

House _____ Amendment NO. _____

Offered By _____

1 AMEND Senate Substitute for Senate Committee Substitute for Senate Bill No. 767, Page 1, in the
2 Title, Line 2, by deleting the word "the"; and
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4 Further amend said page, bill, and section, Lines 3 through 4, by deleting all of said lines and
5 inserting in lieu thereof the words "health care"; and
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7 Further amend said bill and page, Section 44.035, Line 7, by inserting after all of said section and
8 line the following:
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10 "191.990. 1. The MO HealthNet division and the department of health and senior services
11 shall collaborate to coordinate goals and benchmarks in each agency's plans to reduce the incidence
12 of diabetes in Missouri, improve diabetes care, and control complications associated with diabetes.

13 2. The MO HealthNet division and the department of health and senior services shall submit
14 a report to the general assembly by January first of each odd-numbered year on the following:

15 (1) The prevalence and financial impact of diabetes of all types on the state of Missouri.
16 Items in this assessment shall include an estimate of the number of people with diagnosed and
17 undiagnosed diabetes, the number of individuals with diabetes impacted or covered by the agency
18 programs addressing diabetes, the financial impact of diabetes, and its complications on Missouri
19 based on the most recently published cost estimates for diabetes;

20 (2) An assessment of the benefits of implemented programs and activities aimed at
21 controlling diabetes and preventing the disease;

22 (3) A description of the level of coordination existing between the agencies, their contracted
23 partners, and other stakeholders on activities, programs, and messaging on managing, treating, or
24 preventing all forms of diabetes and its complications;

25 (4) The development or revision of detailed action plans for battling diabetes with a range of
26 actionable items for consideration by the general assembly. The plans shall identify proposed action
27 steps to reduce the impact of diabetes, prediabetes, and related diabetes complications. The plan also
28 shall identify expected outcomes of the action steps proposed in the following biennium while also
29 establishing benchmarks for controlling and preventing diabetes; and

30 (5) The development of a detailed budget blueprint identifying needs, costs, and resources
31 required to implement the plan identified in subdivision (4) of this subsection. This blueprint shall
32 include a budget range for all options presented in the plan identified in subdivision (4) of this
33 subsection for consideration by the general assembly.

34 3. The requirements of subsections 1 and 2 of this section shall be limited to diabetes
35 information, data, initiatives, and programs within each agency prior to the effective date of this
36 section, unless there is unobligated funding for diabetes in each agency that may be used for new

Action Taken _____ Date _____

research, data collection, reporting, or other requirements of subsections 1 and 2 of this section.

191.1140. 1. Subject to appropriations, the University of Missouri shall manage the “Show-Me Extension for Community Health Care Outcomes (ECHO) Program”. The department of health and senior services shall collaborate with the University of Missouri in utilizing the program to expand the capacity to safely and effectively treat chronic, common, and complex diseases in rural and underserved areas of the state and to monitor outcomes of such treatment.

2. The program is designed to utilize current telehealth technology to disseminate knowledge of best practices for the treatment of chronic, common, and complex diseases from a multidisciplinary team of medical experts to local primary care providers who will deliver the treatment protocol to patients, which will alleviate the need of many patients to travel to see specialists and will allow patients to receive treatment more quickly.

3. The program shall utilize local community health care workers with knowledge of local social determinants as a force multiplier to obtain better patient compliance and improved health outcomes.

196.990. 1. As used in this section, the following terms shall mean:

(1) "Administer", the direct application of an epinephrine auto-injector to the body of an individual;

(2) "Authorized entity", any entity or organization at or in connection with which allergens capable of causing anaphylaxis may be present, including but not limited to restaurants, recreation camps, youth sports leagues, amusement parks, and sports arenas;

(3) "Epinephrine auto-injector", a single-use device used for the automatic injection of a premeasured dose of epinephrine into the human body;

(4) "Physician", a physician licensed in this state under chapter 334;

(5) "Provide", the supply of one or more epinephrine auto-injectors to an individual;

(6) "Self-administration", a person's discretionary use of an epinephrine auto-injector.

2. A physician may prescribe epinephrine auto-injectors in the name of an authorized entity for use in accordance with this section, and pharmacists, physicians, and other persons authorized to dispense prescription medications may dispense epinephrine auto-injectors under a prescription issued in the name of an authorized entity.

