SENATE STATE OF MINNESOTA EIGHTY-EIGHTH SESSION

S.F. No. 1484

(SENATE AUTHORS: SHERAN, Hayden and Nienow)

DATE	D-PG	OFFICIAL STATUS
03/20/2013	1356	Introduction and first reading
		Referred to Health, Human Services and Housing
03/19/2014	6387	Chief author stricken, shown as co-author Hayden
		Chief author added Sheran
03/24/2014	6664a	Comm report: To pass as amended and re-refer to Finance
04/10/2014	8131a	Comm report: To pass as amended
	8199	Second reading
04/29/2014	8508a	Special Order: Amended
	8511	Third reading Passed

relating to health; making changes to dental licensing provisions; authorizing the administration of influenza vaccine by qualified dentists under certain circumstances; providing penalties; modifying grounds for disciplinary action by the Board of Nursing; modifying the health professionals services program; modifying the compensation paid to the health-related licensing board members; making changes to the Minnesota prescription monitoring program; adding and modifying definitions; changing the requirements for pharmacist participation in immunizations; changing the powers and duties of the Board of Pharmacy; changing licensing requirements for businesses regulated by the Board of Pharmacy; clarifying requirements for compounding; allowing certain educational institutions to purchase legend drugs in limited circumstances; allowing certain entities to handle drugs in preparation for emergency use; clarifying the requirement that drug manufacturers report certain payments to the Board of Pharmacy; adding certain substances to the schedules for controlled substances; requiring a report; appropriating money; amending Minnesota Statutes 2012, sections 148.261, subdivisions 1, 4, by adding a subdivision; 150A.01, subdivision 8a; 150A.06, subdivisions 1, 1a, 1c, 1d, 2, 2a, 2d, 3, 8; 150A.091, subdivisions 3, 8, 16; 150A.10; 151.01; 151.06; 151.211; 151.26; 151.34; 151.35; 151.361, subdivision 2; 151.37, as amended; 151.44; 151.58, subdivisions 2, 3, 5; 152.02, subdivision 8b; 152.126, as amended; 214.09, subdivision 3; 214.32, by adding a subdivision; 214.33, subdivision 3; Minnesota

A bill for an act

BE IT ENACTED BY THE LEGISLATURE OF THE STATE OF MINNESOTA:

Statutes 2013 Supplement, sections 151.252, by adding a subdivision; 152.02,

subdivision 2; 364.09; proposing coding for new law in Minnesota Statutes,

1.27 ARTICLE 1

chapters 150A; 151.

1.28 **HEALTH-RELATED LICENSING BOARDS**

1.29 Section 1. Minnesota Statutes 2012, section 148.261, subdivision 1, is amended to read:

Subdivision 1. **Grounds listed.** The board may deny, revoke, suspend, limit, or condition the license and registration of any person to practice professional, advanced practice registered, or practical nursing under sections 148.171 to 148.285, or to otherwise

1.1

1.2

1.3

1.4

1.5

1.6

1.7

1.8

19

1.10

1.11

1.12

1.13

1.14

1.15

1 16

1.17

1 18

1.19

1.20

1.21

1.22

1 23

1.24

1.25

1.26

1 30

1 31

2.2

2.3

2.4

2.5

2.6

2.7

2.8

2.9

2.10

2.11

2.12

2.13

2.14

2.15

2.16

2.17

2.18

2.19

2.20

2.21

2.22

2.23

2.24

2.25

2.26

2.27

2.28

2.29

2.30

2.31

2.32

2.33

2.34

2.35

discipline a licensee or applicant as described in section 148.262. The following are grounds for disciplinary action:

- (1) Failure to demonstrate the qualifications or satisfy the requirements for a license contained in sections 148.171 to 148.285 or rules of the board. In the case of a person applying for a license, the burden of proof is upon the applicant to demonstrate the qualifications or satisfaction of the requirements.
- (2) Employing fraud or deceit in procuring or attempting to procure a permit, license, or registration certificate to practice professional or practical nursing or attempting to subvert the licensing examination process. Conduct that subverts or attempts to subvert the licensing examination process includes, but is not limited to:
- (i) conduct that violates the security of the examination materials, such as removing examination materials from the examination room or having unauthorized possession of any portion of a future, current, or previously administered licensing examination;
- (ii) conduct that violates the standard of test administration, such as communicating with another examinee during administration of the examination, copying another examinee's answers, permitting another examinee to copy one's answers, or possessing unauthorized materials; or
- (iii) impersonating an examinee or permitting an impersonator to take the examination on one's own behalf.
- (3) Conviction of a felony or gross misdemeanor reasonably related to the practice of professional, advanced practice registered, or practical nursing. Conviction as used in this subdivision includes a conviction of an offense that if committed in this state would be considered a felony or gross misdemeanor without regard to its designation elsewhere, or a criminal proceeding where a finding or verdict of guilt is made or returned but the adjudication of guilt is either withheld or not entered.
- (4) Revocation, suspension, limitation, conditioning, or other disciplinary action against the person's professional or practical nursing license or advanced practice registered nursing credential, in another state, territory, or country; failure to report to the board that charges regarding the person's nursing license or other credential are pending in another state, territory, or country; or having been refused a license or other credential by another state, territory, or country.
- (5) Failure to or inability to perform professional or practical nursing as defined in section 148.171, subdivision 14 or 15, with reasonable skill and safety, including failure of a registered nurse to supervise or a licensed practical nurse to monitor adequately the performance of acts by any person working at the nurse's direction.

3.1	
3.2	fi
3.3	ir
3.4	C
3.5	a
3.6	p
3.7	ir

38

3.9

3.10

3.11

3.12

3.13

3.14

3.15

3.16

3.17

3.18

3.19

3.20

3.21

3.22

3.23

3.24

3.25

3.26

3.27

3.28

3.29

3.30

3.31

3.32

3.33

3.34

3.35

3.36

(6) Engaging in unprofessional conduct, including, but not limited to, a departure rom or failure to conform to board rules of professional or practical nursing practice that nterpret the statutory definition of professional or practical nursing as well as provide riteria for violations of the statutes, or, if no rule exists, to the minimal standards of cceptable and prevailing professional or practical nursing practice, or any nursing ractice that may create unnecessary danger to a patient's life, health, or safety. Actual injury to a patient need not be established under this clause.

- (7) Failure of an advanced practice registered nurse to practice with reasonable skill and safety or departure from or failure to conform to standards of acceptable and prevailing advanced practice registered nursing.
- (8) Delegating or accepting the delegation of a nursing function or a prescribed health care function when the delegation or acceptance could reasonably be expected to result in unsafe or ineffective patient care.
- (9) Actual or potential inability to practice nursing with reasonable skill and safety to patients by reason of illness, use of alcohol, drugs, chemicals, or any other material, or as a result of any mental or physical condition.
- (10) Adjudication as mentally incompetent, mentally ill, a chemically dependent person, or a person dangerous to the public by a court of competent jurisdiction, within or without this state.
- (11) Engaging in any unethical conduct, including, but not limited to, conduct likely to deceive, defraud, or harm the public, or demonstrating a willful or careless disregard for the health, welfare, or safety of a patient. Actual injury need not be established under this clause.
- (12) Engaging in conduct with a patient that is sexual or may reasonably be interpreted by the patient as sexual, or in any verbal behavior that is seductive or sexually demeaning to a patient, or engaging in sexual exploitation of a patient or former patient.
- (13) Obtaining money, property, or services from a patient, other than reasonable fees for services provided to the patient, through the use of undue influence, harassment, duress, deception, or fraud.
- (14) Revealing a privileged communication from or relating to a patient except when otherwise required or permitted by law.
- (15) Engaging in abusive or fraudulent billing practices, including violations of federal Medicare and Medicaid laws or state medical assistance laws.
- (16) Improper management of patient records, including failure to maintain adequate patient records, to comply with a patient's request made pursuant to sections 144.291 to 144.298, or to furnish a patient record or report required by law.

4.2

4.3

4.4

4.5

4.6

4.7

48

4.9

4.10

4.11

4.12

4.13

4.14

4.15

4.16

4.17

4.18

4.19

4.20

4.21

4.22

4.23

4.24

4.25

4.26

4.27

4.28

4.29

4.30

4.31

4.32

4.33

4.34

4.35

4.36

(17) Knowingly aiding, assisting, advising, or allowing an unlicensed person to engage in the unlawful practice of professional, advanced practice registered, or practical nursing.

S1484-3

3rd Engrossment

- (18) Violating a rule adopted by the board, an order of the board, or a state or federal law relating to the practice of professional, advanced practice registered, or practical nursing, or a state or federal narcotics or controlled substance law.
- (19) Knowingly providing false or misleading information that is directly related to the care of that patient unless done for an accepted therapeutic purpose such as the administration of a placebo.
- (20) Aiding suicide or aiding attempted suicide in violation of section 609.215 as established by any of the following:
- (i) a copy of the record of criminal conviction or plea of guilty for a felony in violation of section 609.215, subdivision 1 or 2;
- (ii) a copy of the record of a judgment of contempt of court for violating an injunction issued under section 609.215, subdivision 4;
- (iii) a copy of the record of a judgment assessing damages under section 609.215, subdivision 5; or
- (iv) a finding by the board that the person violated section 609.215, subdivision 1 or 2. The board shall investigate any complaint of a violation of section 609.215, subdivision 1 or 2.
- (21) Practicing outside the scope of practice authorized by section 148.171, subdivision 5, 10, 11, 13, 14, 15, or 21.
- (22) Practicing outside the specific field of nursing practice for which an advanced practice registered nurse is certified unless the practice is authorized under section 148.284.
- (23) Making a false statement or knowingly providing false information to the board, failing to make reports as required by section 148.263, or failing to cooperate with an investigation of the board as required by section 148.265.
 - (24) Engaging in false, fraudulent, deceptive, or misleading advertising.
- (25) Failure to inform the board of the person's certification status as a nurse anesthetist, nurse-midwife, nurse practitioner, or clinical nurse specialist.
- (26) Engaging in clinical nurse specialist practice, nurse-midwife practice, nurse practitioner practice, or registered nurse anesthetist practice without current certification by a national nurse certification organization acceptable to the board, except during the period between completion of an advanced practice registered nurse course of study and certification, not to exceed six months or as authorized by the board.
 - (27) Engaging in conduct that is prohibited under section 145.412.

5.1	(28) Failing to report employment to the board as required by section 148.211,
5.2	subdivision 2a, or knowingly aiding, assisting, advising, or allowing a person to fail to
5.3	report as required by section 148.211, subdivision 2a.
5.4	(29) Discharge from the health professionals services program as described in
5.5	sections 214.31 to 214.37, or any other alternative monitoring or diversion program for
5.6	reasons other than satisfactory completion of the program as set forth in the participation
5.7	agreement.
5.8	Sec. 2. Minnesota Statutes 2012, section 148.261, is amended by adding a subdivision
5.9	to read:
5.10	Subd. 1a. Conviction of a felony-level criminal sexual offense. (a) Except as
5.11	provided in paragraph (e), the board may not grant or renew a license to practice nursing
5.12	to any person who has been convicted on or after August 1, 2014, of any of the provisions
5.13	of sections 609.342, subdivision 1, 609.343, subdivision 1, 609.344, subdivision 1,
5.14	paragraphs (c) to (o), or 609.345, subdivision 1, paragraphs (c) to (o), or a similar statute
5.15	in another jurisdiction.
5.16	(b) A license to practice nursing is automatically revoked if the licensee is convicted
5.17	of an offense listed in paragraph (a) of this section.
5.18	(c) A license to practice nursing that has been denied or revoked under this
5.19	subdivision is not subject to chapter 364.
5.20	(d) For purposes of this subdivision, "conviction" means a plea of guilty, a verdict of
5.21	guilty by a jury, or a finding of guilty by the court, unless the court stays imposition or
5.22	execution of the sentence and final disposition of the case is accomplished at a nonfelony
5.23	<u>level.</u>
5.24	(e) The board may establish criteria whereby an individual convicted of an offense
5.25	listed in paragraph (a) of this subdivision may become licensed provided that the criteria:
5.26	(1) utilize a rebuttable presumption that the applicant is not suitable for licensing;
5.27	(2) provide a standard for overcoming the presumption; and
5.28	(3) require that a minimum of ten years has elapsed since the applicant's sentence
5.29	was discharged.
5.30	The board shall not consider an application under this paragraph if the board
5.31	determines that the victim involved in the offense was a patient or a client of the applicant
5.32	at the time of the offense.

