

LEGISLATURE OF NEBRASKA
ONE HUNDRED SIXTH LEGISLATURE
SECOND SESSION

LEGISLATIVE BILL 922

Introduced by Kolterman, 24.

Read first time January 10, 2020

Committee: Health and Human Services

1 A BILL FOR AN ACT relating to prescriptions; to amend sections 28-414,
2 28-414.01, 38-2870, and 38-2891, Revised Statutes Cumulative
3 Supplement, 2018, and section 38-101, Revised Statutes Supplement,
4 2019; to define a term; to require electronic issuance of
5 prescriptions for controlled substances; to provide exceptions; to
6 harmonize provisions; to provide an operative date; and to repeal
7 the original sections.
8 Be it enacted by the people of the State of Nebraska,

1 Section 1. Section 28-414, Revised Statutes Cumulative Supplement,
2 2018, is amended to read:

3 28-414 (1) Except as otherwise provided in this section or section
4 28-412 or when administered directly by a practitioner to an ultimate
5 user, a controlled substance listed in Schedule II of section 28-405
6 shall not be dispensed without a prescription from a practitioner
7 authorized to prescribe. Beginning January 1, 2021, all such
8 prescriptions shall be subject to section 4 of this act. No prescription
9 for a controlled substance listed in Schedule II of section 28-405 shall
10 be filled more than six months from the date of issuance. A prescription
11 for a controlled substance listed in Schedule II of section 28-405 shall
12 not be refilled.

13 (2) A prescription for controlled substances listed in Schedule II
14 of section 28-405 must contain the following information prior to being
15 filled by a pharmacist or dispensing practitioner: (a) Patient's name and
16 address, (b) name of the drug, device, or biological, (c) strength of the
17 drug or biological, if applicable, (d) dosage form of the drug or
18 biological, (e) quantity of the drug, device, or biological prescribed,
19 (f) directions for use, (g) date of issuance, (h) prescribing
20 practitioner's name and address, and (i) Drug Enforcement Administration
21 number of the prescribing practitioner. If the prescription is a written
22 paper prescription, the paper prescription must contain the prescribing
23 practitioner's manual signature. If the prescription is an electronic
24 prescription, the electronic prescription must contain all of the
25 elements in subdivisions (a) through (i) of this subsection, must be
26 digitally signed, and must be transmitted to and received by the pharmacy
27 electronically to meet all of the requirements of the Controlled
28 Substances Act, 21 U.S.C. 801 et seq., as it existed on January 1, 2014,
29 pertaining to electronic prescribing of controlled substances.

30 (3)(a) In emergency situations, a controlled substance listed in
31 Schedule II of section 28-405 may be dispensed pursuant to an oral

1 prescription reduced to writing in accordance with subsection (2) of this
2 section, except for the prescribing practitioner's signature, and bearing
3 the word "emergency".

4 (b) For purposes of this section, emergency situation means a
5 situation in which a prescribing practitioner determines that (i)
6 immediate administration of the controlled substance is necessary for
7 proper treatment of the patient, (ii) no appropriate alternative
8 treatment is available, including administration of a drug which is not a
9 controlled substance listed in Schedule II of section 28-405, and (iii)
10 it is not reasonably possible for the prescribing practitioner to provide
11 a signed, written or electronic prescription to be presented to the
12 person dispensing the controlled substance prior to dispensing.

13 (4)(a) In nonemergency situations:

14 (i) A controlled substance listed in Schedule II of section 28-405
15 may be dispensed pursuant to a facsimile of a written, signed paper
16 prescription if the original written, signed paper prescription is
17 presented to the pharmacist for review before the controlled substance is
18 dispensed, except as provided in subdivision (a)(ii) or (iii) of this
19 subsection;

20 (ii) A narcotic drug listed in Schedule II of section 28-405 may be
21 dispensed pursuant to a facsimile of a written, signed paper prescription
22 (A) to be compounded for direct parenteral administration to a patient
23 for the purpose of home infusion therapy or (B) for administration to a
24 patient enrolled in a hospice care program and bearing the words "hospice
25 patient"; and

26 (iii) A controlled substance listed in Schedule II of section 28-405
27 may be dispensed pursuant to a facsimile of a written, signed paper
28 prescription for administration to a resident of a long-term care
29 facility.

