (a) Each participating
- 686 wholesaler may, subject to the provisions of this section and sections 11
- 687 and 14 of this act, import into this state a legend drug from a
- 688 participating Canadian supplier, and distribute such legend drug to a
- 689 pharmacy or institutional pharmacy, as both terms are defined in
- section 20-571 of the general statutes, or a qualified laboratory in this
- 691 state, under the program if:
- 692 (1) Such participating wholesaler:
- (A) Is registered with the federal Secretary of Health and Human
- 694 Services pursuant to Section 510(b) of the Food, Drug and Cosmetic Act,
- 695 21 USC 360(b), as amended from time to time; and
- (B) Holds a valid labeler code that has been issued to such
- 697 participating wholesaler by the United States Food and Drug
- 698 Administration, or any successor agency; and
- 699 (2) Such legend drug:
- 700 (A) May be imported into this state in accordance with applicable federal patent laws;
- 702 (B) Meets the United States Food and Drug Administration's, or any

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- successor agency's, standards concerning drug safety, effectiveness,
- 704 misbranding and adulteration; and
- 705 (C) Is not:
- 706 (i) A controlled substance, as defined in 21 USC 802, as amended from 707 time to time;
- 708 (ii) A biological product, as defined in 42 USC 262, as amended from time to time;
- 710 (iii) An infused drug;
- 711 (iv) An intravenously injected drug;
- 712 (v) A drug that is inhaled during surgery; or
- 713 (vi) A drug that is a parenteral drug, the importation of which is 714 determined by the federal Secretary of Health and Human Services to
- 715 pose a threat to the public health.
- 716 (b) Each participating wholesaler shall:
- 717 (1) Comply with all applicable track-and-trace requirements, and
- 718 make available to the Commissioner of Consumer Protection all track-
- and-trace records not later than forty-eight hours after the commissioner
- 720 requests such records;
- 721 (2) Not import, distribute, dispense or sell in this state any legend
- 722 drugs under the program except in accordance with the provisions of
- 723 this section and sections 11 and 14 of this act;
- 724 (3) Not distribute, dispense or sell outside of this state any legend 725 drugs that are imported into this state under the program;
- 726 (4) Ensure the safety and quality of the legend drugs that are 727 imported and distributed in this state under the program;
- 728 (5) For each initial shipment of a legend drug that is imported into

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- 729 this state by such participating wholesaler, ensure that a qualified
- 730 laboratory engaged by such participating wholesaler tests a statistically
- valid sample size for each batch of such legend drug in such shipment
- for authenticity and degradation in a manner that is consistent with the
- 733 Food, Drug and Cosmetic Act;
- 734 (6) For each shipment of a legend drug that is imported into this state
- by such participating wholesaler, and sampled and tested pursuant to
- 736 subdivision (5) of this subsection, ensure that a qualified laboratory
- 737 engaged by such participating wholesaler tests a statistically valid
- 738 sample of such legend drug in such shipment for authenticity and
- 739 degradation in a manner that is consistent with the Food, Drug and
- 740 Cosmetic Act;
- 741 (7) Certify to the Commissioner of Consumer Protection that each
- 742 legend drug imported into this state under the program:
- 743 (A) Is approved for marketing in the United States and not
- 744 adulterated or misbranded; and
- 745 (B) Meets all labeling requirements under 21 USC 352, as amended
- 746 from time to time;
- 747 (8) Maintain laboratory records, including, but not limited to,
- 748 complete data derived from all tests necessary to ensure that each
- 749 legend drug imported into this state under the program satisfies the
- 750 requirements of subdivisions (5) and (6) of this subsection;
- 751 (9) Maintain documentation demonstrating that the testing required
- 752 by subdivisions (5) and (6) of this subsection was conducted at a
- 753 qualified laboratory in accordance with the Food, Drug and Cosmetic
- 754 Act and all other applicable federal and state laws and regulations
- 755 concerning laboratory qualifications;
- 756 (10) Maintain the following information for each legend drug that
- such participating wholesaler imports and distributes in this state under
- 758 the program, and submit such information to the Commissioner of

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- 759 Consumer Protection upon request by the commissioner:
- 760 (A) The name and quantity of the active ingredient of such legend 761 drug;
- 762 (B) A description of the dosage form of such legend drug;
- 763 (C) The date on which such participating wholesaler received such legend drug;
- 765 (D) The quantity of such legend drug that such participating 766 wholesaler received;
- 767 (E) The point of origin and destination of such legend drug;
- 768 (F) The price paid by such participating wholesaler for such legend drug;
- 770 (G) A report for any legend drug that fails laboratory testing under 771 subdivision (5) or (6) of this subsection; and
- 772 (H) Such additional information and documentation that the 773 commissioner deems necessary to ensure the protection of the public 774 health; and
- 775 (11) Maintain all information and documentation that is submitted to 776 the Commissioner of Consumer Protection pursuant to this subsection 777 for a period of not less than three years.
- Sec. 13. (NEW) (*Effective July 1, 2021*) Each participating Canadian supplier shall:
- 780 (1) Comply with all applicable track-and-trace requirements;
- 781 (2) Not distribute, dispense or sell outside of this state any legend 782 drugs that are imported into this state under the program; and
- 783 (3) Maintain the following information and documentation and, 784 upon request by the Commissioner of Consumer Protection, submit

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- such information and documentation to the commissioner for each legend drug that such participating Canadian supplier exports into this state under the program:
- 788 (A) The original source of such legend drug, including, but not limited to:
- 790 (i) The name of the manufacturer of such legend drug;
- 791 (ii) The date on which such legend drug was manufactured; and
- 792 (iii) The location where such legend drug was manufactured;
- 793 (B) The date on which such legend drug was shipped to a participating wholesaler;
- 795 (C) The quantity of such legend drug that was shipped to a participating wholesaler;
- 797 (D) The quantity of each lot of such legend drug that such 798 participating Canadian supplier originally received and the source of 799 such lot;
- 800 (E) The lot or control number and the batch number assigned to such legend drug by the manufacturer; and
- (F) Such additional information and documentation that the commissioner deems necessary to ensure the protection of the public health.
- Sec. 14. (NEW) (*Effective July 1, 2021*) (a) The Commissioner of Consumer Protection shall issue a written order:
- (1) Suspending importation and distribution of a legend drug under the program if the commissioner discovers that such distribution or importation violates any provision of sections 11 to 13, inclusive, of this act or any other applicable state or federal law or regulation;
- 811 (2) Suspending all importation and distribution of legend drugs by a