Sec. 3. Minnesota Statutes 2012, section 148.261, subdivision 4, is amended to read:

6.1	Subd. 4. Evidence. In disciplinary actions alleging a violation of subdivision 1,
6.2	clause (3) or (4), or subdivision 1a, a copy of the judgment or proceeding under the seal
6.3	of the court administrator or of the administrative agency that entered the same shall be
6.4	admissible into evidence without further authentication and shall constitute prima facie
6.5	evidence of the violation concerned.
6.6	Sec. 4. Minnesota Statutes 2012, section 150A.01, subdivision 8a, is amended to read:
6.7	Subd. 8a. Resident dentist. "Resident dentist" means a person who is licensed to
6.8	practice dentistry as an enrolled graduate student or student of an advanced education
6.9	program accredited by the American Dental Association Commission on Dental
6.10	Accreditation.
6.11	Sec. 5. [150A.055] ADMINISTRATION OF INFLUENZA IMMUNIZATIONS.
6.12	Subdivision 1. Practice of dentistry. A person licensed to practice dentistry under
6.13	sections 150A.01 to 150A.14 shall be deemed to be practicing dentistry while participating
6.14	in the administration of an influenza vaccination.
6.15	Subd. 2. Qualified dentists. (a) The influenza immunization shall be administered
6.16	only to patients 19 years of age and older and only by licensed dentists who:
6.17	(1) have immediate access to emergency response equipment, including but not
6.18	limited to oxygen administration equipment, epinephrine, and other allergic reaction
6.19	response equipment; and
6.20	(2) are trained in or have successfully completed a program approved by the
6.21	Minnesota Board of Dentistry, specifically for the administration of immunizations. The
6.22	training or program must include:
6.23	(i) educational material on the disease of influenza and vaccination as prevention
6.24	of the disease;
6.25	(ii) contraindications and precautions;
6.26	(iii) intramuscular administration;
6.27	(iv) communication of risk and benefits of influenza vaccination and legal
6.28	requirements involved;
6.29	(v) reporting of adverse events;
6.30	(vi) documentation required by federal law; and
6.31	(vii) storage and handling of vaccines.
6.32	(b) Any dentist giving influenza vaccinations under this section shall comply
6.33	with guidelines established by the federal Advisory Committee on Immunization
6.34	Practices relating to vaccines and immunizations, which includes, but is not limited to,

7.2

7.3

7.4

7.5

7.6

7.7

7.8

7.9

7.10

7.11

7 12

7 13

7.14

7.15

7.16

7.17

7.18

7.19

7.20

7.21

7.22

7.23

7.24

7.25

7.26

7.27

7 28

7.29

7.30

7.31

7.32

7.33

7.34

vaccine storage and handling, vaccine administration and documentation, and vaccine contraindications and precautions.

DM

Subd. 3. Coordination of care. After a dentist qualified under subdivision 2 has administered an influenza vaccine to a patient, the dentist shall report the administration of the immunization to the Minnesota Immunization Information Connection or otherwise notify the patient's primary physician or clinic of the administration of the immunization.

EFFECTIVE DATE. This section is effective January 1, 2015, and applies to influenza immunizations performed on or after that date.

Sec. 6. Minnesota Statutes 2012, section 150A.06, subdivision 1, is amended to read: Subdivision 1. **Dentists.** A person of good moral character who has graduated from a dental program accredited by the Commission on Dental Accreditation of the American Dental Association, having submitted an application and fee as prescribed by the board, may be examined by the board or by an agency pursuant to section 150A.03, subdivision 1, in a manner to test the applicant's fitness to practice dentistry. A graduate of a dental college in another country must not be disqualified from examination solely because of the applicant's foreign training if the board determines that the training is equivalent to or higher than that provided by a dental college accredited by the Commission on Dental Accreditation of the American Dental Association. In the case of examinations conducted pursuant to section 150A.03, subdivision 1, applicants shall take the examination prior to applying to the board for licensure. The examination shall include an examination of the applicant's knowledge of the laws of Minnesota relating to dentistry and the rules of the board. An applicant is ineligible to retake the clinical examination required by the board after failing it twice until further education and training are obtained as specified by the board by rule. A separate, nonrefundable fee may be charged for each time a person applies. An applicant who passes the examination in compliance with subdivision 2b, abides by professional ethical conduct requirements, and meets all other requirements of the board shall be licensed to practice dentistry and granted a general dentist license by the board.

Sec. 7. Minnesota Statutes 2012, section 150A.06, subdivision 1a, is amended to read: Subd. 1a. Faculty dentists. (a) Faculty members of a school of dentistry must be licensed in order to practice dentistry as defined in section 150A.05. The board may issue to members of the faculty of a school of dentistry a license designated as either a "limited faculty license" or a "full faculty license" entitling the holder to practice dentistry within the terms described in paragraph (b) or (c). The dean of a school of dentistry and program directors of a Minnesota dental hygiene or dental assisting school accredited by

8.1 th
8.2 to
8.3 lie
8.4 de
8.5 hy
8.6 lie
8.7 a
8.8 re
8.9 fo
8.10 th

8.11

8.12

8.13

8.14

8.15

8.16

8.17

8.18

8.19

8.20

8.21

8.22

8.23

8.24

8.25

8.26

8.27

8.28

8.29

8.30

8.31

8.32

8.33

8.34

the Commission on Dental Accreditation of the American Dental Association shall certify to the board those members of the school's faculty who practice dentistry but are not licensed to practice dentistry in Minnesota. A faculty member who practices dentistry as defined in section 150A.05, before beginning duties in a school of dentistry or a dental hygiene or dental assisting school, shall apply to the board for a limited or full faculty license. Pursuant to Minnesota Rules, chapter 3100, and at the discretion of the board, a limited faculty license must be renewed annually and a full faculty license must be renewed biennially. The faculty applicant shall pay a nonrefundable fee set by the board for issuing and renewing the faculty license. The faculty license is valid during the time the holder remains a member of the faculty of a school of dentistry or a dental hygiene or dental assisting school and subjects the holder to this chapter.

- (b) The board may issue to dentist members of the faculty of a Minnesota school of dentistry, dental hygiene, or dental assisting accredited by the Commission on Dental Accreditation of the American Dental Association, a license designated as a limited faculty license entitling the holder to practice dentistry within the school and its affiliated teaching facilities, but only for the purposes of teaching or conducting research. The practice of dentistry at a school facility for purposes other than teaching or research is not allowed unless the dentist was a faculty member on August 1, 1993.
- (c) The board may issue to dentist members of the faculty of a Minnesota school of dentistry, dental hygiene, or dental assisting accredited by the Commission on Dental Accreditation of the American Dental Association a license designated as a full faculty license entitling the holder to practice dentistry within the school and its affiliated teaching facilities and elsewhere if the holder of the license is employed 50 percent time or more by the school in the practice of teaching or research, and upon successful review by the board of the applicant's qualifications as described in subdivisions 1, 1c, and 4 and board rule. The board, at its discretion, may waive specific licensing prerequisites.
 - Sec. 8. Minnesota Statutes 2012, section 150A.06, subdivision 1c, is amended to read:
- Subd. 1c. **Specialty dentists.** (a) The board may grant <u>a one or more</u> specialty <u>license licenses</u> in the specialty areas of dentistry that are recognized by the <u>American Dental Association</u> Commission on Dental Accreditation.
 - (b) An applicant for a specialty license shall:
- (1) have successfully completed a postdoctoral specialty education program accredited by the Commission on Dental Accreditation of the American Dental Association, or have announced a limitation of practice before 1967;

- (2) have been certified by a specialty examining board approved by the Minnesota Board of Dentistry, or provide evidence of having passed a clinical examination for licensure required for practice in any state or Canadian province, or in the case of oral and maxillofacial surgeons only, have a Minnesota medical license in good standing;
- (3) have been in active practice or a postdoctoral specialty education program or United States government service at least 2,000 hours in the 36 months prior to applying for a specialty license;
- (4) if requested by the board, be interviewed by a committee of the board, which may include the assistance of specialists in the evaluation process, and satisfactorily respond to questions designed to determine the applicant's knowledge of dental subjects and ability to practice;
- (5) if requested by the board, present complete records on a sample of patients treated by the applicant. The sample must be drawn from patients treated by the applicant during the 36 months preceding the date of application. The number of records shall be established by the board. The records shall be reasonably representative of the treatment typically provided by the applicant for each specialty area;
- (6) at board discretion, pass a board-approved English proficiency test if English is not the applicant's primary language;
 - (7) pass all components of the National Board Dental Examinations;
 - (8) pass the Minnesota Board of Dentistry jurisprudence examination;
 - (9) abide by professional ethical conduct requirements; and
- 9.22 (10) meet all other requirements prescribed by the Board of Dentistry.
 - (c) The application must include:

9.2

9.3

9.4

9.5

9.6

9.7

98

9.9

9.10

9.11

9.12

9.13

9.14

9.15

9.16

9.17

9.18

9.19

9.20

9.21

9.23

9.25

9.26

9.27

9.28

9.29

9.30

9.31

9.32

9.33

9.34

9.35

- 9.24 (1) a completed application furnished by the board;
 - (2) at least two character references from two different dentists for each specialty area, one of whom must be a dentist practicing in the same specialty area, and the other from the director of the each specialty program attended;
 - (3) a licensed physician's statement attesting to the applicant's physical and mental condition;
 - (4) a statement from a licensed ophthalmologist or optometrist attesting to the applicant's visual acuity;
 - (5) a nonrefundable fee; and
 - (6) a notarized, unmounted passport-type photograph, three inches by three inches, taken not more than six months before the date of application.
 - (d) A specialty dentist holding a <u>one or more</u> specialty <u>license</u> is limited to practicing in the dentist's designated specialty area or areas. The scope of practice must be

10.2

10.3

10.4

10.5

10.6

10.7

10.8

10.9

10.10

10.11

10.12

10.13

10.14

10.15

10.16

10.17

10.18

10.19

10.20

10.21

10.22

10.23

10.24

10.25

10.26

10.27

10.28

10.29

10.30

10.31

10.32

10.33

10.34

defined by each national specialty board recognized by the American Dental Association Commission on Dental Accreditation.

DM

- (e) A specialty dentist holding a general dentist dental license is limited to practicing in the dentist's designated specialty area or areas if the dentist has announced a limitation of practice. The scope of practice must be defined by each national specialty board recognized by the American Dental Association Commission on Dental Accreditation.
- (f) All specialty dentists who have fulfilled the specialty dentist requirements and who intend to limit their practice to a particular specialty area or areas may apply for a one or more specialty license licenses.