30 (b) For purposes of subdivisions (a)(ii) and (iii) of this
31 subsection, a facsimile of a written, signed paper prescription shall

1 serve as the original written prescription and shall be maintained in
2 accordance with subsection (1) of section 28-414.03.

3 (5)(a) A prescription for a controlled substance listed in Schedule
4 II of section 28-405 may be partially filled if the pharmacist does not
5 supply the full quantity prescribed and he or she makes a notation of the
6 quantity supplied on the face of the prescription or in the electronic
7 record. The remaining portion of the prescription may be filled no later
8 than thirty days after the date on which the prescription is written. The
9 pharmacist shall notify the prescribing practitioner if the remaining
10 portion of the prescription is not or cannot be filled within such
11 period. No further quantity may be supplied after such period without a
12 new written, signed paper prescription or electronic prescription.

13 (b) A prescription for a controlled substance listed in Schedule II
14 of section 28-405 written for a patient in a long-term care facility or
15 for a patient with a medical diagnosis documenting a terminal illness may
16 be partially filled. Such prescription shall bear the words "terminally
17 ill" or "long-term care facility patient" on its face or in the
18 electronic record. If there is any question whether a patient may be
19 classified as having a terminal illness, the pharmacist shall contact the
20 prescribing practitioner prior to partially filling the prescription.
21 Both the pharmacist and the prescribing practitioner have a corresponding
22 responsibility to assure that the controlled substance is for a
23 terminally ill patient. For each partial filling, the dispensing
24 pharmacist shall record on the back of the prescription or on another
25 appropriate record, uniformly maintained and readily retrievable, the
26 date of the partial filling, quantity dispensed, remaining quantity
27 authorized to be dispensed, and the identification of the dispensing
28 pharmacist. The total quantity of controlled substances listed in
29 Schedule II which is dispensed in all partial fillings shall not exceed
30 the total quantity prescribed. A prescription for a Schedule II
31 controlled substance for a patient in a long-term care facility or a

1 patient with a medical diagnosis documenting a terminal illness is valid
2 for sixty days from the date of issuance or until discontinuance of the
3 prescription, whichever occurs first.

4 Sec. 2. Section 28-414.01, Revised Statutes Cumulative Supplement,
5 2018, is amended to read:

6 28-414.01 (1) Except as otherwise provided in this section or when
7 administered directly by a practitioner to an ultimate user, a controlled
8 substance listed in Schedule III, IV, or V of section 28-405 shall not be
9 dispensed without a written, oral, or electronic medical order. Such
10 medical order is valid for six months after the date of issuance.
11 Original prescription information for any controlled substance listed in
12 Schedule III, IV, or V of section 28-405 may be transferred between
13 pharmacies for purposes of refill dispensing pursuant to section 38-2871.

14 (2) A prescription for controlled substances listed in Schedule III,
15 IV, or V of section 28-405 must contain the following information prior
16 to being filled by a pharmacist or dispensing practitioner: (a) Patient's
17 name and address, (b) name of the drug, device, or biological, (c)
18 strength of the drug or biological, if applicable, (d) dosage form of the
19 drug or biological, (e) quantity of the drug, device, or biological
20 prescribed, (f) directions for use, (g) date of issuance, (h) number of
21 refills, including pro re nata or PRN refills, not to exceed five refills
22 within six months after the date of issuance, (i) prescribing
23 practitioner's name and address, and (j) Drug Enforcement Administration
24 number of the prescribing practitioner. Beginning January 1, 2021, all
25 such prescriptions shall be subject to section 4 of this act. If the
26 prescription is a written paper prescription, the paper prescription must
27 contain the prescribing practitioner's manual signature. If the
28 prescription is an electronic prescription, the electronic prescription
29 must contain all of the elements in subdivisions (a) through (j) of this
30 subsection, must be digitally signed, and must be transmitted to and
31 received by the pharmacy electronically to meet all of the requirements

1 of 21 C.F.R. 1311, as the regulation existed on January 1, 2014,
2 pertaining to electronic prescribing of controlled substances.

3 (3) A controlled substance listed in Schedule III, IV, or V of
4 section 28-405 may be dispensed pursuant to a facsimile of a written,
5 signed paper prescription. The facsimile of a written, signed paper
6 prescription shall serve as the original written prescription for
7 purposes of this subsection and shall be maintained in accordance with
8 subsection (2) of section 28-414.03.