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- participating wholesaler under the program if the commissioner discovers that the participating wholesaler has violated any provision of section 11 or 12 of this act or any other applicable state or federal law or regulation;
- (3) Suspending all importation and distribution of legend drugs by a participating Canadian supplier under the program if the commissioner discovers that the participating Canadian supplier has violated any provision of section 11 or 13 of this act or any other applicable state or federal law or regulation; or
- 821 (4) Requiring the recall or seizure of any legend drug that was 822 imported and distributed under the program and has been identified as 823 adulterated, within the meaning of section 21a-105 of the general 824 statutes, or misbranded.

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- (b) The Commissioner of Consumer Protection shall send a notice to each participating Canadian supplier and participating wholesaler affected by an order issued pursuant to subsection (a) of this section notifying such participating Canadian supplier or participating wholesaler that:
- 830 (1) The commissioner has issued such order, and providing the legal 831 and factual basis for such order; and
 - (2) Such participating Canadian supplier or participating wholesaler may request, in writing, a hearing before the commissioner, provided such request is received by the commissioner not later than thirty days after the date of such notice.
 - (c) If a participating Canadian supplier or participating wholesaler timely requests a hearing pursuant to subsection (b) of this section, the Commissioner of Consumer Protection shall, not later than thirty days after the receipt of the request, convene the hearing as a contested case in accordance with the provisions of chapter 54 of the general statutes. Not later than sixty days after the receipt of such request, the commissioner shall issue a final decision vacating, modifying or

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- Raised Bill No. 1006 843 affirming the commissioner's order. A participating Canadian supplier 844 or participating wholesaler aggrieved by a final decision may appeal 845 such decision in accordance with the provisions of section 4-183 of the 846 general statutes. 847 Sec. 15. (NEW) (Effective July 1, 2021) The Commissioner of Consumer 848 Protection may, in consultation with the Commissioner of Public 849 Health, adopt regulations in accordance with the provisions of chapter 850 54 of the general statutes to implement the provisions of sections 10 to 851 14, inclusive, of this act. 852 Sec. 16. Section 38a-8b of the general statutes is repealed and the 853 following is substituted in lieu thereof (*Effective January 1, 2022*): 854 (a) For the purposes of this section: 855 (1) "Attachment point" means the dollar value of claims incurred by 856 a policyholder at which the insurer that issues or delivers a medical stop-loss insurance policy to the policyholder incurs liability to such 857 858
- policyholder for payment under such medical stop-loss insurance 859 policy;
- (2) "Employee" has the same meaning as provided in section 38a-564; 860
- 861 (3) "Expected claims" means the dollar value of claims that, in the absence of a medical stop-loss insurance policy, the policyholder of a 862 863 medical stop-loss insurance policy is projected to incur under such policyholder's health benefit plan; 864
- 865 (4) "Lasering" means assigning a different attachment point or deductible, or denying coverage altogether, under a medical stop-loss 866 867 insurance policy for an enrollee or a dependent because the enrollee or 868 dependent has a high-cost preexisting condition or another identified 869 risk;
- 870 "Medical stop-loss insurance" means stop-loss insurance 871 purchased by a person, other than a health carrier or health care 872 provider, and providing coverage for catastrophic, excess or unexpected

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873	losses incurred by the policyholder, and due and owing to a third party,
874	under a health benefit plan not providing coverage for retirees;
875	(6) "Medical stop-loss insurer" means an insurer that is licensed
876	pursuant to section 38a-41 to sell, issue and deliver medical stop-loss
877	insurance in this state;
878	(7) "Retiree stop-loss insurance" means stop-loss insurance purchased
879	by a person, other than a health carrier or health care provider, and
880	providing coverage for catastrophic, excess or unexpected losses
881	incurred by the policyholder, and due and owing to a third party, under
882	a health benefit plan providing coverage for retirees; and
883	(8) "Stop-loss insurance" means insurance, other than reinsurance,
884	providing coverage for catastrophic, excess or unexpected losses
885	incurred by the policyholder, and due and owing to a third party, under
886	another insurance policy or a health benefit plan.
887	(b) No [stop loss] stop-loss insurance policy [may] shall be issued or
888	delivered in this state unless a copy of the [stop loss] $\underline{\text{stop-loss insurance}}$
889	policy form has been submitted to, and approved by, the Insurance
890	Commissioner. [pursuant to regulations that the commissioner may
891	adopt in accordance with chapter 54. Such regulations, if adopted, shall
892	include, but need not be limited to, a definition of a stop loss policy and
893	the standards for filing and review of stop loss policies.]
894	(c) (1) Except as provided in subdivision (4) of subsection (d) of this
895	section, no medical stop-loss insurer shall issue or deliver, and the
896	Insurance Commissioner shall not approve, a medical stop-loss
897	insurance policy in this state on or after January 1, 2022, if the medical
898	stop-loss insurance policy:
899	(A) Imposes an annual attachment point that is less than twenty
900	thousand dollars for claims incurred per enrolled employee or
901	dependent;

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(B) Imposes an annual aggregate attachment point:

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903	(i) That is less than the greatest of the following amounts for an
904	insured group consisting of not more than fifty employees, as calculated
905	in the manner set forth in subdivision (2) of this subsection:
906	(I) Four thousand dollars multiplied by the number of employees in
907	such insured group;
908	(II) One hundred twenty per cent of the expected claims for such
909	insured group; or
910	(III) Twenty thousand dollars; or
911	(ii) That is less than one hundred ten per cent of the expected claims
912	for an insured group consisting of more than fifty employees, as
913	calculated in the manner set forth in subdivision (2) of this subsection;
914	(C) Provides direct coverage for an enrollee's or dependent's health
915	care expenses;
916	(D) Provides for a determination regarding whether a benefit is:
917	(i) Medically necessary;
918	(ii) Usual or customary; or
919	(iii) Experimental or investigational;
920	(E) Imposes a case management requirement or an annual dollar
921	limitation for an enrolled employee, dependent or benefit;
922	(F) Requires an enrolled employee or dependent to use a provider
923	network or provides a benefit incentive for an enrolled employee or
924	dependent to use a provider participating in a provider network;
925	(G) Provides the medical stop-loss insurer with a right to examine an
926	enrolled employee or dependent;
927	(H) Permits the medical stop-loss insurer to:
928	(i) Deny a claim if the policyholder is legally obligated to pay the

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929	claim under such policyholder's health benefit plan;
930	(ii) Rescind such medical stop-loss insurance policy for any reason
931	other than fraud or intentional misrepresentation;
932	(iii) Terminate such medical stop-loss insurance policy, in the sole
933	discretion of such medical stop-loss insurer, in any manner that is
934	inconsistent with applicable laws concerning cancellation or
935	nonrenewal of medical stop-loss insurance policies; or
936	(iv) Increase the rates imposed under such medical stop-loss
937	insurance policy, in the sole discretion of such medical stop-loss insurer,
938	during the term of such medical stop-loss insurance policy;
939	(I) Requires an enrolled employee to be actively at work; or
940	(J) Contains any provision that is misleading, deceptive or contrary
941	to any provision of the general statutes or the public interest.
942	(2) (A) For the purposes of subparagraph (B) of subdivision (1) of this
943	subsection, the number of employees in an insured group shall be
944	determined by adding:
945	(i) The number of the policyholder's full-time employees for each
946	month who work a normal work week of thirty hours or more; and
947	(ii) The number of the policyholder's full-time equivalent employees,
948	calculated for each month by dividing by one hundred twenty the
949	aggregate number of hours worked for such month by employees who
950	work a normal work week of less than thirty hours, and averaging such
951	total for the calendar year.
952	(B) If a policyholder was not in existence throughout the preceding
953	calendar year, the number of employees shall be based on the average
954	number of employees that such policyholder reasonably expects to
955	employ in the current calendar year.
956	(d) Each insurer that underwrites a medical stop-loss insurance

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957 policy issued or delivered in this state on or after January 1, 2022, may 958 use lasering in underwriting such medical stop-loss insurance policy, provided: 959 960 (1) If such insurer uses lasering in underwriting such medical stoploss insurance policy, such insurer and any insurance producer who 961 sells, solicits or negotiates such medical stop-loss insurance policy on 962 963 behalf of such insurer includes in each application for coverage under such medical stop-loss insurance policy: 964 965 (A) A statement disclosing the increased financial risk that each 966 prospective policyholder under such medical stop-loss insurance policy 967 will bear because such insurer intends to use lasering in underwriting 968 such medical stop-loss insurance policy, and any alternatives available 969 to each such prospective policyholder with respect to such insurer's 970 intended use of lasering in underwriting such medical stop-loss 971 insurance policy; 972 (B) A statement by such insurer or insurance producer, as applicable, 973 affirming that such insurer or insurance producer fully explained to 974 each prospective policyholder under such medical stop-loss insurance 975 policy the increased financial risk described in subparagraph (A) of this subdivision and that each such prospective policyholder understands 976 977 such increased financial risk; and 978 (C) The signature of such insurer, insurance producer and each 979 prospective policyholder below the statement required under 980 subparagraph (B) of this subdivision; 981 (2) If such insurer uses lasering on the effective date of such medical stop-loss insurance policy, such insurer shall not change such lasering 982 983 during the term of such medical stop-loss insurance policy; 984 (3) If such insurer does not use lasering on the effective date of such

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medical stop-loss insurance policy, such insurer shall not use lasering

during the term of such medical stop-loss insurance policy; and

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- 987 (4) The attachment point for an enrolled employee under such medical stop-loss insurance policy shall not exceed an amount that is equal to three hundred per cent of the attachment point for such medical stop-loss insurance policy.
- 991 (e) No retiree stop-loss insurance policy issued or delivered in this 992 state on or after January 1, 2022, shall be subject to the provisions of 993 subsection (c) or (d) of this section, and the Insurance Commissioner 994 shall review and approve, on a case-by case basis, such retiree stop-loss 995 insurance policies for issuance and delivery in this state on or after said 996 date.
- 997 <u>(f) The Insurance Commissioner may adopt regulations, in</u> 998 accordance with chapter 54, to carry out the purposes of this section.

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- Sec. 17. Subparagraph (C) of subdivision (3) of subsection (m) of section 5-259 of the general statutes is repealed and the following is substituted in lieu thereof (*Effective January 1, 2022*):
 - (C) The Comptroller may offer to nonstate public employers that choose to purchase prescription drugs pursuant to subparagraph (A) of this subdivision the option to purchase [stop loss] <u>stop-loss</u> coverage from an insurer at a rate negotiated by the Comptroller.
- Sec. 18. Subdivision (1) of subsection (c) of section 7-464 of the general statutes is repealed and the following is substituted in lieu thereof (*Effective January 1, 2022*):
 - (1) In no event shall any commercial insurance company which provides health insurance benefits to the employees of a town, city or borough and their covered dependents and family members, including, but not limited to, [stop loss] stop-loss insurance beyond a municipal self-funded medical expense amount, be entitled to any reimbursement from a tortfeasor recovery. The provisions of this subsection shall be construed to only permit a self-insured town, city or borough to recover medical expenses paid from its own revenues. The provisions of this subsection shall not be construed to permit a self-insured town, city or

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1018 borough to recover medical expenses paid from an insured plan, 1019 whether insured in whole or in part.