3. An authorized entity may acquire and stock a supply of epinephrine auto-injectors under a prescription issued in accordance with this section. Such epinephrine auto-injectors shall be stored in a location readily accessible in an emergency and in accordance with the epinephrine auto-injector's instructions for use and any additional requirements established by the department of health and senior services by rule. An authorized entity shall designate employees or agents who have completed the training required under this section to be responsible for the storage, maintenance, and general oversight of epinephrine auto-injectors acquired by the authorized entity.

4. An employee or agent of an authorized entity or any other person who has completed the training required under this section may use epinephrine auto-injectors prescribed under this section on the premises of or in connection with the authorized entity to:

(1) Provide an epinephrine auto-injector to any individual who the employee, agent, or other person believes in good faith is experiencing anaphylaxis for immediate self-administration, regardless of whether the individual has a prescription for an epinephrine auto-injector or has previously been diagnosed with an allergy;

(2) Administer an epinephrine auto-injector to any individual who the employee, agent, or other person believes in good faith is experiencing anaphylaxis, regardless of whether the individual has a prescription for an epinephrine auto-injector or has previously been diagnosed with an allergy.

5. An employee, agent, or other person described in subsection 4 of this section shall successfully complete an anaphylaxis training program prior to providing or administering an

epinephrine auto-injector made available by an authorized entity and at least every two years following successful completion of the initial anaphylaxis training program. Such training shall be conducted by a nationally recognized organization experienced in training laypersons in emergency health treatment or other entity or person approved by the department of health and senior services. Training may be conducted online or in person and, at a minimum, shall cover:

(1) Techniques on how to recognize symptoms of severe allergic reactions, including anaphylaxis;

(2) Standards and procedures for the storage and administration of an epinephrine auto-injector; and

(3) Emergency follow-up procedures.

The entity that conducts the training shall issue a certificate, on a form developed or approved by the department of health and senior services, to each person who successfully completes the anaphylaxis training program.

6. The following persons and entities shall not be liable for any injuries or related damages that result from the administration of, or failure to administer an epinephrine auto-injector to a person eighteen years of age or older in accordance with this section that may constitute ordinary negligence:

(1) An authorized entity that either does or does not possess and make available epinephrine auto-injectors and its employees, agents, and other trained persons;

(2) Any person who uses an epinephrine auto-injector made available under this section;

(3) A physician that prescribes epinephrine auto-injectors to an authorized entity; or

(4) Any person or entity that conducts the training described in subsection 5 of this section.

Such immunity does not apply to acts or omissions constituting gross negligence or willful or wanton conduct. The administration of an epinephrine auto-injector in accordance with this section shall not be considered the practice of medicine. The immunity from liability provided under this subsection is in addition to and not in lieu of that provided under section 537.037. An authorized entity located in this state shall not be liable for any injuries or related damages that result from the provision or administration of an epinephrine auto-injector by its employees or agents outside of this state if the entity or its employees or agents are not liable for such injuries or related damages under the laws of the state in which such provision or administration occurred.

7. An authorized entity that possesses and makes available epinephrine auto-injectors shall submit to the department of health and senior services, on a form developed by the department, a report of each incident on the authorized entity's premises involving the administration of an epinephrine auto-injector. The department shall annually publish a report that summarizes all reports submitted to it under this subsection, but shall not include any identifying information regarding the persons to whom such epinephrine auto-injectors were administered.

8. An authorized entity that acquires a stock supply of epinephrine auto-injectors under a prescription issued in accordance with this section may make such epinephrine auto-injectors available to individuals other than the trained persons described in subsection 4 of this section if the epinephrine auto-injectors are stored in a locked secure container in accordance with manufacturer specifications and are made available only upon remote authorization by a physician via audio, televideo, or other similar means of electronic communication. Consultation with a physician for such purpose shall not be considered the practice of telemedicine or otherwise be construed as violating any law or rule regulating the physician's professional practice.

208.790. 1. The applicant shall have or intend to have a fixed place of residence in Missouri, with the present intent of maintaining a permanent home in Missouri for the indefinite future. The burden of establishing proof of residence within this state is on the applicant. The requirement also applies to persons residing in long-term care facilities located in the state of Missouri.

