LEGISLATURE OF NEBRASKA

ONE HUNDRED NINTH LEGISLATURE

FIRST SESSION

LEGISLATIVE BILL 697

Read first time January 22, 2025

Committee: Health and Human Services

- 1 A BILL FOR AN ACT relating to the Pharmacy Practice Act; to amend
- 2 sections 38-2849 and 38-2884, Reissue Revised Statutes of Nebraska,
- and section 38-2867.01, Revised Statutes Cumulative Supplement,
- 4 2024; to change requirements for the Board of Pharmacy; to change
- 5 requirements relating to compounding and delegated dispensing
- 6 permits; and to repeal the original sections.
- 7 Be it enacted by the people of the State of Nebraska,

- **Section 1.** Section 38-2849, Reissue Revised Statutes of Nebraska, is
- 2 amended to read:
- 3 38-2849 The board shall be composed of eight five members, including
- 4 five four actively practicing pharmacists, at least one of whom practices
- 5 within the confines of a hospital, one pharmacy technician, and two one
- 6 public members member who are is interested in the health of the people
- 7 of Nebraska.
- 8 Sec. 2. Section 38-2867.01, Revised Statutes Cumulative Supplement,
- 9 2024, is amended to read:
- 10 38-2867.01 (1) Any person authorized to compound shall compound in
- 11 compliance with the standards of chapters 795 and 797 of The United
- 12 States Pharmacopeia and The National Formulary, as such chapters existed
- 13 on January 1, 2023, and shall compound (a) as the result of a
- 14 practitioner's medical order or initiative occurring in the course of
- 15 practice based upon the relationship between the practitioner, patient,
- 16 and pharmacist, (b) for the purpose of, or as an incident to, research,
- 17 teaching, or chemical analysis and not for sale or dispensing, or (c) for
- 18 office use only and not for resale by an outsourcing facility operating
- 19 pursuant to 21 U.S.C. 353b or section 71-470.
- 20 (2) Compounding in a hospital pharmacy may occur for any hospital
- 21 which is part of the same health care system under common ownership or
- 22 which is a member of or an affiliated member of a formal network or
- 23 partnership agreement.
- 24 (3)(a) Any authorized person may reconstitute a commercially
- 25 available drug product in accordance with directions contained in
- 26 approved labeling provided by the product's manufacturer and other
- 27 manufacturer directions consistent with labeling.
- 28 (b) Any authorized person using beyond-use dating must follow the
- 29 approved product manufacturer's labeling or the standards of The United
- 30 States Pharmacopeia and The National Formulary if the product
- 31 manufacturer's labeling does not specify beyond-use dating.

- 1 (c) Any authorized person engaged in activities listed in this
- 2 subsection is not engaged in compounding, except that any variance from
- 3 the approved product manufacturer's labeling will result in the person
- 4 being engaged in compounding.
- 5 (4) Any authorized person splitting a scored tablet along scored
- 6 lines or adding flavoring to a commercially available drug product is not
- 7 engaged in compounding.
- 8 (5) No person shall compound:
- 9 (a) A drug that has been identified by the federal Food and Drug
- 10 Administration as withdrawn or removed from the market because the drug
- 11 was found to be unsafe or ineffective;
- 12 (b) A drug that is essentially a copy of an approved drug unless
- 13 there is a drug shortage as determined by the board or unless a patient
- 14 has an allergic reaction to the approved drug; or
- 15 (c) A drug that has been identified by the federal Food and Drug
- 16 Administration or the board as a product which may not be compounded.
- 17 Sec. 3. Section 38-2884, Reissue Revised Statutes of Nebraska, is
- 18 amended to read:
- 19 38-2884 Under a delegated dispensing permit for a public health
- 20 clinic, approved formulary drugs and devices may be dispensed by a public
- 21 health clinic worker or a health care professional licensed in Nebraska
- 22 to practice medicine and surgery or licensed in Nebraska as a registered
- 23 nurse, licensed practical nurse, or physician assistant without the
- 24 onsite services of a pharmacist if:
- 25 (1) The initial dispensing of all prescriptions for approved
- 26 formulary drugs and devices is conducted by a health care professional
- 27 licensed in Nebraska to practice medicine and surgery or pharmacy or
- 28 licensed in Nebraska as a registered nurse, licensed practical nurse, or
- 29 physician assistant;
- 30 (2) The drug or device is dispensed pursuant to a prescription
- 31 written onsite by a practitioner or by a practitioner licensed in

1 Nebraska working in affiliation with a public health clinic pursuant to a

- 2 delegated dispensing permit;
- 3 (3) The only prescriptions to be refilled under the delegated
- 4 dispensing permit are prescriptions for contraceptives;
- 5 (4) Prescriptions are accompanied by patient instructions and
- 6 written information approved by the director;
- 7 (5) The dispensing of authorized refills of contraceptives is done
- 8 by a licensed health care professional listed in subdivision (1) of this
- 9 section or by a public health clinic worker;
- 10 (6) All drugs or devices are prepackaged by the manufacturer or at a
- 11 public health clinic by a pharmacist into the quantity to be prescribed
- 12 and dispensed at the public health clinic;
- 13 (7) All drugs and devices stored, received, or dispensed under the
- 14 authority of public health clinics are properly labeled at all times. For
- 15 purposes of this subdivision, properly labeled means that the label
- 16 affixed to the container prior to dispensing contains the following
- 17 information:
- 18 (a) The name of the manufacturer;
- 19 (b) The lot number and expiration date from the manufacturer or, if
- 20 repackaged by a pharmacist, the lot number and calculated expiration
- 21 date;
- 22 (c) Directions for patient use;
- 23 (d) The quantity of drug in the container;
- 24 (e) The name, strength, and dosage form of the drug; and
- 25 (f) Auxiliary labels as needed for proper adherence to any
- 26 prescription;
- 27 (8) The following additional information is added to the label of
- 28 each container when the drug or device is dispensed:
- 29 (a) The patient's name;
- 30 (b) The name of the prescribing health care professional;
- 31 (c) The prescription number;

LB697 2025

- 1 (d) The date dispensed; and
- 2 (e) The name and address of the public health clinic;
- 3 (9) The only drugs and devices allowed to be dispensed or stored by
- 4 public health clinics appear on the formulary approved pursuant to
- 5 section 38-2881; and
- 6 (10) At any time that dispensing is occurring from a public health
- 7 clinic, the delegating pharmacist for the public health clinic or on-call
- 8 pharmacist in Nebraska is available, either in person or by telephone, to
- 9 answer questions from clients, staff, public health clinic workers, or
- 10 volunteers. This availability shall be confirmed and documented at the
- 11 beginning of each day that dispensing will occur. The delegating
- 12 pharmacist or on-call pharmacist shall inform the public health clinic if
- 13 he or she will not be available during the time that his or her
- 14 availability is required. If a pharmacist is unavailable, no dispensing
- 15 shall occur.
- 16 Sec. 4. Original sections 38-2849 and 38-2884, Reissue Revised
- 17 Statutes of Nebraska, and section 38-2867.01, Revised Statutes Cumulative
- 18 Supplement, 2024, are repealed.