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- 1020 Sec. 19. Subparagraph (F) of subdivision (18) of section 38a-465 of the general statutes is repealed and the following is substituted in lieu 1022 thereof (*Effective January 1, 2022*):
- 1023 (F) An authorized or eligible insurer that provides [stop loss] stop-1024 <u>loss</u> coverage to a provider, purchaser, financing entity, special purpose 1025 entity or related provider trust;
- 1026 Sec. 20. Subsection (c) of section 38a-465d of the general statutes is 1027 repealed and the following is substituted in lieu thereof (*Effective January* 1028 1, 2022):
 - (c) Except as otherwise required or permitted by law, no person, including, but not limited to, a provider, broker, insurance company, insurance producer, information bureau, rating agency or company, or any other person with actual knowledge of an insured's identity, shall disclose such identity or information where there is a reasonable basis to conclude such information could be used to identify the insured or the insured's financial or medical information to any other person unless such disclosure: (1) Is necessary to effect a life settlement contract between the owner and a provider and the owner and insured have provided prior written consent to such disclosure; (2) is provided in response to an investigation or examination by the commissioner or any other governmental office or agency or pursuant to the requirements of section 38a-465i; (3) is necessary to effectuate the sale of life settlement contracts or interests therein as investments, provided the sale is conducted in accordance with applicable state and federal securities laws, and provided further the owner and the insured have both provided prior written consent to the disclosure; (4) is a term of or condition to the transfer of a policy by one provider to another provider, in which case the provider receiving such information shall comply with the confidentiality requirements specified in this subsection; (5) is necessary to allow the provider or broker or their authorized

LCO No. 3680 **36** of 55 representatives to make contacts for the purpose of determining health status. For the purpose of this section, "authorized representative" does not include any person who has or may have a financial interest in the settlement contract other than a provider, licensed broker, financing entity, related provider trust or special purpose entity. Each provider or broker shall require its authorized representative to agree in writing to comply with the privacy provisions of this part; or (6) is required to purchase [stop loss] stop-loss coverage.

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- Sec. 21. Subparagraph (A) of subdivision (2) of subsection (b) of section 38a-478*l* of the general statutes is repealed and the following is substituted in lieu thereof (*Effective January 1*, 2022):
- (A) "State medical loss ratio" means the ratio of incurred claims to earned premiums for the prior calendar year for managed care plans issued in the state. Claims shall be limited to medical expenses for services and supplies provided to enrollees and shall not include expenses for [stop loss] stop-loss coverage, reinsurance, enrollee educational programs or other cost containment programs or features;
- Sec. 22. Subsection (c) of section 38a-720h of the general statutes is repealed and the following is substituted in lieu thereof (*Effective January* 1069 1, 2022):
- 1070 (c) The third-party administrator shall disclose to the insurer or other person utilizing the services of the third-party administrator all charges, fees and commissions that the third-party administrator receives arising from services it provides for the insurer or other person utilizing the services of the third-party administrator, including any fees or commissions paid by insurers providing reinsurance or [stop loss] stop-loss coverage.
- Sec. 23. (NEW) (*Effective from passage*) (a) For the purposes of this section:
- 1079 (1) "Affordable Care Act" has the same meaning as provided in section 38a-1080 of the general statutes;

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1081 (2) "Exchange" means the Connecticut Health Insurance Exchange 1082 established under section 38a-1081 of the general statutes, as amended 1083 by this act; and

- (3) "Office" means the Office of Health Strategy established under section 19a-754a of the general statutes, as amended by this act.
- 1086 (b) The office shall, in conjunction with the Office of Policy and
 1087 Management, the Insurance Department and the Health Reinsurance
 1088 Association created under section 38a-556 of the general statutes, seek a
 1089 state innovation waiver under Section 1332 of the Affordable Care Act
 1090 to establish a reinsurance program pursuant to subsection (d) of this
 1091 section.
 - (c) Subject to the approval of a waiver described in subsection (b) of this section, the office, not later than September 1, 2022, for plan year 2023 and annually thereafter for the subsequent plan year, shall:
 - (1) Determine the amount needed, not to exceed twenty-one million two hundred ten thousand dollars, annually, to fund the reinsurance program established pursuant to subsection (d) of this section; and
 - (2) Inform the Office of Policy and Management of the amount determined pursuant to subdivision (1) of this subsection.
 - (d) The amount described in subsection (c) of this section shall be utilized to establish a reinsurance program for the individual health insurance market designed to lower premiums on health benefit plans sold in such market, on and off the exchange, provided the federal government approves the waiver described in subsection (b) of this section. Any such reinsurance program shall be administered by the Health Reinsurance Association. The Treasurer shall annually pay the amount as described in subsection (c) of this section for the purpose of administering such reinsurance program.
 - (e) If the waiver described in subsection (b) of this section terminates and the office does not obtain another waiver pursuant to subsection (a)

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- of this section, the Treasurer shall cease paying the amount described in
- 1112 subsection (c) of this section for the purpose of administering the
- 1113 reinsurance program established pursuant to subsection (d) of this
- 1114 section.
- 1115 Sec. 24. (NEW) (Effective from passage) (a) Not later than January 31,
- 1116 2022, the Auditors of Public Accounts shall annually conduct an audit
- of each health care plan administered or offered by this state to persons
- other than state employees during the preceding calendar year.
- 1119 (b) Not later than March 1, 2022, and annually thereafter, the
- 1120 Auditors of Public Accounts shall submit a report, in accordance with
- the provisions of section 11-4a of the general statutes, disclosing the
- results of the audit conducted pursuant to subsection (a) of this section
- for the preceding calendar year to the joint standing committees of the
- 1124 General Assembly having cognizance of matters relating to
- appropriations, finance, revenue and bonding and human services.
- (c) The Auditors of Public Accounts may, in their discretion, engage
- the services of such third-party actuaries, professionals and specialists
- that the Auditors of Public Accounts deem necessary to assist the
- Auditors of Public Accounts to perform their duties under this section.
- 1130 Sec. 25. Section 38a-1081 of the general statutes is repealed and the
- following is substituted in lieu thereof (*Effective October 1, 2021*):
- 1132 (a) There is hereby created as a body politic and corporate,
- 1133 constituting a public instrumentality and political subdivision of the
- 1134 state created for the performance of an essential public and
- 1135 governmental function, to be known as the Connecticut Health
- 1136 Insurance Exchange. The Connecticut Health Insurance Exchange shall
- 1137 not be construed to be a department, institution or agency of the state.
- 1138 The exchange shall serve both qualified individuals and qualified
- 1139 employers.
- (b) (1) (A) The powers of the exchange shall be vested in and
- exercised by a board of directors, which, until June 19, 2013, shall consist