2. The department shall promulgate rules outlining standards for documenting proof of residence in Missouri. Documents used to show proof of residence shall include the applicant's name and address in the state of Missouri.

3. Applicant household income limits for eligibility shall be subject to appropriations, but in no event shall applicants have household income that is greater than one hundred eighty-five percent of the federal poverty level for the applicable family size for the applicable year as converted to the MAGI equivalent net income standard.

4. The department shall promulgate rules outlining standards for documenting proof of household income.

208.798. The provisions of sections 208.780 to 208.798 shall terminate on August 28, [2014] 2017.

338.010. 1. The "practice of pharmacy" means the interpretation, implementation, and evaluation of medical prescription orders, including any legend drugs under 21 U.S.C. Section 353; receipt, transmission, or handling of such orders or facilitating the dispensing of such orders; the designing, initiating, implementing, and monitoring of a medication therapeutic plan as defined by the prescription order so long as the prescription order is specific to each patient for care by a pharmacist; the compounding, dispensing, labeling, and administration of drugs and devices pursuant to medical prescription orders and administration of viral influenza, pneumonia, shingles, hepatitis A, hepatitis B, diphtheria, tetanus, pertussis, and meningitis vaccines by written protocol authorized by a physician for persons twelve years of age or older as authorized by rule or the administration of pneumonia, shingles, hepatitis A, hepatitis B, diphtheria, tetanus, pertussis, and meningitis vaccines by written protocol authorized by a physician for a specific patient as authorized by rule; the participation in drug selection according to state law and participation in drug utilization reviews; the proper and safe storage of drugs and devices and the maintenance of proper records thereof; consultation with patients and other health care practitioners, and veterinarians and their clients about legend drugs, about the safe and effective use of drugs and devices; and the offering or performing of those acts, services, operations, or transactions necessary in the conduct, operation, management and control of a pharmacy. No person shall engage in the practice of pharmacy unless he is licensed under the provisions of this chapter. This chapter shall not be construed to prohibit the use of auxiliary personnel under the direct supervision of a pharmacist from assisting the pharmacist in any of his or her duties. This assistance in no way is intended to relieve the pharmacist from his or her responsibilities for compliance with this chapter and he or she will be responsible for the actions of the auxiliary personnel acting in his or her assistance. This chapter shall also not be construed to prohibit or interfere with any legally registered practitioner of medicine, dentistry, or podiatry, or veterinary medicine only for use in animals, or the practice of optometry in accordance with and as provided in sections 195.070 and 336.220 in the compounding, administering, prescribing, or dispensing of his or her own prescriptions.

2. Any pharmacist who accepts a prescription order for a medication therapeutic plan shall have a written protocol from the physician who refers the patient for medication therapy services. The written protocol and the prescription order for a medication therapeutic plan shall come from the physician only, and shall not come from a nurse engaged in a collaborative practice arrangement under section 334.104, or from a physician assistant engaged in a supervision agreement under section 334.735.

3. Nothing in this section shall be construed as to prevent any person, firm or corporation from owning a pharmacy regulated by sections 338.210 to 338.315, provided that a licensed pharmacist is in charge of such pharmacy.

4. Nothing in this section shall be construed to apply to or interfere with the sale of nonprescription drugs and the ordinary household remedies and such drugs or medicines as are

1 normally sold by those engaged in the sale of general merchandise.

2 5. No health carrier as defined in chapter 376 shall require any physician with which they
3 contract to enter into a written protocol with a pharmacist for medication therapeutic services.

4 6. This section shall not be construed to allow a pharmacist to diagnose or independently
5 prescribe pharmaceuticals.