Sec. 9. Minnesota Statutes 2012, section 150A.06, subdivision 1d, is amended to read: Subd. 1d. **Dental therapists.** A person of good moral character who has graduated with a baccalaureate degree or a master's degree from a dental therapy education program that has been approved by the board or accredited by the American Dental Association Commission on Dental Accreditation or another board-approved national accreditation organization may apply for licensure.

The applicant must submit an application and fee as prescribed by the board and a diploma or certificate from a dental therapy education program. Prior to being licensed, the applicant must pass a comprehensive, competency-based clinical examination that is approved by the board and administered independently of an institution providing dental therapy education. The applicant must also pass an examination testing the applicant's knowledge of the Minnesota laws and rules relating to the practice of dentistry. An applicant who has failed the clinical examination twice is ineligible to retake the clinical examination until further education and training are obtained as specified by the board. A separate, nonrefundable fee may be charged for each time a person applies. An applicant who passes the examination in compliance with subdivision 2b, abides by professional ethical conduct requirements, and meets all the other requirements of the board shall be licensed as a dental therapist.

Sec. 10. Minnesota Statutes 2012, section 150A.06, subdivision 2, is amended to read:
Subd. 2. **Dental hygienists.** A person of good moral character, who has graduated
from a dental hygiene program accredited by the Commission on Dental Accreditation of
the American Dental Association and established in an institution accredited by an agency
recognized by the United States Department of Education to offer college-level programs,
may apply for licensure. The dental hygiene program must provide a minimum of two
academic years of dental hygiene education. The applicant must submit an application and

11.2

11.3

11.4

11.5

11.6

11.7

11.8

11.9

11.10

11.11

11.12

11.13

11.14

11.15

11.16

11.17

11.18

11.19

11.20

11.21

11.22

11.23

11.24

11.25

11.26

11.27

11.28

11.29

11.30

11.31

11.32

11.33

11.34

fee as prescribed by the board and a diploma or certificate of dental hygiene. Prior to being licensed, the applicant must pass the National Board of Dental Hygiene examination and a board approved examination designed to determine the applicant's clinical competency. In the case of examinations conducted pursuant to section 150A.03, subdivision 1, applicants shall take the examination before applying to the board for licensure. The applicant must also pass an examination testing the applicant's knowledge of the laws of Minnesota relating to the practice of dentistry and of the rules of the board. An applicant is ineligible to retake the clinical examination required by the board after failing it twice until further education and training are obtained as specified by board rule. A separate, nonrefundable fee may be charged for each time a person applies. An applicant who passes the examination in compliance with subdivision 2b, abides by professional ethical conduct requirements, and meets all the other requirements of the board shall be licensed as a dental hygienist.

DM

Sec. 11. Minnesota Statutes 2012, section 150A.06, subdivision 2a, is amended to read: Subd. 2a. Licensed dental assistant. A person of good moral character, who has graduated from a dental assisting program accredited by the Commission on Dental Accreditation of the American Dental Association, may apply for licensure. The applicant must submit an application and fee as prescribed by the board and the diploma or certificate of dental assisting. In the case of examinations conducted pursuant to section 150A.03, subdivision 1, applicants shall take the examination before applying to the board for licensure. The examination shall include an examination of the applicant's knowledge of the laws of Minnesota relating to dentistry and the rules of the board. An applicant is ineligible to retake the licensure examination required by the board after failing it twice until further education and training are obtained as specified by board rule. A separate, nonrefundable fee may be charged for each time a person applies. An applicant who passes the examination in compliance with subdivision 2b, abides by professional ethical conduct requirements, and meets all the other requirements of the board shall be licensed as a dental assistant.

Sec. 12. Minnesota Statutes 2012, section 150A.06, subdivision 2d, is amended to read: Subd. 2d. Continuing education and professional development waiver. (a) The board shall grant a waiver to the continuing education requirements under this chapter for a licensed dentist, licensed dental therapist, licensed dental hygienist, or licensed dental assistant who documents to the satisfaction of the board that the dentist, dental therapist, dental hygienist, or licensed dental assistant has retired from active practice in the state and limits the provision of dental care services to those offered without compensation

12.2

12.3

12.4

12.5

126

12.7

12.8

12.9

12.10

12.11

12.12

12.13

12.14

12.15

12.16

12.17

12.18

12.19

12.20

12.21

12.22

12.23

12.24

12.25

12.26

12.27

12.28

12.29

12.30

12.31

12.32

12.33

12.34

12.35

in a public health, community, or tribal clinic or a nonprofit organization that provides services to the indigent or to recipients of medical assistance, general assistance medical care, or MinnesotaCare programs.

- (b) The board may require written documentation from the volunteer and retired dentist, dental therapist, dental hygienist, or licensed dental assistant prior to granting this waiver.
- (c) The board shall require the volunteer and retired dentist, dental therapist, dental hygienist, or licensed dental assistant to meet the following requirements:
- (1) a licensee seeking a waiver under this subdivision must complete and document at least five hours of approved courses in infection control, medical emergencies, and medical management for the continuing education cycle; and
- (2) provide documentation of current CPR certification from completion of the American Heart Association healthcare provider course, or the American Red Cross professional rescuer course, or an equivalent entity.
 - Sec. 13. Minnesota Statutes 2012, section 150A.06, subdivision 3, is amended to read:
- Subd. 3. Waiver of examination. (a) All or any part of the examination for dentists or dental hygienists, except that pertaining to the law of Minnesota relating to dentistry and the rules of the board, may, at the discretion of the board, be waived for an applicant who presents a certificate of having passed all components of the National Board Dental Examinations or evidence of having maintained an adequate scholastic standing as determined by the board, in dental school as to dentists, or dental hygiene school as to dental hygienists.
- (b) The board shall waive the clinical examination required for licensure for any dentist applicant who is a graduate of a dental school accredited by the Commission on Dental Accreditation of the American Dental Association, who has passed all components of the National Board Dental Examinations, and who has satisfactorily completed a Minnesota-based postdoctoral general dentistry residency program (GPR) or an advanced education in general dentistry (AEGD) program after January 1, 2004. The postdoctoral program must be accredited by the Commission on Dental Accreditation of the American Dental Association, be of at least one year's duration, and include an outcome assessment evaluation assessing the resident's competence to practice dentistry. The board may require the applicant to submit any information deemed necessary by the board to determine whether the waiver is applicable. The board may waive the clinical examination for an applicant who meets the requirements of this paragraph and has satisfactorily completed an accredited postdoctoral general dentistry residency program located outside of Minnesota.

13.2

13.3

13.4

13.5

136

13.7

13.8

13.9

13.10

13.11

13.12

13.13

13.14

13.15

13.16

13.17

13.18

13.19

13.20

13.21

13.22

13.23

13.24

13.25

13.26

13.27

13.28

13.29

13.30

13.31

13.33

13.34

13.35

	Sec.	14.	Minnesota	Statutes	2012,	section	150A.06,	, subdivision	8,	is amended to	read
--	------	-----	-----------	----------	-------	---------	----------	---------------	----	---------------	------

- Subd. 8. **Licensure by credentials.** (a) Any dental assistant may, upon application and payment of a fee established by the board, apply for licensure based on an evaluation of the applicant's education, experience, and performance record in lieu of completing a board-approved dental assisting program for expanded functions as defined in rule, and may be interviewed by the board to determine if the applicant:
- (1) has graduated from an accredited dental assisting program accredited by the Commission of on Dental Accreditation of the American Dental Association, or is currently certified by the Dental Assisting National Board;
- (2) is not subject to any pending or final disciplinary action in another state or Canadian province, or if not currently certified or registered, previously had a certification or registration in another state or Canadian province in good standing that was not subject to any final or pending disciplinary action at the time of surrender;
- (3) is of good moral character and abides by professional ethical conduct requirements;
- (4) at board discretion, has passed a board-approved English proficiency test if English is not the applicant's primary language; and
- (5) has met all expanded functions curriculum equivalency requirements of a Minnesota board-approved dental assisting program.
- (b) The board, at its discretion, may waive specific licensure requirements in paragraph (a).
- (c) An applicant who fulfills the conditions of this subdivision and demonstrates the minimum knowledge in dental subjects required for licensure under subdivision 2a must be licensed to practice the applicant's profession.
- (d) If the applicant does not demonstrate the minimum knowledge in dental subjects required for licensure under subdivision 2a, the application must be denied. If licensure is denied, the board may notify the applicant of any specific remedy that the applicant could take which, when passed, would qualify the applicant for licensure. A denial does not prohibit the applicant from applying for licensure under subdivision 2a.
- (e) A candidate whose application has been denied may appeal the decision to the board according to subdivision 4a.
- Sec. 15. Minnesota Statutes 2012, section 150A.091, subdivision 3, is amended to read:
 - Subd. 3. **Initial license or permit fees.** Along with the application fee, each of the following applicants shall submit a separate prorated initial license or permit fee. The prorated initial fee shall be established by the board based on the number of months of the

14.1	applicant's initial term as described in Minnesota Rules, part 3100.1700, subpart 1a, not to
14.2	exceed the following monthly nonrefundable fee amounts:
14.3	(1) dentist or full faculty dentist, \$14 times the number of months of the initial
14.4	term_\$168;
14.5	(2) dental therapist, \$10 times the number of months of the initial term \$120;
14.6	(3) dental hygienist, \$5 times the number of months of the initial term \$60;
14.7	(4) licensed dental assistant, \$3 times the number of months of the initial term
14.8	<u>\$36</u> ; and
14.9	(5) dental assistant with a permit as described in Minnesota Rules, part 3100.8500,
14.10	subpart 3, \$1 times the number of months of the initial term \$12.
14.11	Sec. 16. Minnesota Statutes 2012, section 150A.091, subdivision 8, is amended to read:
14.12	Subd. 8. Duplicate license or certificate fee. Each applicant shall submit, with
14.13	a request for issuance of a duplicate of the original license, or of an annual or biennial
14.14	renewal certificate for a license or permit, a fee in the following amounts:
14.15	(1) original dentist, full faculty dentist, dental therapist, dental hygiene, or dental
14.16	assistant license, \$35; and
14.17	(2) annual or biennial renewal certificates, \$10-; and
14.18	(3) wallet-sized license and renewal certificate, \$15.
14.19	Sec. 17. Minnesota Statutes 2012, section 150A.091, subdivision 16, is amended to
14.20	read:
14.21	Subd. 16. Failure of professional development portfolio audit. A licensee shall
14.22	submit a fee as established by the board not to exceed the amount of \$250 after failing two
14.23	consecutive professional development portfolio audits and, thereafter, for each failed (a) If
14.24	<u>a licensee fails a</u> professional development portfolio audit under Minnesota Rules, part
14.25	3100.5300, the board is authorized to take the following actions:
14.26	(1) for the first failure, the board may issue a warning to the licensee;
14.27	(2) for the second failure within ten years, the board may assess a penalty of not
14.28	more than \$250; and
14.29	(3) for any additional failures within the ten year period, the board may assess a
14.30	penalty of not more than \$1000.
14.31	(b) In addition to the penalty fee, the board may initiate the complaint process to
14.32	address multiple failed audits.

Sec. 18. Minnesota Statutes 2012, section 150A.10, is amended to read:

150A.10 ALLIED DENTAL PERSONNEL.