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- of twelve voting members. The appointment of the initial board members shall be as follows:
- (i) The Governor shall appoint two board members, one of whom shall have expertise in the area of individual health insurance coverage and shall serve for a term of three years and one of whom shall have expertise in issues relating to small employer health insurance coverage and shall serve for a term of two years;

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- (ii) The president pro tempore of the Senate shall appoint one board member who shall have expertise in the area of health care finance and shall serve for a term of four years;
 - (iii) The speaker of the House of Representatives shall appoint one board member who shall have expertise in the area of health care benefits plan administration and shall serve for a term of four years;
- 1155 (iv) The majority leader of the Senate shall appoint one board 1156 member who shall have expertise in the health care delivery systems 1157 and shall serve for a term of two years;
- (v) The majority leader of the House of Representatives shall appoint one board member who shall have expertise in the area of health care economics and shall serve for a term of two years;
 - (vi) The minority leader of the Senate shall appoint one board member who shall have expertise in health care access issues faced by self-employed individuals and shall serve for a term of three years;
- (vii) The minority leader of the House of Representatives shall appoint one board member who shall have expertise concerning barriers to individual health care coverage and shall serve for a term of two years;
 - (viii) The Commissioner of Social Services, the Special Advisor to the Governor on Healthcare Reform, the Secretary of the Office of Policy and Management and the Healthcare Advocate, or their designees, who shall serve as ex-officio, voting board members; and

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1172 (ix) The Insurance Commissioner and the Commissioner of Public 1173 Health, or their designees, who shall serve as ex-officio, nonvoting 1174 board members.

- (B) On and after June 19, 2013, the board of directors shall consist of eleven voting members and three nonvoting members as follows: (i) The board members appointed pursuant to subparagraphs (A)(i) to (A)(vii), inclusive, of this subdivision, except that each such board member appointed or reappointed on or after October 1, 2021, shall have expertise in the area of insurance; (ii) the Commissioner of Social Services, the Secretary of the Office of Policy and Management and the Healthcare Advocate, or their designees, who shall serve as ex-officio, voting board members; and (iii) the Insurance Commissioner and the Commissioners of Public Health and Mental Health and Addiction Services, or their designees, who shall serve as ex-officio, nonvoting board members. The provisions of this subparagraph shall not affect the terms of the board members set forth in subparagraphs (A)(i) to (A)(vii), inclusive, of this subdivision.
- (2) (A) No board member shall be employed by, a consultant to, a member of the board of directors of, affiliated with or otherwise a representative of (i) an insurer, (ii) an insurance producer or broker, (iii) a health care provider, or (iv) a health care facility or health or medical clinic while serving on the board of the exchange. For purposes of this subdivision, "health care provider" means any person that is licensed in this state, or operates or owns a facility or institution in this state, to provide health care or health care professional services in this state, or an officer, employee or agent thereof acting in the course and scope of such officer's, employee's or agent's employment.
- (B) No board member shall be a member of, a member of the board of, a consultant to or an employee of a trade association of (i) insurers, (ii) insurance producers or brokers, (iii) health care providers, or (iv) health care facilities or health or medical clinics while serving on the board of the exchange.

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(C) No board member shall be a health care provider unless such member receives no compensation for rendering services as a health care provider and does not have an ownership interest in a professional health care practice.

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- (c) (1) All initial appointments shall be made not later than July 1, 2011. Following the expiration of such initial terms, subsequent board member terms shall be for four years, except that no board member shall serve more than eight years. Any board member appointed to the board before October 1, 2021, who has served eight or more years on the board may complete such board member's term. Any vacancy shall be filled by the appointing authority for the balance of the unexpired term. If an appointing authority fails to make an initial appointment, or an appointment to fill a vacancy within ninety days of the date of such vacancy, the appointed board members may make such appointment by a majority vote. Any board member previously appointed to the board or appointed to fill a vacancy may be reappointed in accordance with this section unless such reappointment would cause the board member to serve on the board for more than eight years. Any board member may be removed for misfeasance, malfeasance or wilful neglect of duty at the sole direction of the appointing authority.
 - (2) As a condition of qualifying as a member of the board of directors, each appointee shall, before entering upon such member's duties, take and subscribe the oath or affirmation required under section 1 of article eleventh of the Constitution of the state. A record of each such oath shall be filed in the office of the Secretary of the State.
- (3) Appointed board members may not designate a representative to perform in their absence their respective duties under sections 38a-1080 to 38a-1092, inclusive. The Governor shall select a chairperson from among the board members and the board members shall annually elect a vice-chairperson. Meetings of the board of directors shall be held at such times as shall be specified in the bylaws adopted by the board and at such other time or times as the chairperson deems necessary. Any board member who fails to attend more than fifty per cent of all

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meetings held during any calendar year shall be deemed to have resigned from the board.

- (4) Six board members shall constitute a quorum for the transaction of any business or the exercise of any power of the exchange. For the transaction of any business or the exercise of any power of the exchange, the exchange may act by a majority of the board members present at any meeting at which a quorum is in attendance. No vacancy in the membership of the board of directors shall impair the right of such board members to exercise all the rights and perform all the duties of the board. Except as otherwise provided in sections 38a-1080 to 38a-1092, inclusive, any action taken by the board under the provisions of sections 38a-1080 to 38a-1092, inclusive, may be authorized by resolution approved by a majority of the board members present at any regular or special meeting, which resolution shall take effect immediately unless otherwise provided in the resolution.
- (5) Board members shall receive no compensation for their services but shall receive actual and necessary expenses incurred in the performance of their official duties.
- (6) Subject to the provisions of subdivision (2) of subsection (b) of this section, board members may engage in private employment or in a profession or business, subject to any applicable laws, rules and regulations of the state or federal government regarding official ethics or conflicts of interest.
- (7) Notwithstanding any provision of the general statutes, it shall not constitute a conflict of interest for a trustee, director, partner or officer of any person, firm or corporation, or any individual having a financial interest in a person, firm or corporation, to serve as a board member of the exchange, provided such trustee, director, partner, officer or individual shall abstain from deliberation, action or vote by the exchange in specific request to such person, firm or corporation.
- (8) Each board member shall execute a surety bond in the penal sum of fifty thousand dollars, or, in lieu thereof, the chairperson of the board

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shall execute a blanket position bond or procure an equivalent insurance product covering each board member, the chief executive officer and the employees of the exchange, each surety bond or equivalent insurance product to be conditioned upon the faithful performance of the duties of the office or offices covered, to be issued by an insurance company authorized to transact business in this state for surety or such equivalent insurance product. The cost of each such bond or insurance product shall be paid by the exchange.