9 (4) A prescription for a controlled substance listed in Schedule
10 III, IV, or V of section 28-405 may be partially filled if (a) each
11 partial filling is recorded in the same manner as a refilling, (b) the
12 total quantity dispensed in all partial fillings does not exceed the
13 total quantity prescribed, and (c) each partial filling is dispensed
14 within six months after the prescription was issued.

15 Sec. 3. Section 38-101, Revised Statutes Supplement, 2019, is
16 amended to read:

17 38-101 Sections 38-101 to 38-1,145 and section 4 of this act and the
18 following practice acts shall be known and may be cited as the Uniform
19 Credentialing Act:

- 20 (1) The Advanced Practice Registered Nurse Practice Act;
- 21 (2) The Alcohol and Drug Counseling Practice Act;
- 22 (3) The Athletic Training Practice Act;
- 23 (4) The Audiology and Speech-Language Pathology Practice Act;
- 24 (5) The Certified Nurse Midwifery Practice Act;
- 25 (6) The Certified Registered Nurse Anesthetist Practice Act;
- 26 (7) The Chiropractic Practice Act;
- 27 (8) The Clinical Nurse Specialist Practice Act;
- 28 (9) The Cosmetology, Electrology, Esthetics, Nail Technology, and
29 Body Art Practice Act;
- 30 (10) The Dentistry Practice Act;
- 31 (11) The Dialysis Patient Care Technician Registration Act;

- 1 (12) The Emergency Medical Services Practice Act;
- 2 (13) The Environmental Health Specialists Practice Act;
- 3 (14) The Funeral Directing and Embalming Practice Act;
- 4 (15) The Genetic Counseling Practice Act;
- 5 (16) The Hearing Instrument Specialists Practice Act;
- 6 (17) The Licensed Practical Nurse-Certified Practice Act until
- 7 November 1, 2017;
- 8 (18) The Massage Therapy Practice Act;
- 9 (19) The Medical Nutrition Therapy Practice Act;
- 10 (20) The Medical Radiography Practice Act;
- 11 (21) The Medicine and Surgery Practice Act;
- 12 (22) The Mental Health Practice Act;
- 13 (23) The Nurse Practice Act;
- 14 (24) The Nurse Practitioner Practice Act;
- 15 (25) The Nursing Home Administrator Practice Act;
- 16 (26) The Occupational Therapy Practice Act;
- 17 (27) The Optometry Practice Act;
- 18 (28) The Perfusion Practice Act;
- 19 (29) The Pharmacy Practice Act;
- 20 (30) The Physical Therapy Practice Act;
- 21 (31) The Podiatry Practice Act;
- 22 (32) The Psychology Practice Act;
- 23 (33) The Respiratory Care Practice Act;
- 24 (34) The Surgical First Assistant Practice Act;
- 25 (35) The Veterinary Medicine and Surgery Practice Act; and
- 26 (36) The Water Well Standards and Contractors' Practice Act.

27 If there is any conflict between any provision of sections 38-101 to
28 38-1,145 and section 4 of this act and any provision of a practice act,
29 the provision of the practice act shall prevail.

30 The Revisor of Statutes shall assign the Uniform Credentialing Act,
31 including the practice acts enumerated in subdivisions (1) through (35)

1 of this section, to articles within Chapter 38.

2 Sec. 4. (1) For purposes of this section, prescriber means a health
3 care practitioner authorized to prescribe controlled substances in the
4 practice for which credentialed under the Uniform Credentialing Act.

5 (2) Except as otherwise provided in subsection (3) of this section,
6 no prescriber shall, in this state, issue any prescription as defined in
7 section 38-2840 for a controlled substance as defined in section 28-401
8 unless such prescription is issued (a) using electronic prescription
9 technology, (b) from the prescriber issuing the prescription to a
10 pharmacy, and (c) in accordance with all requirements of state law and
11 the rules and regulations adopted and promulgated pursuant to such state
12 law.

