

LEGISLATURE OF NEBRASKA  
ONE HUNDRED NINTH LEGISLATURE  
FIRST SESSION

**LEGISLATIVE BILL 697**

Introduced by Strommen, 47; Bosn, 25; DeBoer, 10; DeKay, 40; Dover, 19;  
Dungan, 26; Fredrickson, 20; Guereca, 7; Hallstrom, 1;  
Holdcroft, 36; Hunt, 8; Ibach, 44; Kauth, 31; Lonowski, 33;  
Quick, 35; Rountree, 3; Sorrentino, 39; Spivey, 13; Storer,  
43.

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,  
3 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,

1           **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;

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.