15.1

15.2

15.3

15.4

15.5

15.6

15.7

15.8

15.9

15.10

15.11

15.12

15.13

15.14

15.15

15.16

15.17

15.18

15.19

15.20

15.21

15.22

15.23

15.24

15.25

15.26

15.27

15.28

15.29

Subdivision 1. **Dental hygienists.** Any licensed dentist, licensed dental therapist, public institution, or school authority may obtain services from a licensed dental hygienist. The licensed dental hygienist may provide those services defined in section 150A.05, subdivision 1a. The services provided shall not include the establishment of a final diagnosis or treatment plan for a dental patient. All services shall be provided under supervision of a licensed dentist. Any licensed dentist who shall permit any dental service by a dental hygienist other than those authorized by the Board of Dentistry, shall be deemed to be violating the provisions of sections 150A.01 to 150A.12, and any unauthorized dental service by a dental hygienist shall constitute a violation of sections 150A.01 to 150A.12.

- Subd. 1a. **Limited authorization for dental hygienists.** (a) Notwithstanding subdivision 1, a dental hygienist licensed under this chapter may be employed or retained by a health care facility, program, or nonprofit organization to perform dental hygiene services described under paragraph (b) without the patient first being examined by a licensed dentist if the dental hygienist:
- (1) has been engaged in the active practice of clinical dental hygiene for not less than 2,400 hours in the past 18 months or a career total of 3,000 hours, including a minimum of 200 hours of clinical practice in two of the past three years;
- (2) has entered into a collaborative agreement with a licensed dentist that designates authorization for the services provided by the dental hygienist;
- (3) has documented participation in courses in infection control and medical emergencies within each continuing education cycle; and
- (4) maintains current CPR certification from completion of the American Heart Association healthcare provider course, or the American Red Cross professional rescuer course, or an equivalent entity.
- (b) The dental hygiene services authorized to be performed by a dental hygienist under this subdivision are limited to:
 - (1) oral health promotion and disease prevention education;
- 15.30 (2) removal of deposits and stains from the surfaces of the teeth;
- 15.31 (3) application of topical preventive or prophylactic agents, including fluoride 15.32 varnishes and pit and fissure sealants;
- 15.33 (4) polishing and smoothing restorations;
- 15.34 (5) removal of marginal overhangs;
- 15.35 (6) performance of preliminary charting;
- 15.36 (7) taking of radiographs; and

16.2

16.3

16.4

16.5

16.6

16.7

16.8

16.9

16.10

16.11

16.12

16.13

16.14

16.15

16.16

16.17

16.18

16.19

16.20

16.21

16.22

16.23

16.24

16.25

16.26

16.27

16.28

16.29

16.30

16.31

16.32

16.33

16.34

16.35

16.36

(8) performance of scaling and root planing.

The dental hygienist may administer injections of local anesthetic agents or nitrous oxide inhalation analgesia as specifically delegated in the collaborative agreement with a licensed dentist. The dentist need not first examine the patient or be present. If the patient is considered medically compromised, the collaborative dentist shall review the patient record, including the medical history, prior to the provision of these services. Collaborating dental hygienists may work with unlicensed and licensed dental assistants who may only perform duties for which licensure is not required. The performance of dental hygiene services in a health care facility, program, or nonprofit organization as authorized under this subdivision is limited to patients, students, and residents of the facility, program, or organization.

- (c) A collaborating dentist must be licensed under this chapter and may enter into a collaborative agreement with no more than four dental hygienists unless otherwise authorized by the board. The board shall develop parameters and a process for obtaining authorization to collaborate with more than four dental hygienists. The collaborative agreement must include:
- (1) consideration for medically compromised patients and medical conditions for which a dental evaluation and treatment plan must occur prior to the provision of dental hygiene services;
- (2) age- and procedure-specific standard collaborative practice protocols, including recommended intervals for the performance of dental hygiene services and a period of time in which an examination by a dentist should occur;
 - (3) copies of consent to treatment form provided to the patient by the dental hygienist;
- (4) specific protocols for the placement of pit and fissure sealants and requirements for follow-up care to assure the efficacy of the sealants after application; and
- (5) a procedure for creating and maintaining dental records for the patients that are treated by the dental hygienist. This procedure must specify where these records are to be located.
- The collaborative agreement must be signed and maintained by the dentist, the dental hygienist, and the facility, program, or organization; must be reviewed annually by the collaborating dentist and dental hygienist; and must be made available to the board upon request.
- (d) Before performing any services authorized under this subdivision, a dental hygienist must provide the patient with a consent to treatment form which must include a statement advising the patient that the dental hygiene services provided are not a substitute for a dental examination by a licensed dentist. If the dental hygienist makes any referrals

17.2

17.3

17.4

17.5

17.6

17.7

17.8

17.9

17.10

17.11

17.12

17.13

17.14

17.15

17.16

17.17

17.18

17.19

17.20

17.21

17.22

17.23

17.24

17.25

17.26

17.27

17.28

17.29

17.30

17.31

17.32

17.33

17.34

17.35

17.36

to the patient for further dental procedures, the dental hygienist must fill out a referral form and provide a copy of the form to the collaborating dentist.

- (e) For the purposes of this subdivision, a "health care facility, program, or nonprofit organization" is limited to a hospital; nursing home; home health agency; group home serving the elderly, disabled, or juveniles; state-operated facility licensed by the commissioner of human services or the commissioner of corrections; and federal, state, or local public health facility, community clinic, tribal clinic, school authority, Head Start program, or nonprofit organization that serves individuals who are uninsured or who are Minnesota health care public program recipients.
- (f) For purposes of this subdivision, a "collaborative agreement" means a written agreement with a licensed dentist who authorizes and accepts responsibility for the services performed by the dental hygienist. The services authorized under this subdivision and the collaborative agreement may be performed without the presence of a licensed dentist and may be performed at a location other than the usual place of practice of the dentist or dental hygienist and without a dentist's diagnosis and treatment plan, unless specified in the collaborative agreement.
- Subd. 2. **Dental assistants.** Every licensed dentist and dental therapist who uses the services of any unlicensed person for the purpose of assistance in the practice of dentistry or dental therapy shall be responsible for the acts of such unlicensed person while engaged in such assistance. The dentist or dental therapist shall permit the unlicensed assistant to perform only those acts which are authorized to be delegated to unlicensed assistants by the Board of Dentistry. The acts shall be performed under supervision of a licensed dentist or dental therapist. A licensed dental therapist shall not supervise more than four registered licensed or unlicensed dental assistants at any one practice setting. The board may permit differing levels of dental assistance based upon recognized educational standards, approved by the board, for the training of dental assistants. The board may also define by rule the scope of practice of licensed and unlicensed dental assistants. The board by rule may require continuing education for differing levels of dental assistants, as a condition to their license or authority to perform their authorized duties. Any licensed dentist or dental therapist who permits an unlicensed assistant to perform any dental service other than that authorized by the board shall be deemed to be enabling an unlicensed person to practice dentistry, and commission of such an act by an unlicensed assistant shall constitute a violation of sections 150A.01 to 150A.12.
- Subd. 3. **Dental technicians.** Every licensed dentist and dental therapist who uses the services of any unlicensed person, other than under the dentist's or dental therapist's supervision and within the same practice setting, for the purpose of constructing, altering,

18.2

18.3

18.4

18.5

18.6

18.7

18.8

18.9

18.10

18.11

18.12

18.13

18.14

18.15

18.16

18.17

18 18

18.19

18.20

18.21

18.22

18.23

18.24

18.25

18.26

18.27

18.28

18.29

18.30

18.31

18.32

18.33

repairing or duplicating any denture, partial denture, crown, bridge, splint, orthodontic,
prosthetic or other dental appliance, shall be required to furnish such unlicensed person
with a written work order in such form as shall be prescribed by the rules of the board. The
work order shall be made in duplicate form, a duplicate copy to be retained in a permanent
file of the dentist or dental therapist at the practice setting for a period of two years, and
the original to be retained in a permanent file for a period of two years by the unlicensed
person in that person's place of business. The permanent file of work orders to be kept
by the dentist, dental therapist, or unlicensed person shall be open to inspection at any
reasonable time by the board or its duly constituted agent.

- Subd. 4. **Restorative procedures.** (a) Notwithstanding subdivisions 1, 1a, and 2, a licensed dental hygienist or licensed dental assistant may perform the following restorative procedures:
 - (1) place, contour, and adjust amalgam restorations;
 - (2) place, contour, and adjust glass ionomer;
 - (3) adapt and cement stainless steel crowns; and
- (4) place, contour, and adjust class I and class V supragingival composite restorations where the margins are entirely within the enamel-; and
- (5) place, contour, and adjust class II and class V supragingival composite restorations on primary teeth.
 - (b) The restorative procedures described in paragraph (a) may be performed only if:
- (1) the licensed dental hygienist or licensed dental assistant has completed a board-approved course on the specific procedures;
- (2) the board-approved course includes a component that sufficiently prepares the licensed dental hygienist or licensed dental assistant to adjust the occlusion on the newly placed restoration;
- (3) a licensed dentist or licensed advanced dental therapist has authorized the procedure to be performed; and
- (4) a licensed dentist or licensed advanced dental therapist is available in the clinic while the procedure is being performed.
- (c) The dental faculty who teaches the educators of the board-approved courses specified in paragraph (b) must have prior experience teaching these procedures in an accredited dental education program.
 - Sec. 19. Minnesota Statutes 2012, section 214.09, subdivision 3, is amended to read:
- Subd. 3. **Compensation.** (a) Members of the boards may be compensated at the rate of \$55 a day spent on board activities, when authorized by the board, plus expenses

19.2

19.3

19.4

19.5

19.6

19.7

19.8

19.9

19.10

19.11

19.12

19.13

19.14

19.15

19.16

19.17

19.18

19.19

19.20

19.21

19.22

19.23

19.24

19.25

19.26

19.27

19.28

19.29

19.30

19.31

19.32

19.33

in Members of health-related licensing boards may be compensated at the rate of \$75 a day spent on board activities and members of nonhealth-related licensing boards may be compensated at the rate of \$55 a day spent on board activities when authorized by the board, plus expenses in the same manner and amount as authorized by the commissioner's plan adopted under section 43A.18, subdivision 2. Members who, as a result of time spent attending board meetings, incur child care expenses that would not otherwise have been incurred, may be reimbursed for those expenses upon board authorization.