6 7. The state board of registration for the healing arts, under section 334.125, and the state
7 board of pharmacy, under section 338.140, shall jointly promulgate rules regulating the use of
8 protocols for prescription orders for medication therapy services and administration of viral
9 influenza vaccines. Such rules shall require protocols to include provisions allowing for timely
10 communication between the pharmacist and the referring physician, and any other patient protection
11 provisions deemed appropriate by both boards. In order to take effect, such rules shall be approved
12 by a majority vote of a quorum of each board. Neither board shall separately promulgate rules
13 regulating the use of protocols for prescription orders for medication therapy services and
14 administration of viral influenza vaccines. Any rule or portion of a rule, as that term is defined in
15 section 536.010, that is created under the authority delegated in this section shall become effective
16 only if it complies with and is subject to all of the provisions of chapter 536 and, if applicable,
17 section 536.028. This section and chapter 536 are nonseverable and if any of the powers vested with
18 the general assembly pursuant to chapter 536 to review, to delay the effective date, or to disapprove
19 and annul a rule are subsequently held unconstitutional, then the grant of rulemaking authority and
20 any rule proposed or adopted after August 28, 2007, shall be invalid and void.

21 8. The state board of pharmacy may grant a certificate of medication therapeutic plan
22 authority to a licensed pharmacist who submits proof of successful completion of a board-approved
23 course of academic clinical study beyond a bachelor of science in pharmacy, including but not
24 limited to clinical assessment skills, from a nationally accredited college or university, or a
25 certification of equivalence issued by a nationally recognized professional organization and
26 approved by the board of pharmacy.

27 9. Any pharmacist who has received a certificate of medication therapeutic plan authority
28 may engage in the designing, initiating, implementing, and monitoring of a medication therapeutic
29 plan as defined by a prescription order from a physician that is specific to each patient for care by a
30 pharmacist.

31 10. Nothing in this section shall be construed to allow a pharmacist to make a therapeutic
32 substitution of a pharmaceutical prescribed by a physician unless authorized by the written protocol
33 or the physician's prescription order.

34 11. "Veterinarian", "doctor of veterinary medicine", "practitioner of veterinary medicine",
35 "DVM", "VMD", "BVSe", "BVMS", "BSe (VetScience)", "VMB", "MRCVS", or an equivalent title
36 means a person who has received a doctor's degree in veterinary medicine from an accredited school
37 of veterinary medicine or holds an Educational Commission for Foreign Veterinary Graduates
38 (EDFVG) certificate issued by the American Veterinary Medical Association (AVMA).

39 12. In addition to other requirements established by the joint promulgation of rules by the
40 board of pharmacy and the state board of registration for the healing arts:

41 (1) A pharmacist shall administer vaccines in accordance with treatment guidelines
42 established by the Centers for Disease Control and Prevention (CDC);

43 (2) A pharmacist who is administering a vaccine shall request a patient to remain in the
44 pharmacy a safe amount of time after administering the vaccine to observe any adverse reactions.
45 Such pharmacist shall have adopted emergency treatment protocols;

46 (3) In addition to other requirements by the board, a pharmacist shall receive additional
47 training as required by the board and evidenced by receiving a certificate from the board upon
48 completion, and shall display the certification in his or her pharmacy where vaccines are delivered.

13. A pharmacist shall provide a written report within fourteen days of administration of a vaccine to the patient's primary health care provider, if provided by the patient, containing:

- (1) The identity of the patient;
- (2) The identity of the vaccine or vaccines administered;
- (3) The route of administration;
- (4) The anatomic site of the administration;
- (5) The dose administered; and
- (6) The date of administration.

338.059. 1. It shall be the duty of a licensed pharmacist or a physician to affix or have affixed by someone under the pharmacist's or physician's supervision a label to each and every container provided to a consumer in which is placed any prescription drug upon which is typed or written the following information:

- (1) The date the prescription is filled;
- (2) The sequential number or other unique identifier;
- (3) The patient's name;
- (4) The prescriber's directions for usage;
- (5) The prescriber's name;
- (6) The name and address of the pharmacy;
- (7) The exact name and dosage of the drug dispensed;
- (8) There may be one line under the information provided in subdivisions (1) to (7) of this subsection stating "Refill" with a blank line or squares following or the words "No Refill";
- (9) When a generic substitution is dispensed, the name of the manufacturer or an abbreviation thereof shall appear on the label or in the pharmacist's records as required in section 338.100.

2. The label of any drug which is sold at wholesale in this state and which requires a prescription to be dispensed at retail shall contain the name of the manufacturer, expiration date, if applicable, batch or lot number and national drug code.