- (9) No board member of the exchange shall, for one year after the end of such member's service on the board, accept employment with any health carrier that offers a qualified health benefit plan through the exchange.
- (d) (1) With respect to the initial appointment of a chief executive officer of the exchange, the board of directors shall nominate three candidates to the Governor, who shall make a selection from such nominations. After such initial appointment, the board shall select and appoint subsequent chief executive officers.
- (2) The chief executive officer shall be responsible for administering the exchange's programs and activities in accordance with the policies and objectives established by the board. The chief executive officer (A) may employ such other employees as shall be designated by the board of directors, and (B) shall attend all meetings of the board, keep a record of all proceedings and maintain and be custodian of all records, books, documents and papers filed with or compiled by the exchange.
- (e) (1) (A) No employee of the exchange shall be employed by, a consultant to, a member of the board of directors of, affiliated with or otherwise a representative of (i) an insurer, (ii) an insurance producer or broker, (iii) a health care provider, or (iv) a health care facility or health or medical clinic while serving on the staff of the exchange. For purposes of this subdivision, "health care provider" means any person that is licensed in this state, or operates or owns a facility or institution in this state, to provide health care or health care professional services in this

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state, or an officer, employee or agent thereof acting in the course and scope of such officer's, employee's or agent's employment.

- (B) No employee of the exchange shall be a member of, a member of the board of, a consultant to or an employee of a trade association of (i) insurers, (ii) insurance producers or brokers, (iii) health care providers, or (iv) health care facilities or health or medical clinics while serving on the staff of the exchange.
- (C) No employee of the exchange shall be a health care provider unless (i) (I) such employee receives no compensation for rendering services as a health care provider, or (II) the chief executive officer approves the hiring of such provider as an employee on the basis that such provider fills an area of need of expertise for the exchange, and (ii) such employee does not have an ownership interest in a professional health care practice.
 - (2) No employee of the exchange shall, for one year after terminating employment with the exchange, accept employment with any health carrier that offers a qualified health benefit plan through the exchange.
 - (3) Any employee of the exchange whose primary purpose is to assist individuals or small employers in selecting health insurance plans offered through the exchange to purchase shall be licensed as an insurance producer under chapter 701a not later than eighteen months after such employee begins employment with the exchange.
 - (4) Any employee of the exchange may enroll in a group hospitalization and medical and surgical insurance plan under subsection (a) of section 5-259, <u>as amended by this act</u>, provided the exchange reimburses the appropriate state agencies for all costs incurred by such enrollment.
 - (f) The board may consult with such parties, public or private, as it deems desirable or necessary in exercising its duties under sections 38a-1080 to 38a-1093, inclusive, as amended by this act.

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- 1331 (g) The board may create such advisory committees as it deems 1332 necessary to provide input on issues that may include, but are not 1333 limited to, customer service needs and insurance producer concerns.
- Sec. 26. Section 38a-1083 of the general statutes is repealed and the following is substituted in lieu thereof (*Effective October 1, 2021*):

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- (a) For purposes of sections 38a-1080 to 38a-1093, inclusive, <u>as</u> <u>amended by this act</u>, "purposes of the exchange" means the purposes of and the pursuit of the goals of the exchange expressed in and pursuant to this section and the performance of the duties and responsibilities of the exchange set forth in sections 38a-1084 to 38a-1087, inclusive, which are hereby determined to be public purposes for which public funds may be expended. The powers enumerated in this section shall be interpreted broadly to effectuate the purposes of the exchange and shall not be construed as a limitation of powers.
- (b) The goals of the exchange shall be to reduce the number of individuals without health insurance in this state and assist individuals and small employers in the procurement of health insurance by, among other services, offering easily comparable and understandable information about health insurance options.
 - (c) The exchange is authorized and empowered to:
- 1351 (1) Have perpetual succession as a body politic and corporate and to 1352 adopt bylaws for the regulation of its affairs and the conduct of its 1353 business;
- 1354 (2) Adopt an official seal and alter the same at pleasure;
- 1355 (3) Maintain an office in the state at such place or places as it may designate;
- (4) Employ such assistants, agents, managers and other employees asmay be necessary or desirable;
- 1359 (5) Acquire, lease, purchase, own, manage, hold and dispose of real

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and personal property, and lease, convey or deal in or enter into agreements with respect to such property on any terms necessary or incidental to the carrying out of these purposes, provided all such acquisitions of real property for the exchange's own use with amounts appropriated by this state to the exchange or with the proceeds of bonds supported by the full faith and credit of this state shall be subject to the approval of the Secretary of the Office of Policy and Management and the provisions of section 4b-23;

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- (6) Receive and accept, from any source, aid or contributions, including money, property, labor and other things of value;
- 1370 (7) Charge assessments or user fees to health carriers that are capable 1371 of offering a qualified health plan through the exchange, [or] implement 1372 and change methods of calculating such assessments and fees and 1373 otherwise generate funding necessary to support the operations of the 1374 exchange, [and impose] provided each such proposed assessment or fee 1375 to be charged, any proposed increase in the amount of any such 1376 assessment or fee to be imposed and any proposed method, or change 1377 to any method, used to calculate any such assessment or fee to be 1378 implemented on or after October 1, 2021, shall be:
 - (A) The subject of a public meeting of the board of directors held for the purpose of receiving public comment concerning such proposed assessment, fee, increase, method or change in method before such assessment or fee is charged, increase is imposed or method, or change in method, is implemented; and
- 1384 (B) Subject to prior legislative approval under subsection (d) of this section;
- 1386 (8) Impose interest and penalties on [such] health carriers for delinquent payments of [such] assessments or <u>user</u> fees;
- [(8)] (9) Procure insurance against loss in connection with its property and other assets in such amounts and from such insurers as it deems desirable;