- (b) Members who are state employees or employees of the political subdivisions of the state must not receive the daily payment for activities that occur during working hours for which they are also compensated by the state or political subdivision. However, a state or political subdivision employee may receive the daily payment if the employee uses vacation time or compensatory time accumulated in accordance with a collective bargaining agreement or compensation plan for board activity. Members who are state employees or employees of the political subdivisions of the state may receive the expenses provided for in this subdivision unless the expenses are reimbursed by another source. Members who are state employees or employees of political subdivisions of the state may be reimbursed for child care expenses only for time spent on board activities that are outside their working hours.
- (c) Each board must adopt internal standards prescribing what constitutes a day spent on board activities for purposes of making daily payments under this subdivision.
- Sec. 20. Minnesota Statutes 2012, section 214.32, is amended by adding a subdivision to read:
- Subd. 6. **Duties of a participating board.** Upon receiving a report from the program manager in accordance with section 214.33, subdivision 3, that a regulated person has been discharged from the program due to noncompliance based on allegations that the regulated person has engaged in conduct that might cause risk to the public, the participating board may temporarily suspend the regulated person's professional license until the completion of a disciplinary investigation. The board must complete the disciplinary investigation within 60 days of receipt of the report from the program. If the investigation is not completed by the board within 60 days, the temporary suspension shall be lifted, unless the regulated person requests a delay in the disciplinary proceedings for any reason, upon which the temporary suspension shall remain in place until the completion of the investigation.
 - Sec. 21. Minnesota Statutes 2012, section 214.33, subdivision 3, is amended to read:

20.1	Subd. 3. Program manager. (a) The program manager shall report to the
20.2	appropriate participating board a regulated person who:
20.3	(1) does not meet program admission criteria-;
20.4	(2) violates the terms of the program participation agreement, or;
20.5	(3) leaves or is discharged from the program except upon fulfilling the terms for
20.6	successful completion of the program as set forth in the participation agreement-;
20.7	(4) is subject to the provisions of sections 214.17 to 214.25;
20.8	(5) causes identifiable patient harm;
20.9	(6) unlawfully substitutes or adulterates medications;
20.10	(7) writes a prescription or causes a prescription to be dispensed in the name of a
20.11	person, other than the prescriber, or veterinary patient for the personal use of the prescriber;
20.12	(8) alters a prescription without the knowledge of the prescriber for the purpose of
20.13	obtaining a drug for personal use;
20.14	(9) unlawfully uses a controlled or mood-altering substance or uses alcohol while
20.15	providing patient care or during the period of time in which the regulated person may be
20.16	contacted to provide patient care or is otherwise on duty, if current use is the reason for
20.17	participation in the program or the use occurs while the regulated person is participating
20.18	in the program; or
20.19	The program manager shall report to the appropriate participating board a regulated
20.20	person who (10) is alleged to have committed violations of the person's practice act that
20.21	are outside the authority of the health professionals services program as described in
20.22	sections 214.31 to 214.37.
20.23	(b) The program manager shall inform any reporting person of the disposition of the
20.24	person's report to the program.
20.25	EFFECTIVE DATE. This section is effective August 1, 2014, and applies to
20.26	violations that occur after the effective date.
20.27	Sec. 22. Minnesota Statutes 2013 Supplement, section 364.09, is amended to read:
20.28	364.09 EXCEPTIONS.
20.29	(a) This chapter does not apply to the licensing process for peace officers; to law
20.30	enforcement agencies as defined in section 626.84, subdivision 1, paragraph (f); to fire
20.31	protection agencies; to eligibility for a private detective or protective agent license; to the
20.32	licensing and background study process under chapters 245A and 245C; to eligibility
20.33	for school bus driver endorsements; to eligibility for special transportation service
20.34	endorsements; to eligibility for a commercial driver training instructor license, which is

SF1484	REVISOR	DM	S1484-3	3rd Engrossment
DITIOI	TEE TISOIT	2111	511015	ora Engrosoment

governed by section 171.35 and rules adopted under that section; to emergency medical
services personnel, or to the licensing by political subdivisions of taxicab drivers, if the
applicant for the license has been discharged from sentence for a conviction within the ter
years immediately preceding application of a violation of any of the following:

- (1) sections 609.185 to 609.21, 609.221 to 609.223, 609.342 to 609.3451, or 617.23, subdivision 2 or 3;
- (2) any provision of chapter 152 that is punishable by a maximum sentence of 15 years or more; or
- (3) a violation of chapter 169 or 169A involving driving under the influence, leaving the scene of an accident, or reckless or careless driving. 21.10
- 21.11 This chapter also shall not apply to eligibility for juvenile corrections employment, where the offense involved child physical or sexual abuse or criminal sexual conduct. 21.12
 - (b) This chapter does not apply to a school district or to eligibility for a license issued or renewed by the Board of Teaching or the commissioner of education.
 - (c) Nothing in this section precludes the Minnesota Police and Peace Officers Training Board or the state fire marshal from recommending policies set forth in this chapter to the attorney general for adoption in the attorney general's discretion to apply to law enforcement or fire protection agencies.
 - (d) This chapter does not apply to a license to practice medicine that has been denied or revoked by the Board of Medical Practice pursuant to section 147.091, subdivision 1a.
 - (e) This chapter does not apply to any person who has been denied a license to practice chiropractic or whose license to practice chiropractic has been revoked by the board in accordance with section 148.10, subdivision 7.
 - (f) This chapter does not apply to any license, registration, or permit that has been denied or revoked by the Board of Nursing in accordance with section 148.261, subdivision 1a.
- (f) (g) This chapter does not supersede a requirement under law to conduct a 21.27 criminal history background investigation or consider criminal history records in hiring 21.28 for particular types of employment. 21.29

21.30 21.31 21.32		_	APPROPRIAT vailable for th Ending June	e Year
21.32			2014	2015
21.34	Sec. 23. APPROPRIATIONS	<u>\$</u>	<u>\$</u>	
21.35	Board of Behavioral Health and Therapy		<u>-0-</u>	8,000

21.2

21.3

21.4

21.5

21.6

21.7

21.8

21.9

21.13

21.14

21.15

21.16

21.17

21.18

21.19

21.20

21.21

21.22

21.23

21.24

21.25

	SF 1484	REVISOR	DM	51484-3		3rd Engrossment		
22.1	This appropriation is from the state							
22.2	government special revenue fund for board							
22.3	member per diem payments and licensing							
22.4	activity.							
22.5	Board of Cl	niropractic Examin	ers		<u>-0-</u>	10,000		
22.6	This appropr	riation is from the s	state					
22.7	government	special revenue fund	d for board					
22.8	member per	diem payments.						
22.9	Board of Do	entistry			<u>-0-</u>	39,000		
22.10	This appropr	riation is from the s	<u>state</u>					
22.11	government	special revenue fund	d for board					
22.12	member per	diem payments.						
22.13	Board of Di	etetics and Nutrition	on Practice		<u>-0-</u>	1,000		
22.14	This appropr	riation is from the s	<u>state</u>					
22.15	government	special revenue fund	d for board					
22.16	member per	diem payments.						
22.17	Board of M	arriage and Family	Therapy		<u>-0-</u>	4,000		
22.18	This appropr	riation is from the s	state					
22.19	government	special revenue fund	d for board					
22.20	member per	diem payments and	licensing					
22.21	activity.							
22.22	Board of M	edical Practice			<u>-0-</u>	38,000		
22.23	This appropr	riation is from the s	state					
22.24	government	special revenue fund	d for board					
22.25	member per	diem payments.						
22.26	Board of Nu	ursing			<u>-0-</u>	266,000		
22.27	This appropr	riation is from the s	state					
22.28	government	special revenue fund	d for board					
22.29	member per	diem payments and	licensing					
22.30	activity.							
22.31	Board of Nu	ursing Home Admi	<u>nistrators</u>		<u>-0-</u>	2,000		

SF1484

REVISOR

DM

S1484-3

3rd Engrossment

	SI 1404 KEVISOK	DW	31404-3		31d Eligiossilicht
23.1	This appropriation is from the	state			
23.2	government special revenue fur	nd for board			
23.3	member per diem payments.				
23.4	Board of Optometry			<u>-0-</u>	<u>1,000</u>
23.5	This appropriation is from the	state			
23.6	government special revenue fur	nd for board			
23.7	member per diem payments.				
23.8	Board of Pharmacy			<u>-0-</u>	2,000
23.9	This appropriation is from the	state			
23.10	government special revenue fur	nd for board			
23.11	member per diem payments.				
23.12	Board of Physical Therapy			<u>-0-</u>	<u>4,000</u>
23.13	This appropriation is from the	state			
23.14	government special revenue fur	nd for board			
23.15	member per diem payments.				
23.16	Board of Podiatric Medicine			<u>-0-</u>	<u>1,000</u>
23.17	This appropriation is from the	<u>state</u>			
23.18	government special revenue fur	nd for board			
23.19	member per diem payments.				
23.20	Board of Psychology			<u>-0-</u>	15,000
23.21	This appropriation is from the	state			
23.22	government special revenue fur	nd for board			
23.23	member per diem payments.				
23.24	Board of Social Work			<u>-0-</u>	17,000
23.25	This appropriation is from the	<u>state</u>			
23.26	government special revenue fur	nd for board			
23.27	member per diem payments and	d licensing			
23.28	activity.				
23.29	Board of Veterinary Medicine	2		<u>-0-</u>	<u>2,000</u>
23.30	This appropriation is from the	state			
23.31	government special revenue fur	nd for board			
23.32	member per diem payments.				

SF1484

REVISOR

DM

S1484-3

3rd Engrossment

24.1 ARTICLE 2

24.2 **BOARD OF PHARMACY**

Section 1. Minnesota Statutes 2012, section 151.01, is amended to read:

151.01 DEFINITIONS.

24.3

244

24.5

24.6

24.7

24.8

24.9

24.10

24.11

24.12

24.13

24.14

24.15

24.16

24.17

24.18

24.19

24.20

24.21

24.22

24.23

24.24

24.25

24.26

24.27

24.28

24.29

24.30

24.31

24.32

24.33

24.34

24.35

Subdivision 1. **Words, terms, and phrases.** Unless the language or context clearly indicates that a different meaning is intended, the following words, terms, and phrases, for the purposes of this chapter, shall be given the meanings subjoined to them.

- Subd. 2. **Pharmacy.** "Pharmacy" means an established a place of business in which prescriptions, prescription drugs, medicines, chemicals, and poisons are prepared, compounded, or dispensed, vended, or sold to or for the use of patients by or under the supervision of a pharmacist and from which related clinical pharmacy services are delivered.
- Subd. 2a. **Limited service pharmacy.** "Limited service pharmacy" means a pharmacy that has been issued a restricted license by the board to perform a limited range of the activities that constitute the practice of pharmacy.
- Subd. 3. **Pharmacist.** The term "pharmacist" means an individual with a currently valid license issued by the Board of Pharmacy to practice pharmacy.
- Subd. 5. **Drug.** The term "drug" means all medicinal substances and preparations recognized by the United States Pharmacopoeia and National Formulary, or any revision thereof, <u>vaccines and biologicals</u>, and all substances and preparations intended for external and internal use in the diagnosis, cure, mitigation, treatment, or prevention of disease in humans or other animals, and all substances and preparations, other than food, intended to affect the structure or any function of the bodies of humans or other animals. The term drug shall also mean any compound, substance, or derivative that is not approved for human consumption by the United States Food and Drug Administration or specifically permitted for human consumption under Minnesota law and, when introduced into the body, induces an effect similar to that of a Schedule I or Schedule II controlled substance listed in section 152.02, subdivisions 2 and 3, or Minnesota Rules, parts 6800.4210 and 6800.4220, regardless of whether the substance is marketed for the purpose of human consumption.
- Subd. 6. **Medicine.** The term "medicine" means any remedial agent that has the property of curing, preventing, treating, or mitigating diseases, or that is used for that purpose.
- Subd. 7. **Poisons.** The term "poisons" means any substance which that, when introduced into the system, directly or by absorption, produces violent, morbid, or fatal changes, or which that destroys living tissue with which it comes in contact.

