As Amended by Senate Committee

Session of 2021

SENATE BILL No. 200

By Committee on Public Health and Welfare

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AN ACT concerning pharmacists and pharmacy; relating to the state board 1 of pharmacy; expanding the pharmacist's scope of practice to include 2 point-of-care testing for and treatment of initiation of therapy for 3 4 certain health conditions; authorizing the collaborative drug therapy 5 management advisory committee to adopt a statewide protocol for such therapy; amending K.S.A. 65-1626a and repealing the existing 6 7 section. 8 9 Be it enacted by the Legislature of the State of Kansas: 10 New Section 1. A pharmacist may independently test or screen for and initiate therapy pursuant to a state-wide protocol established by the 11 state board of pharmacy for health conditions of individuals who are-12 eligible to receive the testing or servening service. For purposes of this 13 14 section, a health condition is a condition that is generally managed with minimal treatment or self-care and includes, but is not limited to: 15 16 (a) Influenza: 17 (b) streptococcus; (c) COVID-19; 18 19 (d) pre-exposure prophylaxis; 20 (e) post-exposure prophylaxis; or (f) a condition for which its diagnostic test is waived under the-21 22 federal clinical laboratory improvement amendments of 1988. 23 New Section 1. (a) A pharmacist may initiate therapy within the 24 framework of a statewide protocol for the following health conditions: 25 (1) Influenza; 26 (2) streptococcal pharyngitis; or 27 (3) urinary tract infection. 28 (b) The collaborative drug therapy management advisory committee established pursuant to K.S.A. 65-1677, and amendments thereto, may 29 adopt a statewide protocol for each condition listed in subsection (a). 30 31 The committee shall consider appropriateness of therapy based upon: 32 (1) The patient's age; 33 (2) approval by the United States food and drug administration;

34 (3) in-person interactions; and <u>(4) the method of administration</u> In establishing such statewide
 protocols, the committee shall specify:

- 3 (1) The medications or categories of medications included in the 4 protocol for each health condition;
- 5 (2) the training or qualifications required for pharmacists to 6 implement the protocols;

7 (3) requirements for documentation and maintenance of records,
8 including patient inclusion and exclusion criteria, medical referral
9 criteria, patient assessment tools based on current clinical guidelines,
10 follow-up monitoring or care plans and the pharmacist's adherence to
11 the applicable protocols; and

(4) communication requirements, including, but not limited to,
 notification to the patient's personal or primary care provider.

14 (c) The board may deny an application or renewal or revoke or 15 suspend the license of a pharmacist upon a finding that the 16 pharmacist has violated the provisions of this section or failed to 17 practice within the framework of statewide protocols established 18 pursuant to this section by the collaborative drug therapy 19 management advisory committee.

Sec. 2. K.S.A. 65-1626a is hereby amended to read as follows: 65-1626a. (a) For the purpose of the pharmacy act of the state of Kansas, the following-persons *individuals* shall be deemed to be engaged in the practice of pharmacy:

(1) PersonsIndividuals who publicly profess to be a pharmacist, or
 publicly profess to assume the duties incident to being a pharmacist and
 their knowledge of drugs or drug actions, or both; and

(2) personsindividuals who attach to their name any words or
abbreviation indicating that they are a pharmacist licensed to practice
pharmacy in Kansas.

30 (b) As used in this section:

31 (1) "Practice of pharmacy" means:

(A) The interpretation and evaluation of prescription orders;

(B) the compounding, dispensing and labeling of drugs and devices
 pursuant to prescription orders;

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(C) the administering of vaccine pursuant to a vaccination protocol;

(D) the participation in drug selection according to state law and
 participation in drug utilization reviews;

(E) the proper and safe storage of prescription drugs and prescription
 devices and the maintenance of proper records thereof in accordance with
 law;

41 *(F)* consultation with patients and other health care practitioners 42 about the safe and effective use of prescription drugs and prescription 43 devices; (G) performance of collaborative drug therapy management pursuant
 to a written collaborative practice agreement with one or more physicians
 who have an established physician-patient relationship;-and

4 *(H)* participation in the offering or performing of those acts, services, 5 operations or transactions necessary in the conduct, operation, 6 management and control of a pharmacy; *and*

7 (I) the initiation of drugs, drug categories or devices that are-8 initiated in accordance with the product's federal food and drug-9 administration-approved labeling and limited to conditions listed below-10 that:

11 12 *(i) Do not require a new diagnosis;*

(ii) are minor and generally self-limiting;

(iii) have a test that is used to guide diagnosis or clinical decision making and is waived under the federal clinical laboratory improvement
 amendments of 1988; or

(iv) in the professional judgment of the pharmacist, threaten the health or safety of the patient should the prescription not be immediately.
 dispensed. In such cases, only sufficient quantity may be provided until the
 patient is able to be seen by another provider initiation of therapy for the

20 conditions specified in section 1, and amendments thereto.

21 Nothing in this section shall be construed to add any additional-22 requirements for registration or for a permit under the pharmaey act of the 23 state of Kansas or for approval under subsection (g) of K.S.A. 65-1643; and amendments thereto, or to prevent persons other than pharmacists-24 from engaging in drug utilization review, or to require persons lawfully in 25 possession of prescription drugs or prescription devices to meet any-26 27 storage or record keeping requirements except such storage and record keeping requirements as may be otherwise provided by law or to affect any 28 29 person consulting with a health care practitioner about the safe andeffective use of prescription drugs or prescription devices. 30

(2) "Collaborative drug therapy management" means a practice of 31 pharmacy where a pharmacist performs certain pharmaceutical-related 32 patient care functions for a specific patient which have been delegated to 33 the pharmacist by a physician through a collaborative practice agreement. 34 A physician who enters into a collaborative practice agreement is 35 36 responsible for the care of the patient following initial diagnosis and 37 assessment and for the direction and supervision of the pharmacist 38 throughout the collaborative drug therapy management process. Nothing in 39 this subsection shall be construed to permit a pharmacist to alter a physician's orders or directions, diagnose or treat any disease, 40 41 independently prescribe drugs or independently practice medicine and 42 surgery.

(3) "Collaborative practice agreement" means a written agreement or

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protocol between one or more pharmacists and one or more physicians that 1 provides for collaborative drug therapy management. Such collaborative 2 3 practice agreement shall contain certain specified conditions or limitations 4 pursuant to the collaborating physician's order, standing order, delegation 5 or protocol. A collaborative practice agreement shall be: (A) Consistent 6 with the normal and customary specialty, competence and lawful practice 7 of the physician; and (B) appropriate to the pharmacist's training and 8 experience.

9 (4) "Physician" means a person licensed to practice medicine and 10 surgery in this state.

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(c) Nothing in this section shall be construed to:

(1) Add any additional requirements for registration or for a permit
under the pharmacy act of the state of Kansas or for approval under
K.S.A. 65-1643(g), and amendments thereto;

15 (2) prevent persons other than pharmacists from engaging in drug 16 utilization review;

(3) require persons lawfully in possession of prescription drugs or
prescription devices to meet any storage or record keeping requirements
except such storage and record keeping requirements as may be otherwise
provided by law; or

(4) affect any person consulting with a healthcare practitioner about the safe and effective use of prescription drugs or prescription devices.

23 Sec. 3. K.S.A. 65-1626a is hereby repealed.

24 Sec. 4. This act shall take effect and be in force from and after its 25 publication in the statute book.