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- [(9)] (10) Invest any funds not needed for immediate use or disbursement in obligations issued or guaranteed by the United States of America or the state and in obligations that are legal investments for savings banks in the state;

 [(10)] (11) Issue bonds, bond anticipation notes and other obligations
 - [(10)] (11) Issue bonds, bond anticipation notes and other obligations of the exchange for any of its corporate purposes, and to fund or refund the same and provide for the rights of the holders thereof, and to secure the same by pledge of revenues, notes and mortgages of others;
- [(11)] (12) Borrow money for the purpose of obtaining working capital;

- [(12)] (13) Account for and audit funds of the exchange and any recipients of funds from the exchange;
 - [(13)] (14) Make and enter into any contract or agreement necessary or incidental to the performance of its duties and execution of its powers, [. The] provided any proposed severance or nondisclosure agreement to be entered into on or after October 1, 2021, shall be subject to prior legislative approval under subsection (d) of this section. Except as otherwise provided in this subdivision, the contracts entered into by the exchange shall not be subject to the approval of any other state department, office or agency, provided copies of all contracts of the exchange shall be maintained by the exchange as public records, subject to the proprietary rights of any party to the contract;
 - [(14)] (15) To the extent permitted under its contract with other persons, consent to any termination, modification, forgiveness or other change of any term of any contractual right, payment, royalty, contract or agreement of any kind to which the exchange is a party;
 - [(15)] (16) Award grants to trained and certified individuals and institutions that will assist individuals, families and small employers and their employees in enrolling in appropriate coverage through the exchange. Applications for grants from the exchange shall be made on a form prescribed by the board;

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1422	[(16)] (17) Limit the number of plans offered, and use selective criteria
1423	in determining which plans to offer, through the exchange, provided
1424	individuals and employers have an adequate number and selection of
1425	choices;
1426	[(17)] (18) Evaluate jointly with the Health Care Cabinet established
1427	pursuant to section 19a-725 the feasibility of implementing a basic
1428	health program option as set forth in Section 1331 of the Affordable Care
1429	Act;
1430	[(18)] (19) Establish one or more subsidiaries, in accordance with
1431	section 38a-1093, as amended by this act, to further the purposes of the
1432	exchange;
1433	[(19)] (20) Make loans to each subsidiary established pursuant to
1434	section 38a-1093, as amended by this act, from the assets of the exchange
1435	and the proceeds of bonds, bond anticipation notes and other
1436	obligations issued by the exchange or assign or transfer to such
1437	subsidiary any of the rights, moneys or other assets of the exchange,
1438	provided such assignment or transfer is not in violation of state or
1439	federal law;
1440	[(20)] (21) Sue and be sued, plead and be impleaded;
1441	[(21)] (22) Adopt regular procedures that are not in conflict with other
1442	provisions of the general statutes, for exercising the power of the
1443	exchange; and
1444	[(22)] (23) Do all acts and things necessary and convenient to carry
1445	out the purposes of the exchange, provided such acts or things shall no
1446	conflict with the provisions of the Affordable Care Act, regulations
1447	adopted thereunder or federal guidance issued pursuant to the
1448	Affordable Care Act.
1449	(d) The exchange shall submit any proposed assessment or fee to be
1450	charged to health carriers that are capable of offering a qualified health
1451	plan through the exchange, any proposed increase in the amount of any

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such assessment or fee to be imposed, any proposed method, or change in method, used to calculate any such assessment or fee to be implemented and any proposed severance or nondisclosure agreement to be entered into on or after October 1, 2021, to the joint standing committee of the General Assembly having cognizance of matters relating to insurance for the committee's review and approval. If the committee does not approve a submittal within sixty days after receiving the submittal, the proposed assessment, fee, increase, method, change in method or agreement, as the case may be, shall be deemed to have been rejected by the committee.

- [(d)] (e) (1) The chief executive officer of the exchange shall provide to the commissioner the name of any health carrier that fails to pay any assessment or user fee under subdivision (7) of subsection (c) of this section to the exchange. The commissioner shall see that all laws respecting the authority of the exchange pursuant to [said subdivision (7)] subdivisions (7) and (8) of subsection (c) of this section are faithfully executed. The commissioner has all the powers specifically granted under this title and all further powers that are reasonable and necessary to enable the commissioner to enforce the provisions of [said subdivision (7)] subdivisions (7) and (8) of subsection (c) of this section.
- (2) Any health carrier aggrieved by an administrative action taken by the commissioner under subdivision (1) of this subsection may appeal therefrom in accordance with the provisions of section 4-183, except venue for such appeal shall be in the judicial district of New Britain.
- Sec. 27. Subsection (b) of section 38a-1093 of the general statutes is repealed and the following is substituted in lieu thereof (*Effective October* 1478 1, 2021):
 - (b) Each subsidiary shall have and may exercise the powers of the exchange and such additional powers as are set forth in such resolution, except the powers of the exchange set forth in subdivisions (7), [(12), (15), (16), (17) and (21)] (8), (13), (16), (17), (18) and (22) of subsection (c) of section 38a-1083, as amended by this act, shall be reserved to the

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1484 exchange and shall not be exercisable by any subsidiary of the exchange.