25.1	Subd. 8. Chemical. The term "chemical" means all medicinal or industrial
25.2	substances, whether simple or compound, or obtained through the process of the science
25.3	and art of chemistry, whether of organic or inorganic origin.
25.4	Subd. 9. Board or State Board of Pharmacy. The term "board" or "State Board of
25.5	Pharmacy" means the Minnesota State Board of Pharmacy.
25.6	Subd. 10. Director. The term "director" means the <u>executive</u> director of the
25.7	Minnesota State Board of Pharmacy.
25.8	Subd. 11. Person. The term "person" means an individual, firm, partnership,
25.9	company, corporation, trustee, association, agency, or other public or private entity.
25.10	Subd. 12. Wholesale. The term "wholesale" means and includes any sale for the
25.11	purpose of resale.
25.12	Subd. 13. Commercial purposes. The phrase "commercial purposes" means the
25.13	ordinary purposes of trade, agriculture, industry, and commerce, exclusive of the practices
25.14	of medicine and, pharmacy, and other health care professions.
25.15	Subd. 14. Manufacturing. The term "manufacturing" except in the case of bulk
25.16	compounding, prepackaging or extemporaneous compounding within a pharmacy, means
25.17	and includes the production, quality control and standardization by mechanical, physical,
25.18	chemical, or pharmaceutical means, packing, repacking, tableting, encapsulating, labeling,
25.19	relabeling, filling or by any other process, of all drugs, medicines, chemicals, or poisons,
25.20	without exception, for medicinal purposes. preparation, propagation, conversion, or
25.21	processing of a drug, either directly or indirectly, by extraction from substances of natural
25.22	origin or independently by means of chemical or biological synthesis. Manufacturing
25.23	includes the packaging or repackaging of a drug, or the labeling or relabeling of
25.24	the container of a drug, for resale by pharmacies, practitioners, or other persons.
25.25	Manufacturing does not include the prepackaging, extemporaneous compounding, or
25.26	anticipatory compounding of a drug within a licensed pharmacy or by a practitioner,
25.27	nor the labeling of a container within a pharmacy or by a practitioner for the purpose of
25.28	dispensing a drug to a patient pursuant to a valid prescription.
25.29	Subd. 14a. Manufacturer. The term "manufacturer" means any person engaged
25.30	in manufacturing.
25.31	Subd. 14b. Outsourcing facility. "Outsourcing facility" means a facility that is
25.32	registered by the United States Food and Drug Administration pursuant to United States
25.33	Code, title 21, section 353b.
25.34	Subd. 15. Pharmacist intern. The term "pharmacist intern" means (1) a natural
25.35	person satisfactorily progressing toward the degree in pharmacy required for licensure, or
25.36	(2) a graduate of the University of Minnesota College of Pharmacy, or other pharmacy

college approved by the board, who is registered by the State Board of Pharmacy for the purpose of obtaining practical experience as a requirement for licensure as a pharmacist, or (3) a qualified applicant awaiting examination for licensure.

Subd. 15a. **Pharmacy technician.** The term "pharmacy technician" means a person not licensed as a pharmacist or a pharmacist intern, who assists the pharmacist in the preparation and dispensing of medications by performing computer entry of prescription data and other manipulative tasks. A pharmacy technician shall not perform tasks specifically reserved to a licensed pharmacist or requiring professional judgment.

Subd. 16. **Prescription** drug order. The term "prescription drug order" means a signed lawful written order, or an, oral, or electronic order reduced to writing, given by of a practitioner licensed to prescribe drugs for patients in the course of the practitioner's practice, issued for an individual patient and containing the following: the date of issue, name and address of the patient, name and quantity of the drug prescribed, directions for use, and the name and address of the prescriber. for a drug for a specific patient.

Prescription drug orders for controlled substances must be prepared in accordance with the provisions of section 152.11 and the federal Controlled Substances Act and the regulations promulgated thereunder.

Subd. 16a. Prescription. The term "prescription" means a prescription drug order that is written or printed on paper, an oral order reduced to writing by a pharmacist, or an electronic order. To be valid, a prescription must be issued for an individual patient by a practitioner within the scope and usual course of the practitioner's practice, and must contain the date of issue, name and address of the patient, name and quantity of the drug prescribed, directions for use, the name and address of the practitioner, and a telephone number at which the practitioner can be reached. A prescription written or printed on paper that is given to the patient or an agent of the patient or that is transmitted by fax must contain the practitioner's manual signature. An electronic prescription must contain the practitioner's electronic signature.

Subd. 16b. Chart order. The term "chart order" means a prescription drug order for a drug that is to be dispensed by a pharmacist, or by a pharmacist intern under the direct supervision of a pharmacist, and administered by an authorized person only during the patient's stay in a hospital or long-term care facility. The chart order shall contain the name of the patient, another patient identifier such as birth date or medical record number, the drug ordered, and any directions that the practitioner may prescribe concerning strength, dosage, frequency, and route of administration. The manual or electronic signature of the practitioner must be affixed to the chart order at the time it is written or at a later date in the case of verbal chart orders.

26.1

26.2

26.3

26.4

26.5

26.6

26.7

26.8

26.9

26.10

26.11

26.12

26.13

26.14

26.15

26.16

26.17

26.18

26.19

26.20

26.21

26.22

26.23

26.24

26.25

26.26

26.27

26.28

26.29

26.30

26.31

26.32

26.33

26.34

26.35

27.2

27.3

27.4

27.5

27.6

27.7

27.8

27.9

27.10

27.11

27.12

27.13

27.14

27.15

27.16

27.17

27.18

27.19

27.20

27.21

27.22

27.23

27.24

27.25

27.26

27.27

27.28

27.29

27.30

27.31

27.32

27.33

27.34

27.35

Subd. 17. Legend drug. "Legend drug" means a drug which that is required by federal law to bear the following statement, "Caution: Federal law prohibits dispensing	
federal law to bear the following statement, "Caution: Federal law prohibits dispensing	
without prescription." be dispensed only pursuant to the prescription of a licensed	
practitioner.	
Subd. 18. Label. "Label" means a display of written, printed, or graphic matter	
upon the immediate container of any drug or medicine; and a requirement made by or	

- upon the immediate container of any drug or medicine; and a requirement made by or under authority of Laws 1969, chapter 933 that. Any word, statement, or other information appearing required by or under the authority of this chapter to appear on the label shall not be considered to be complied with unless such word, statement, or other information also appears appear on the outside container or wrapper, if any there be, of the retail package of such drug or medicine, or is be easily legible through the outside container or wrapper.
- Subd. 19. **Package.** "Package" means any container or wrapping in which any drug or medicine is enclosed for use in the delivery or display of that article to retail purchasers, but does not include:
- (a) shipping containers or wrappings used solely for the transportation of any such article in bulk or in quantity to manufacturers, packers, processors, or wholesale or retail distributors;
- (b) shipping containers or outer wrappings used by retailers to ship or deliver any such article to retail customers if such containers and wrappings bear no printed matter pertaining to any particular drug or medicine.
- Subd. 20. **Labeling.** "Labeling" means all labels and other written, printed, or graphic matter (a) upon a drug or medicine or any of its containers or wrappers, or (b) accompanying such article.
- Subd. 21. **Federal act.** "Federal act" means the Federal Food, Drug, and Cosmetic Act, United States Code, title 21, section 301, et seq., as amended.
- Subd. 22. **Pharmacist in charge.** "Pharmacist in charge" means a duly licensed pharmacist in the state of Minnesota who has been designated in accordance with the rules of the State Board of Pharmacy to assume professional responsibility for the operation of the pharmacy in compliance with the requirements and duties as established by the board in its rules.
- Subd. 23. **Practitioner.** "Practitioner" means a licensed doctor of medicine, licensed doctor of osteopathy duly licensed to practice medicine, licensed doctor of dentistry, licensed doctor of optometry, licensed podiatrist, or licensed veterinarian. For purposes of sections 151.15, subdivision 4; 151.252, subdivision 3; 151.37, subdivision 2, paragraphs (b), (e), and (f); and 151.461, "practitioner" also means a physician assistant authorized to prescribe, dispense, and administer under chapter 147A, or an advanced practice nurse

SF1484 3rd Engrossment authorized to prescribe, dispense, and administer under section 148.235. For purposes of 28.1 sections 151.15, subdivision 4; 151.252, subdivision 3; 151.37, subdivision 2, paragraph 28.2 (b); and 151.461, "practitioner" also means a dental therapist authorized to dispense and 28.3 administer under chapter 150A. 28.4 Subd. 24. **Brand name.** "Brand name" means the registered trademark name given 28.5 to a drug product by its manufacturer, labeler or distributor. 28.6 Subd. 25. Generic name. "Generic name" means the established name or official 28.7 name of a drug or drug product. 28.8 Subd. 26. Finished dosage form. "Finished dosage form" means that form of a 28.9 drug which that is or is intended to be dispensed or administered to the patient and requires 28.10 no further manufacturing or processing other than packaging, reconstitution, or labeling. 28.11 Subd. 27. **Practice of pharmacy.** "Practice of pharmacy" means: 28.12 (1) interpretation and evaluation of prescription drug orders; 28.13

- (2) compounding, labeling, and dispensing drugs and devices (except labeling by a manufacturer or packager of nonprescription drugs or commercially packaged legend drugs and devices);
- (3) participation in clinical interpretations and monitoring of drug therapy for assurance of safe and effective use of drugs, including the performance of laboratory tests that are waived under the federal Clinical Laboratory Improvement Act of 1988, United States Code, title 42, section 263a et seq., provided that a pharmacist may interpret the results of laboratory tests but may modify drug therapy only pursuant to a protocol or collaborative practice agreement;
- (4) participation in drug and therapeutic device selection; drug administration for first dosage and medical emergencies; drug regimen reviews; and drug or drug-related research;
- (5) participation in administration of influenza vaccines to all eligible individuals ten years of age and older and all other vaccines to patients 18 years of age and older under standing orders from a physician licensed under chapter 147 or by written protocol with a physician licensed under chapter 147, a physician assistant authorized to prescribe drugs under chapter 147A, or an advanced practice nurse authorized to prescribe drugs under section 148.235, provided that:
 - (i) the protocol includes, at a minimum:
- (A) the name, dose, and route of each vaccine that may be given; 28.32
- (B) the patient population for whom the vaccine may be given; 28.33
- (C) contraindications and precautions to the vaccine; 28.34
- (D) the procedure for handling an adverse reaction; 28.35

28.14

28.15

28.16

28.17

28.18

28.19

28.20

28.21

28.22

28.23

28.24

28.25

28.26

28.27

28.28

28.29

28.30

29.1	(E) the name, signature, and address of the physician, physician assistant, or
29.2	advanced nurse practitioner;
29.3	(F) a telephone number at which the physician, physician assistant, or advanced
29.4	nurse practitioner can be contacted; and
29.5	(G) the date and time period for which the protocol is valid;
29.6	(i) (ii) the pharmacist is trained in has successfully completed a program approved
29.7	by the American Accreditation Council of Pharmaceutical for Pharmacy Education
29.8	specifically for the administration of immunizations or graduated from a college of
29.9	pharmacy in 2001 or thereafter a program approved by the board; and
29.10	(ii) (iii) the pharmacist reports the administration of the immunization to the patient's
29.11	primary physician or clinic or to the Minnesota Immunization Information Connection; and
29.12	(iv) the pharmacist complies with guidelines for vaccines and immunizations
29.13	established by the federal Advisory Committee on Immunization Practices, except that a
29.14	pharmacist does not need to comply with those portions of the guidelines that establish
29.15	immunization schedules when administering a vaccine pursuant to a valid, patient-specific
29.16	order issued by a physician licensed under chapter 147, a physician assistant authorized to
29.17	prescribe drugs under chapter 147A, or an advanced practice nurse authorized to prescribe
29.18	drugs under section 148.235, provided that the order is consistent with the United States
29.19	Food and Drug Administration approved labeling of the vaccine;
29.20	(6) participation in the practice of managing drug therapy and modifying initiation,
29.21	management, modification, and discontinuation of drug therapy, according to section
29.22	151.21, subdivision 1, according to a written protocol or collaborative practice agreement
29.23	between the specific pharmacist: (i) one or more pharmacists and the individual dentist,
29.24	optometrist, physician, podiatrist, or veterinarian who is responsible for the patient's
29.25	care and authorized to independently prescribe drugs one or more dentists, optometrists,
29.26	physicians, podiatrists, or veterinarians; or (ii) one or more pharmacists and one or more
29.27	physician assistants authorized to prescribe, dispense, and administer under chapter 147A,
29.28	or advanced practice nurses authorized to prescribe, dispense, and administer under
29.29	section 148.235. Any significant changes in drug therapy made pursuant to a protocol or
29.30	<u>collaborative practice agreement</u> must be <u>reported</u> <u>documented</u> by the pharmacist <u>to in</u>
29.31	the patient's medical record or reported by the pharmacist to a practitioner responsible
29.32	for the patient's care;
29.33	(7) participation in the storage of drugs and the maintenance of records;
29.34	(8) responsibility for participation in patient counseling on therapeutic values,
29.35	content, hazards, and uses of drugs and devices; and