338.165. 1. As used in this section, the following terms mean:

- (1) "Board", the Missouri board of pharmacy;
- (2) "Hospital", a hospital as defined in section 197.020;
- (3) "Hospital clinic or facility", a clinic or facility under the common control, management, or ownership of the same hospital or hospital system;
- (4) "Medical staff committee", the committee or other body of a hospital or hospital system responsible for formulating policies regarding pharmacy services and medication management;
- (5) "Medication order", an order for a legend drug or device that is:
 - (a) Authorized or issued by an authorized prescriber acting within the scope of his or her professional practice or pursuant to a protocol or standing order approved by the medical staff committee; and
 - (b) To be distributed or administered to the patient by a health care practitioner or lawfully authorized designee at a hospital or a hospital clinic or facility;
- (6) "Patient", an individual receiving medical diagnosis, treatment or care at a hospital or a hospital clinic or facility.

2. The department of health and senior services shall have sole authority and responsibility for the inspection and licensure of hospitals as provided by chapter 197 including, but not limited to all parts, services, functions, support functions and activities which contribute directly or indirectly to patient care of any kind whatsoever. However, the board may inspect a class B pharmacy or any portion thereof that is not under the inspection authority vested in the department of health and senior services by chapter 197 to determine compliance with this chapter or the rules of the board.

1 This section shall not be construed to bar the board from conducting an investigation pursuant to a
2 public or governmental complaint to determine compliance by an individual licensee or registrant of
3 the board with any applicable provisions of this chapter or the rules of the board.

4 3. The department of health and senior services shall have authority to promulgate rules in
5 conjunction with the board governing medication distribution and the provision of medication
6 therapy services by a pharmacist at or within a hospital. Rules may include, but are not limited to,
7 medication management, preparation, compounding, administration, storage, distribution, packaging
8 and labeling. Until such rules are jointly promulgated, hospitals shall comply with all applicable
9 state law and department of health and senior services rules governing pharmacy services and
10 medication management in hospitals. The rulemaking authority granted herein to the department of
11 health and senior services shall not include the dispensing of medication by prescription.

12 4. All pharmacists providing medication therapy services shall obtain a certificate of
13 medication therapeutic plan authority as provided by rule of the board. Medication therapy services
14 may be provided by a pharmacist for patients of a hospital pursuant to a protocol with a physician as
15 required by section 338.010 or pursuant to a protocol approved by the medical staff committee.
16 However, the medical staff protocol shall include a process whereby an exemption to the protocol for
17 a patient may be granted for clinical efficacy should the patient's physician make such request. The
18 medical staff protocol shall also include an appeals process to request a change in a specific protocol
19 based on medical evidence presented by a physician on staff.

20 5. Medication may be dispensed by a class B hospital pharmacy pursuant to a prescription or
21 a medication order.

22 6. A drug distributor license shall not be required to transfer medication from a class B
23 hospital pharmacy to a hospital clinic or facility for patient care or treatment.

24 7. Medication dispensed by a class A pharmacy located in a hospital to a hospital patient for
25 use or administration outside of the hospital under a medical staff-approved protocol for medication
26 therapy shall be dispensed only by a prescription order for medication therapy from an individual
27 physician for a specific patient.

28 8. Medication dispensed by a hospital to a hospital patient for use or administration outside
29 of the hospital shall be labeled as provided by rules jointly promulgated by the department of health
30 and senior services and the board including, medication distributed for administration by or under
31 the supervision of a health care practitioner at a hospital clinic or facility.

32 9. This section shall not be construed to preempt any law or rule governing controlled
33 substances.

34 10. Any rule, as that term is defined in section 536.010, that is created under the authority
35 delegated in this section shall only become effective if it complies with and is subject to all of the
36 provisions of chapter 536 and, if applicable, section 536.028. This section and chapter 536 are
37 nonseverable and if any of the powers vested with the general assembly under chapter 536 to review,
38 to delay the effective date, or to disapprove and annul a rule are subsequently held unconstitutional,
39 then the grant of rulemaking authority and any rule proposed or adopted after August 28, 2014, shall
40 be invalid and void.