- Sec. 28. (*Effective from passage*) (a) There is established a task force to study inequity in the provision of health insurance coverage and health care to minority populations in this state. Such study shall include, but need not be limited to, identifying any means available to promote equity in the provision of health insurance coverage and health care in this state.
- (b) The task force shall consist of the following members:

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- (1) Two appointed by the speaker of the House of Representatives, both of whom are individual consumers of health care and one of whom has purchased coverage through the Connecticut Health Insurance Exchange established pursuant to section 38a-1081 of the general statutes, as amended by this act;
- (2) Two appointed by the president pro tempore of the Senate, one of whom is a dentist licensed pursuant to chapter 379 of the general statutes who has experience working with minority patients at locations in this state that have an occurrence of dental decay that is greater than the state-wide average occurrence of dental decay;
- (3) Two appointed by the majority leader of the House of Representatives, one of whom is the director of a health care facility who has experience serving predominately minority populations and one of whom has experience analyzing data for a health insurer;
- 1506 (4) One appointed by the majority leader of the Senate, who is the 1507 director of a nonprofit business and has experience examining the 1508 causes of racial inequity in the provision of health care;
- 1509 (5) One appointed by the minority leader of the House of 1510 Representatives, who is an individual consumer of health care provided 1511 by state agencies;
- 1512 (6) One appointed by the minority leader of the Senate, who is a 1513 health care provider who has experience working with minority

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- patients at locations in this state that have occurrences of asthma, diabetes and prenatal death that are greater than the state-wide average
- 1516 occurrences of asthma, diabetes and prenatal death;
- 1517 (7) The Insurance Commissioner, or the commissioner's designee;
- 1518 (8) The Commissioner of Public Health, or the commissioner's 1519 designee;
- 1520 (9) The executive director of the Office of Health Strategy, or the 1521 executive director's designee; and
- 1522 (10) Two appointed by the Governor.

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- 1523 (c) All initial appointments to the task force shall be made not later 1524 than thirty days after the effective date of this section. Any vacancy shall 1525 be filled by the appointing authority.
- (d) The members of the task force shall select the chairpersons of the task force, from among the members of the task force, by a vote of the majority of the members of the task force. The Insurance Commissioner shall schedule the first meeting of the task force, which shall be held not later than sixty days after the effective date of this section.
 - (e) 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 task force.
 - (f) Not later than December 1, 2021, the task force shall submit a report on its findings and recommendations to the joint standing committee of the General Assembly having cognizance of matters relating to insurance, in accordance with the provisions of section 11-4a of the general statutes. The task force shall terminate on the date that it submits such report or December 1, 2021, whichever is later.

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

Section 1	July 1, 2021	19a-754a
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		T
Sec. 2	July 1, 2021	New section
Sec. 3	July 1, 2021	New section
Sec. 4	July 1, 2021	New section
Sec. 5	July 1, 2021	New section
Sec. 6	July 1, 2021	New section
Sec. 7	July 1, 2021	New section
Sec. 8	July 1, 2021	New section
Sec. 9	July 1, 2021	New section
Sec. 10	July 1, 2021	New section
Sec. 11	July 1, 2021	New section
Sec. 12	July 1, 2021	New section
Sec. 13	July 1, 2021	New section
Sec. 14	July 1, 2021	New section
Sec. 15	July 1, 2021	New section
Sec. 16	January 1, 2022	38a-8b
Sec. 17	January 1, 2022	5-259(m)(3)(C)
Sec. 18	January 1, 2022	7-464(c)(1)
Sec. 19	January 1, 2022	38a-465(18)(F)
Sec. 20	January 1, 2022	38a-465d(c)
Sec. 21	January 1, 2022	38a-478l(b)(2)(A)
Sec. 22	January 1, 2022	38a-720h(c)
Sec. 23	from passage	New section
Sec. 24	from passage	New section
Sec. 25	October 1, 2021	38a-1081
Sec. 26	October 1, 2021	38a-1083
Sec. 27	October 1, 2021	38a-1093(b)
Sec. 28	from passage	New section

Statement of Purpose:

To: (1) Require the Office of Health Strategy to (A) develop, innovate, direct and oversee health care delivery and payment models, (B) develop and adopt health care quality benchmarks, (C) enhance the transparency of health care entities, (D) monitor the development of accountable care organizations and patient-centered medical homes, and (E) monitor the adoption of alternative payment methodologies; (2) require the executive director of the Office of Health Strategy to (A) establish annual health care cost growth benchmarks and primary care targets, (B) submit an annual report to the General Assembly, (C) establish standards governing submission of data, information and documents by certain persons, (D) prepare and disclose an annual report concerning health status adjusted total medical expenses, (E) at

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least annually, submit a request to the federal Centers for Medicare and Medicaid Services for the health status adjusted total medical expenses of provider organizations that serve Medicare patients, (F) identify and examine any health care entity or payer that exceeds any annual health care cost growth benchmark and take enforcement action against such entity or payer, and (G) develop and adopt annual health care quality benchmarks for health care entities and payers; (3) require certain providers and provider organizations to annually submit certain data, information and documents to the Office of Health Strategy; (4) authorize the Office of Health Strategy to (A) enter into certain contractual agreements with third parties, and (B) adopt certain regulations; (5) subject to approval by the federal government, require the Commissioner of Consumer Protection to establish a Canadian legend drug importation program and authorize the commissioner, in consultation with the Commissioner of Public Health, to adopt regulations to implement such program; (6) adopt the Insurance Commissioner's recommendations concerning stop-loss insurance; (7) subject to approval by the federal government, require the Office of Health Strategy, in conjunction with the Office of Policy and Management, Insurance Department and Health Reinsurance Association, to establish a reinsurance program; (8) require the Auditors of Public Accounts to annually conduct an audit of certain health care plans administered or offered by this state and disclose the results of such audit to the General Assembly; (9) establish term limits for members of the board of directors of the Connecticut Health Insurance Exchange and require that members appointed or reappointed to the board have insurance expertise; (10) require the Connecticut Health Insurance Exchange to (A) conduct a public meeting before charging an assessment or user fee to certain health carriers, increasing the amount of any such assessment or fee or implementing or changing any process used to calculate any such assessment or fee, and (B) receive the approval of the joint standing committee of the General Assembly having cognizance of matters relating to insurance before (i) charging any assessment or user fee to certain health carriers, increasing the amount of any such assessment or fee or implementing or changing any process used to calculate any such assessment or fee, or (ii) entering into any nondisclosure or severance agreement; and (11) establish a task force to study inequity in the provision of health insurance coverage and health care to minority populations in this state.

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

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