DM

30.1	(9) offering or performing those acts, services, operations, or transactions necessary
30.2	in the conduct, operation, management, and control of a pharmacy.
30.3	Subd. 27a. Protocol. "Protocol" means:
30.4	(1) a specific written plan that describes the nature and scope of activities that a
30.5	pharmacist may engage in when initiating, managing, modifying, or discontinuing drug
30.6	therapy as allowed in subdivision 27, clause (6); or
30.7	(2) a specific written plan that authorizes a pharmacist to administer vaccines and
30.8	that complies with subdivision 27, clause (5).
30.9	Subd. 27b. Collaborative practice. "Collaborative practice" means patient care
30.10	activities, consistent with subdivision 27, engaged in by one or more pharmacists who
30.11	have agreed to work in collaboration with one or more practitioners to initiate, manage,
30.12	and modify drug therapy under specified conditions mutually agreed to by the pharmacists
30.13	and practitioners.
30.14	Subd. 27c. Collaborative practice agreement. "Collaborative practice agreement"
30.15	means a written and signed agreement between one or more pharmacists and one or more
30.16	practitioners that allows the pharmacist or pharmacists to engage in collaborative practice.
30.17	Subd. 28. Veterinary legend drug. "Veterinary legend drug" means a drug that is
30.18	required by federal law to bear the following statement: "Caution: Federal law restricts
30.19	this drug to use by or on the order of a licensed veterinarian." be dispensed only pursuant
30.20	to the prescription of a licensed veterinarian.
30.21	Subd. 29. Legend medical gas. "Legend medical gas" means a liquid or gaseous
30.22	substance used for medical purposes and that is required by federal law to bear the
30.23	following statement: "Caution: Federal law prohibits dispensing without a prescription."
30.24	be dispensed only pursuant to the prescription of a licensed practitioner.
30.25	Subd. 30. Dispense or dispensing. "Dispense or dispensing" means the preparation
30.26	or delivery of a drug pursuant to a lawful order of a practitioner in a suitable container
30.27	appropriately labeled for subsequent administration to or use by a patient or other individual
30.28	entitled to receive the drug. interpretation, evaluation, and processing of a prescription
30.29	drug order and includes those processes specified by the board in rule that are necessary
30.30	for the preparation and provision of a drug to a patient or patient's agent in a suitable
30.31	container appropriately labeled for subsequent administration to, or use by, a patient.
30.32	Subd. 31. Central service pharmacy. "Central service pharmacy" means a
30.33	pharmacy that may provide dispensing functions, drug utilization review, packaging,
30.34	labeling, or delivery of a prescription product to another pharmacy for the purpose of

filling a prescription.

31.2

31.3

31.4

31.5

31.6

31.7

31.8

31.9

31.10

31.11

31.12

31.13

31.14

31.15

31.16

31.17

31.18

31.19

31.20

31.21

31.22

31.23

31.24

31.25

31.26

31.27

31.28

31.29

31.30

31.31

31.32

31.33

31.34

31.35

Subd. 32. **Electronic signature.** "Electronic signature" means an electronic sound, symbol, or process attached to or associated with a record and executed or adopted by a person with the intent to sign the record.

Subd. 33. **Electronic transmission.** "Electronic transmission" means transmission of information in electronic form.

Subd. 34. **Health professional shortage area.** "Health professional shortage area" means an area designated as such by the federal Secretary of Health and Human Services, as provided under Code of Federal Regulations, title 42, part 5, and United States Code, title 42, section 254E.

Subd. 35. **Compounding.** "Compounding" means preparing, mixing, assembling, packaging, and labeling a drug for an identified individual patient as a result of a practitioner's prescription drug order. Compounding also includes anticipatory compounding, as defined in this section, and the preparation of drugs in which all bulk drug substances and components are nonprescription substances. Compounding does not include mixing or reconstituting a drug according to the product's labeling or to the manufacturer's directions. Compounding does not include the preparation of a drug for the purpose of, or incident to, research, teaching, or chemical analysis, provided that the drug is not prepared for dispensing or administration to patients. All compounding, regardless of the type of product, must be done pursuant to a prescription drug order unless otherwise permitted in this chapter or by the rules of the board. Compounding does not include a minor deviation from such directions with regard to radioactivity, volume, or stability, which is made by or under the supervision of a licensed nuclear pharmacist or a physician, and which is necessary in order to accommodate circumstances not contemplated in the manufacturer's instructions, such as the rate of radioactive decay or geographical distance from the patient.

Subd. 36. Anticipatory compounding. "Anticipatory compounding" means the preparation by a pharmacy of a supply of a compounded drug product that is sufficient to meet the short-term anticipated need of the pharmacy for the filling of prescription drug orders. In the case of practitioners only, anticipatory compounding means the preparation of a supply of a compounded drug product that is sufficient to meet the practitioner's short-term anticipated need for dispensing or administering the drug to patients treated by the practitioner. Anticipatory compounding is not the preparation of a compounded drug product for wholesale distribution.

Subd. 37. Extemporaneous compounding. "Extemporaneous compounding" means the compounding of a drug product pursuant to a prescription drug order for a specific

patient that is issued in advance of the compounding. Extemporaneous compounding is 32.1 not the preparation of a compounded drug product for wholesale distribution. 32.2 Subd. 38. Compounded positron emission tomography drug. "Compounded 32.3 positron emission tomography drug" means a drug that: 32.4 (1) exhibits spontaneous disintegration of unstable nuclei by the emission of 32.5 positrons and is used for the purpose of providing dual photon positron emission 32.6 tomographic diagnostic images; 32.7 (2) has been compounded by or on the order of a practitioner in accordance with the 32.8 relevant parts of Minnesota Rules, chapters 4731 and 6800, for a patient or for research, 32.9 teaching, or quality control; and 32.10 (3) includes any nonradioactive reagent, reagent kit, ingredient, nuclide generator, 32.11 accelerator, target material, electronic synthesizer, or other apparatus or computer program 32.12 to be used in the preparation of such a drug. 32.13 32.14 Sec. 2. Minnesota Statutes 2012, section 151.06, is amended to read: 151.06 POWERS AND DUTIES. 32.15 Subdivision 1. **Generally; rules.** (a) Powers and duties. The Board of Pharmacy 32.16 shall have the power and it shall be its duty: 32.17 32.18 (1) to regulate the practice of pharmacy; (2) to regulate the manufacture, wholesale, and retail sale of drugs within this state; 32.19 (3) to regulate the identity, labeling, purity, and quality of all drugs and medicines 32.20 dispensed in this state, using the United States Pharmacopeia and the National Formulary, 32.21 or any revisions thereof, or standards adopted under the federal act as the standard; 32.22 (4) to enter and inspect by its authorized representative any and all places where 32.23 drugs, medicines, medical gases, or veterinary drugs or devices are sold, vended, given 32.24 away, compounded, dispensed, manufactured, wholesaled, or held; it may secure samples 32.25 or specimens of any drugs, medicines, medical gases, or veterinary drugs or devices 32.26 after paying or offering to pay for such sample; it shall be entitled to inspect and make 32.27 copies of any and all records of shipment, purchase, manufacture, quality control, and 32.28 sale of these items provided, however, that such inspection shall not extend to financial 32.29 data, sales data, or pricing data; 32.30 (5) to examine and license as pharmacists all applicants whom it shall deem qualified 32.31 to be such; 32.32 (6) to license wholesale drug distributors; 32.33 (7) to deny, suspend, revoke, or refuse to renew take disciplinary action against any 32.34

32.35

registration or license required under this chapter, to any applicant or registrant or licensee

upon any of the following grounds: listed in section 151.071, and in accordance with
the provisions of section 151.071;
(i) fraud or deception in connection with the securing of such license or registration;
(ii) in the case of a pharmacist, conviction in any court of a felony;
(iii) in the case of a pharmacist, conviction in any court of an offense involving
moral turpitude;
(iv) habitual indulgence in the use of narcotics, stimulants, or depressant drugs;
or habitual indulgence in intoxicating liquors in a manner which could cause conduct
endangering public health;
(v) unprofessional conduct or conduct endangering public health;
(vi) gross immorality;
(vii) employing, assisting, or enabling in any manner an unlicensed person to
practice pharmacy;
(viii) conviction of theft of drugs, or the unauthorized use, possession, or sale thereof;
(ix) violation of any of the provisions of this chapter or any of the rules of the State
Board of Pharmacy;
(x) in the case of a pharmacy license, operation of such pharmacy without a
pharmacist present and on duty;
(xi) in the case of a pharmacist, physical or mental disability which could cause
incompetency in the practice of pharmacy;
(xii) in the case of a pharmacist, the suspension or revocation of a license to practice
pharmacy in another state; or
(xiii) in the ease of a pharmacist, aiding suicide or aiding attempted suicide in
violation of section 609.215 as established by any of the following:
(A) a copy of the record of criminal conviction or plea of guilty for a felony in
violation of section 609.215, subdivision 1 or 2;
(B) a copy of the record of a judgment of contempt of court for violating an
injunction issued under section 609.215, subdivision 4;
(C) a copy of the record of a judgment assessing damages under section 609.215,
subdivision 5; or
(D) a finding by the board that the person violated section 609.215, subdivision
1 or 2. The board shall investigate any complaint of a violation of section 609.215,
subdivision 1 or 2;
(8) to employ necessary assistants and adopt rules for the conduct of its business;
(9) to register as pharmacy technicians all applicants who the board determines are
qualified to carry out the duties of a pharmacy technician; and

34.2

34.3

34.4

34.5

34.6

34.7

34.8

34.9

34.10

34.11

34.12

34.13

34.14

34.15

34.16

34.17

34.18

34.19

34.20

34.21

34.22

34.23

34.24

34.25

34.26

34.27

34.28

34.29

34.30

34.31

34.32

34.33

34.34

34.35

34.36

(10) to perform such other duties and exercise such other powers as the provisions of
the act may require-; and
(11) to enter and inspect any business to which it issues a license or registration.

DM

(b) Temporary suspension. In addition to any other remedy provided by law, the board may, without a hearing, temporarily suspend a license for not more than 60 days if the board finds that a pharmacist has violated a statute or rule that the board is empowered to enforce and continued practice by the pharmacist would create an imminent risk of harm to others. The suspension shall take effect upon written notice to the pharmacist, specifying the statute or rule violated. At the time it issues the suspension notice, the board shall schedule a disciplinary hearing to be held under the Administrative Procedure Act. The pharmacist shall be provided with at least 20 days' notice of any hearing held under this subdivision.

(e) (b) Rules. For the purposes aforesaid, it shall be the duty of the board to make and publish uniform rules not inconsistent herewith for carrying out and enforcing the provisions of this chapter. The board shall adopt rules regarding prospective drug utilization review and patient counseling by pharmacists. A pharmacist in the exercise of the pharmacist's professional judgment, upon the presentation of a new prescription by a patient or the patient's caregiver or agent, shall perform the prospective drug utilization review required by rules issued under this subdivision.