41 11. The board shall appoint an advisory committee to review and make recommendations to
42 the board on the merit of all rules and regulations to be jointly promulgated by the board and the
43 department of health and senior services pursuant to the joint rulemaking authority granted by this
44 section. The advisory committee shall consist of:

45 (1) Two representatives designated by the Missouri Hospital Association, one of whom shall
46 be a pharmacist;

47 (2) One pharmacist designated by the Missouri Society of Health System Pharmacists;

48 (3) One pharmacist designated by the Missouri Pharmacy Association;

(4) One pharmacist designated by the department of health and senior services from a hospital with a licensed bed count that does not exceed fifty beds or from a critical access hospital as defined by the department of social services for purposes of MO HealthNet reimbursement;

(5) One pharmacist designated by the department of health and senior services from a hospital with a licensed bed count that exceeds two hundred beds; and

(6) One pharmacist designated by the Board with experience in the provision of hospital pharmacy services.

12. Nothing in this section shall be construed to limit the authority of a licensed health care provider to prescribe, administer, or dispense medications and treatments within the scope of their professional practice.

338.220. 1. It shall be unlawful for any person, copartnership, association, corporation or any other business entity to open, establish, operate, or maintain any pharmacy as defined by statute without first obtaining a permit or license to do so from the Missouri board of pharmacy. A permit shall not be required for an individual licensed pharmacist to perform nondispensing activities outside of a pharmacy, as provided by the rules of the board. A permit shall not be required for an individual licensed pharmacist to administer drugs, vaccines, and biologicals by protocol, as permitted by law, outside of a pharmacy. The following classes of pharmacy permits or licenses are hereby established:

- (1) Class A: Community/ambulatory;
- (2) Class B: Hospital [outpatient] pharmacy;
- (3) Class C: Long-term care;
- (4) Class D: Nonsterile compounding;
- (5) Class E: Radio pharmaceutical;
- (6) Class F: Renal dialysis;
- (7) Class G: Medical gas;
- (8) Class H: Sterile product compounding;
- (9) Class I: Consultant services;
- (10) Class J: Shared service;
- (11) Class K: Internet;
- (12) Class L: Veterinary;
- (13) Class M: Specialty (bleeding disorder);
- (14) Class N: Automated dispensing system (health care facility);
- (15) Class O: Automated dispensing system (ambulatory care);
- (16) Class P: Practitioner office/clinic.

2. Application for such permit or license shall be made upon a form furnished to the applicant; shall contain a statement that it is made under oath or affirmation and that its representations are true and correct to the best knowledge and belief of the person signing same, subject to the penalties of making a false affidavit or declaration; and shall be accompanied by a permit or license fee. The permit or license issued shall be renewable upon payment of a renewal fee. Separate applications shall be made and separate permits or licenses required for each pharmacy opened, established, operated, or maintained by the same owner.

3. All permits, licenses or renewal fees collected pursuant to the provisions of sections 338.210 to 338.370 shall be deposited in the state treasury to the credit of the Missouri board of pharmacy fund, to be used by the Missouri board of pharmacy in the enforcement of the provisions of sections 338.210 to 338.370, when appropriated for that purpose by the general assembly.

4. Class L: veterinary permit shall not be construed to prohibit or interfere with any legally registered practitioner of veterinary medicine in the compounding, administering, prescribing, or dispensing of their own prescriptions, or medicine, drug, or pharmaceutical product to be used for

1 animals.

2 5. Except for any legend drugs under 21 U.S.C. Section 353, the provisions of this section
3 shall not apply to the sale, dispensing, or filling of a pharmaceutical product or drug used for treating
4 animals.

5 6. A "Class B Hospital Pharmacy" shall be defined as a pharmacy owned, managed, or
6 operated by a hospital as defined by section 197.020 or a clinic or facility under common control,
7 management or ownership of the same hospital or hospital system. This section shall not be
8 construed to require a class B hospital pharmacy permit or license for hospitals solely providing
9 services within the practice of pharmacy under the jurisdiction of, and the licensure granted by, the
10 department of health and senior services pursuant to chapter 197.

11 7. Upon application to the board, any hospital that holds a pharmacy permit or license on the
12 effective date of this section shall be entitled to obtain a class B pharmacy permit or license without
13 fee, provided such application shall be submitted to the board on or before January 1, 2015."; and

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15 Further amend said bill by amending the title, enacting clause, and intersectional references
16 accordingly.
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