13 (3) The requirements of subsection (2) of this section shall not
14 apply to prescriptions:

15 (a) Issued by veterinarians;

16 (b) Issued in circumstances where electronic prescribing is not
17 available due to temporary technological or electrical failure;

18 (c) Issued by a prescriber to be dispensed by a pharmacy located
19 outside the state as set forth in rules and regulations adopted and
20 promulgated by the Department of Health and Human Services;

21 (d) Issued when the prescriber and the dispenser are the same
22 entity;

23 (e) Issued that include elements that are not supported by the
24 Prescriber/Pharmacist Interface SCRIPT Standard of the National Council
25 for Prescription Drug Programs as such standard existed on January 1,
26 2020;

27 (f) Issued for a drug for which the federal Food and Drug
28 Administration requires the prescription to contain certain elements that
29 are not able to be accomplished with electronic prescribing;

30 (g) Issued for dispensing a non-patient-specific prescription which
31 is (i) a standing order, (ii) approved protocol for drug therapy, (iii)

1 collaborative drug management, (iv) comprehensive medication management,
2 (v) in response to a public health emergency, or (vi) in other
3 circumstances where the prescriber may issue a non-patient-specific
4 prescription;

5 (h) Issued for a drug for purposes of a research protocol;

6 (i) Issued by a prescriber who has received a waiver or a renewal of
7 a waiver for a specified period determined by the chief medical officer
8 of the Department of Health and Human Services, not to exceed one year,
9 from the requirement to use electronic prescribing, pursuant to a process
10 established in rules and regulations adopted and promulgated by the
11 Department of Health and Human Services, in consultation with the chief
12 medical officer, due to economic hardship, technological limitations that
13 are not reasonably within the control of the prescriber, or other
14 exceptional circumstance demonstrated by the prescriber;

15 (j) Issued under circumstances in which, notwithstanding the
16 prescriber's ability to make an electronic prescription as required by
17 this section, such prescriber reasonably determines (i) that it would be
18 impractical for the patient to obtain substances prescribed by electronic
19 prescription in a timely manner and (ii) that such delay would adversely
20 impact the patient's medical condition; or

21 (k) Issued for drugs requiring compounding.

22 (4) A pharmacist who receives a written, oral, or faxed prescription
23 is not required to verify that the prescription falls under one of the
24 exceptions listed in subsection (3) of this section. A pharmacist may
25 continue to dispense medication from any otherwise valid written, oral,
26 or faxed prescription consistent with the law and rules and regulations
27 as they existed prior to January 1, 2021.

28 (5) A violation of this section shall not be grounds for
29 disciplinary action under the Uniform Credentialing Act.

30 Sec. 5. Section 38-2870, Revised Statutes Cumulative Supplement,
31 2018, is amended to read:

1 38-2870 (1) Beginning January 1, 2021, prescriptions for controlled
2 substances listed in section 28-405 shall be subject to section 4 of this
3 act.

4 (2) ~~(1)~~ All medical orders shall be written, oral, or electronic and
5 shall be valid for the period stated in the medical order, except that
6 (a) if the medical order is for a controlled substance listed in section
7 28-405, such period shall not exceed six months from the date of issuance
8 at which time the medical order shall expire and (b) if the medical order
9 is for a drug or device which is not a controlled substance listed in
10 section 28-405 or is an order issued by a practitioner for pharmaceutical
11 care, such period shall not exceed twelve months from the date of
12 issuance at which time the medical order shall expire.

13 (3) ~~(2)~~ Prescription drugs or devices may only be dispensed by a
14 pharmacist or pharmacist intern pursuant to a medical order, by an
15 individual dispensing pursuant to a delegated dispensing permit, or as
16 otherwise provided in section 38-2850. Notwithstanding any other
17 provision of law to the contrary, a pharmacist or a pharmacist intern may
18 dispense drugs or devices pursuant to a medical order or an individual
19 dispensing pursuant to a delegated dispensing permit may dispense drugs
20 or devices pursuant to a medical order. The Pharmacy Practice Act shall
21 not be construed to require any pharmacist or pharmacist intern to
22 dispense, compound, administer, or prepare for administration any drug or
23 device pursuant to any medical order. A pharmacist or pharmacist intern
24 shall retain the professional right to refuse to dispense.

25 (4) ~~(3)~~ Except as otherwise provided in sections 28-414 and
26 28-414.01, a practitioner or the practitioner's agent may transmit a
27 medical order to a pharmacist or pharmacist intern and an authorized
28 refill to a pharmacist, pharmacist intern, or pharmacy technician by the
29 following means: (a) In writing, (b) orally, (c) by facsimile
30 transmission of a written medical order or electronic transmission of a
31 medical order signed by the practitioner, or (d) by facsimile

1 transmission of a written medical order or electronic transmission of a
2 medical order which is not signed by the practitioner. Such an unsigned
3 medical order shall be verified with the practitioner.

4 (5)(a) ~~(4)(a)~~ Except as otherwise provided in sections 28-414 and
5 28-414.01, any medical order transmitted by facsimile or electronic
6 transmission shall:

7 (i) Be transmitted by the practitioner or the practitioner's agent
8 directly to a pharmacist or pharmacist intern in a licensed pharmacy of
9 the patient's choice; and any authorized refill transmitted by facsimile
10 or electronic transmission shall be transmitted by the practitioner or
11 the practitioner's agent directly to a pharmacist, pharmacist intern, or
12 pharmacy technician. No intervening person shall be permitted access to
13 the medical order to alter such order or the licensed pharmacy chosen by
14 the patient. Such medical order may be transmitted through a third-party
15 intermediary who shall facilitate the transmission of the order from the
16 practitioner or practitioner's agent to the pharmacy;

17 (ii) Identify the transmitter's telephone number or other suitable
18 information necessary to contact the transmitter for written or oral
19 confirmation, the time and date of the transmission, the identity of the
20 pharmacy intended to receive the transmission, and other information as
21 required by law; and

22 (iii) Serve as the original medical order if all other requirements
23 of this subsection are satisfied.

24 (b) Medical orders transmitted by electronic transmission shall be
25 signed by the practitioner either with an electronic signature for legend
26 drugs which are not controlled substances or a digital signature for
27 legend drugs which are controlled substances.

28 (6) ~~(5)~~ The pharmacist shall exercise professional judgment
29 regarding the accuracy, validity, and authenticity of any medical order
30 transmitted by facsimile or electronic transmission.

31 (7) ~~(6)~~ The quantity of drug indicated in a medical order for a

1 resident of a long-term care facility shall be sixty days unless
2 otherwise limited by the prescribing practitioner.

3 Sec. 6. Section 38-2891, Revised Statutes Cumulative Supplement,
4 2018, is amended to read:

5 38-2891 (1) A pharmacy technician shall only perform tasks which do
6 not require the professional judgment of a pharmacist and which are
7 subject to verification to assist a pharmacist in the practice of
8 pharmacy.

9 (2) The functions and tasks which shall not be performed by pharmacy
10 technicians include, but are not limited to:

11 (a) Receiving oral medical orders from a practitioner or his or her
12 agent except as otherwise provided in subsection (4) ~~(3)~~ of section
13 38-2870;

14 (b) Providing patient counseling;

15 (c) Performing any evaluation or necessary clarification of a
16 medical order or performing any functions other than strictly clerical
17 functions involving a medical order;

18 (d) Supervising or verifying the tasks and functions of pharmacy
19 technicians;

20 (e) Interpreting or evaluating the data contained in a patient's
21 record maintained pursuant to section 38-2869;

22 (f) Releasing any confidential information maintained by the
23 pharmacy;

24 (g) Performing any professional consultations; and

25 (h) Drug product selection, with regard to an individual medical
26 order, in accordance with the Nebraska Drug Product Selection Act.

27 (3) The director shall, with the recommendation of the board, waive
28 any of the limitations in subsection (2) of this section for purposes of
29 a scientific study of the role of pharmacy technicians approved by the
30 board. Such study shall be based upon providing improved patient care or
31 enhanced pharmaceutical care. Any such waiver shall state the length of

1 the study and shall require that all study data and results be made
2 available to the board upon the completion of the study. Nothing in this
3 subsection requires the board to approve any study proposed under this
4 subsection.

5 Sec. 7. This act becomes operative on January 1, 2021.

6 Sec. 8. Original sections 28-414, 28-414.01, 38-2870, and 38-2891,
7 Revised Statutes Cumulative Supplement, 2018, and section 38-101, Revised
8 Statutes Supplement, 2019, are repealed.