(d) (c) Substitution; rules. If the United States Food and Drug Administration (FDA) determines that the substitution of drugs used for the treatment of epilepsy or seizures poses a health risk to patients, the board shall adopt rules in accordance with accompanying FDA interchangeability standards regarding the use of substitution for these drugs. If the board adopts a rule regarding the substitution of drugs used for the treatment of epilepsy or seizures that conflicts with the substitution requirements of section 151.21, subdivision 3, the rule shall supersede the conflicting statute. If the rule proposed by the board would increase state costs for state public health care programs, the board shall report to the chairs and ranking minority members of the senate Health and Human Services Budget Division and the house of representatives Health Care and Human Services Finance Division the proposed rule and the increased cost associated with the proposed rule before the board may adopt the rule.

Subd. 1a. Disciplinary action Cease and desist orders. It shall be grounds for disciplinary action by the Board of Pharmacy against the registration of the pharmacy if the Board of Pharmacy determines that any person with supervisory responsibilities at the pharmacy sets policies that prevent a licensed pharmacist from providing drug utilization review and patient counseling as required by rules adopted under subdivision 1. The Board of Pharmacy shall follow the requirements of chapter 14 in any disciplinary actions

35.2

35.3

35.4

35.5

35.6

35.7

35.8

35.9

35.10

35.11

35.12

35.13

35.14

35.15

35.16

35.17

35.18

35.19

35.20

35.21

35.22

35.23

35.24

35.25

35.26

35.27

35.28

35.29

35.30

35.31

35.32

35.33

35.34

35.35

an act or practice constituting a violation of a law, rule, or other order related to the duties and responsibilities entrusted to the board, the board may issue and cause to be served upon the person an order requiring the person to cease and desist from violations.

- (b) The cease and desist order must state the reasons for the issuance of the order and must give reasonable notice of the rights of the person to request a hearing before an administrative law judge. A hearing must be held not later than ten days after the request for the hearing is received by the board. After the completion of the hearing, the administrative law judge shall issue a report within ten days. Within 15 days after receiving the report of the administrative law judge, the board shall issue a further order vacating or making permanent the cease and desist order. The time periods provided in this provision may be waived by agreement of the executive director of the board and the person against whom the cease and desist order was issued. If the person to whom a cease and desist order is issued fails to appear at the hearing after being duly notified, the person is in default, and the proceeding may be determined against that person upon consideration of the cease and desist order, the allegations of which may be considered to be true. Unless otherwise provided, all hearings must be conducted according to chapter 14. The board may adopt rules of procedure concerning all proceedings conducted under this subdivision.
- (c) If no hearing is requested within 30 days of service of the order, the cease and desist order will become permanent.
- (d) A cease and desist order issued under this subdivision remains in effect until it is modified or vacated by the board. The administrative proceeding provided by this subdivision, and subsequent appellate judicial review of that administrative proceeding, constitutes the exclusive remedy for determining whether the board properly issued the cease and desist order and whether the cease and desist order should be vacated or made permanent.
- Subd. 1b. Enforcement of violations of cease and desist orders. (a) Whenever the board under subdivision 1a seeks to enforce compliance with a cease and desist order are considered conclusively established for purposes of proceeding under subdivision 1a for permanent or temporary relief to enforce the cease and desist order. Whenever the board under subdivision 1a seeks to enforce compliance with a cease and desist order when a hearing or hearing request on the cease and desist order is pending, or the time has not yet expired to request a hearing on whether a cease and desist order should be vacated or made permanent, the allegations in the cease and desist order are considered conclusively

36.2

36.3

36.4

36.5

36.6

36.7

36.8

36.9

36.10

36.11

36.12

36.13

36.14

36.15

36.16

36.17

36.18

36.19

36.20

36.21

36.22

36.23

36.24

36.25

36.26

36.27

36.28

36.29

36.30

36.31

36.32

36.33

established for the purposes of proceeding under subdivision 1a for temporary relief to enforce the cease and desist order.

- (b) Notwithstanding this subdivision or subdivision 1a, the person against whom the cease and desist order is issued and who has requested a hearing under subdivision 1a may, within 15 days after service of the cease and desist order, bring an action in Ramsey County District Court for issuance of an injunction to suspend enforcement of the cease and desist order pending a final decision of the board under subdivision 1a to vacate or make permanent the cease and desist order. The court shall determine whether to issue such an injunction based on traditional principles of temporary relief.
- Subd. 2. **Application.** In the case of a facility licensed or registered by the board, the provisions of subdivision 1 shall apply to an individual owner or sole proprietor and shall also apply to the following:
 - (1) In the case of a partnership, each partner thereof;
 - (2) In the case of an association, each member thereof;
- (3) In the case of a corporation, each officer or director thereof and each shareholder owning 30 percent or more of the voting stock of such corporation.
- Subd. 3. Application of Administrative Procedure Act. The board shall comply with the provisions of chapter 14, before it fails to issue, renew, suspends, or revokes any license or registration issued under this chapter.
- Subd. 4. **Reinstatement.** Any license or registration which has been suspended or revoked may be reinstated by the board provided the holder thereof shall pay all costs of the proceedings resulting in the suspension or revocation, and, in addition thereto, pay a fee set by the board.
- Subd. 5. Costs; penalties. The board may impose a civil penalty not exceeding \$10,000 for each separate violation, the amount of the civil penalty to be fixed so as to deprive a licensee or registrant of any economic advantage gained by reason of the violation, to discourage similar violations by the licensee or registrant or any other licensee or registrant, or to reimburse the board for the cost of the investigation and proceeding, including, but not limited to, fees paid for services provided by the Office of Administrative Hearings, legal and investigative services provided by the Office of the Attorney General, court reporters, witnesses, reproduction of records, board members' per diem compensation, board staff time, and travel costs and expenses incurred by board staff and board members.
- 36.34 **EFFECTIVE DATE.** Subdivisions 1a and 1b are effective August 1, 2014, and apply to violations occurring on or after that date.

	Subdivision 1. Forms of disciplinary action. When the board finds that a licensee,
r	registrant, or applicant has engaged in conduct prohibited under subdivision 2, it may
	lo one or more of the following:
	(1) deny the issuance of a license or registration;
	(2) refuse to renew a license or registration;
	(3) revoke the license or registration;
	(4) suspend the license or registration;
	(5) impose limitations, conditions, or both on the license or registration, including
	out not limited to: the limitation of practice designated settings; the imposition of
	etraining or rehabilitation requirements; the requirement of practice under supervision;
	he requirement of participation in a diversion program such as that established pursuant to
	ection 214.31 or the conditioning of continued practice on demonstration of knowledge
	or skills by appropriate examination or other review of skill and competence;
	(6) impose a civil penalty not exceeding \$10,000 for each separate violation, the
	amount of the civil penalty to be fixed so as to deprive a licensee or registrant of any
	economic advantage gained by reason of the violation, to discourage similar violations
	by the licensee or registrant or any other licensee or registrant, or to reimburse the board
1	for the cost of the investigation and proceeding, including but not limited to, fees paid
	for services provided by the Office of Administrative Hearings, legal and investigative
	ervices provided by the Office of the Attorney General, court reporters, witnesses,
	reproduction of records, board members' per diem compensation, board staff time, and
	ravel costs and expenses incurred by board staff and board members; and
	(7) reprimand the licensee or registrant.
	Subd. 2. Grounds for disciplinary action. The following conduct is prohibited and
l	s grounds for disciplinary action:
	(1) failure to demonstrate the qualifications or satisfy the requirements for a license
	or registration contained in this chapter or the rules of the board. The burden of proof is on
_	he applicant to demonstrate such qualifications or satisfaction of such requirements;
	(2) obtaining a license by fraud or by misleading the board in any way during
t	he application process or obtaining a license by cheating, or attempting to subvert
t	he licensing examination process. Conduct that subverts or attempts to subvert the
l	icensing examination process includes, but is not limited to: (i) conduct that violates the
3	security of the examination materials, such as removing examination materials from the

37.36

examination room or having unauthorized possession of any portion of a future, current,

or previously administered licensing examination; (ii) conduct that violates the standard of

38.1 38.2 38.3 38.4 38.5 38.6 38.7 38.8 38.9 38.10 38.11 38.12 38.13 38.14 38.15 38.16 38.17 38.18 38.19 38.20 38.21 38.22 38.23 38.24 38.25 38.26 38.27 38.28 38.29

38.30

38.31

38.32

38.33

38.34

38.35

38.36

test administration, such as communicating with another examinee during administration of the examination, copying another examinee's answers, permitting another examinee to copy one's answers, or possessing unauthorized materials; or (iii) impersonating an examinee or permitting an impersonator to take the examination on one's own behalf;

- (3) for a pharmacist, pharmacy technician, pharmacist intern, applicant for a pharmacist or pharmacy license, or applicant for a pharmacy technician or pharmacist intern registration, conviction of a felony reasonably related to the practice of pharmacy. Conviction as used in this subdivision includes a conviction of an offense that if committed in this state would be deemed a felony without regard to its designation elsewhere, or a criminal proceeding where a finding or verdict of guilt is made or returned but the adjudication of guilt is either withheld or not entered thereon. The board may delay the issuance of a new license or registration if the applicant has been charged with a felony until the matter has been adjudicated;
- (4) for a facility, other than a pharmacy, licensed or registered by the board, if an owner or applicant is convicted of a felony reasonably related to the operation of the facility. The board may delay the issuance of a new license or registration if the owner or applicant has been charged with a felony until the matter has been adjudicated;
- (5) for a controlled substance researcher, conviction of a felony reasonably related to controlled substances or to the practice of the researcher's profession. The board may delay the issuance of a registration if the applicant has been charged with a felony until the matter has been adjudicated;
- (6) disciplinary action taken by another state or by one of this state's health licensing agencies:
- (i) revocation, suspension, restriction, limitation, or other disciplinary action against a license or registration in another state or jurisdiction, failure to report to the board that charges or allegations regarding the person's license or registration have been brought in another state or jurisdiction, or having been refused a license or registration by any other state or jurisdiction. The board may delay the issuance of a new license or registration if an investigation or disciplinary action is pending in another state or jurisdiction until the investigation or action has been dismissed or otherwise resolved; and
- (ii) revocation, suspension, restriction, limitation, or other disciplinary action against a license or registration issued by another of this state's health licensing agencies, failure to report to the board that charges regarding the person's license or registration have been brought by another of this state's health licensing agencies, or having been refused a license or registration by another of this state's health licensing agencies. The board may delay the issuance of a new license or registration if a disciplinary action is pending before

39.1	another of this state's health licensing agencies until the action has been dismissed or
39.2	otherwise resolved;
39.3	(7) for a pharmacist, pharmacy, pharmacy technician, or pharmacist intern, violation
39.4	of any order of the board, of any of the provisions of this chapter or any rules of the
39.5	board or violation of any federal, state, or local law or rule reasonably pertaining to the
39.6	practice of pharmacy;
39.7	(8) for a facility, other than a pharmacy, licensed by the board, violations of any
39.8	order of the board, of any of the provisions of this chapter or the rules of the board or
39.9	violation of any federal, state, or local law relating to the operation of the facility;
39.10	(9) engaging in any unethical conduct; conduct likely to deceive, defraud, or harm
39.11	the public, or demonstrating a willful or careless disregard for the health, welfare, or safety
39.12	of a patient; or pharmacy practice that is professionally incompetent, in that it may create
39.13	unnecessary danger to any patient's life, health, or safety, in any of which cases, proof
39.14	of actual injury need not be established;
39.15	(10) aiding or abetting an unlicensed person in the practice of pharmacy, except
39.16	that it is not a violation of this clause for a pharmacist to supervise a properly registered
39.17	pharmacy technician or pharmacist intern if that person is performing duties allowed
39.18	by this chapter or the rules of the board;
39.19	(11) for an individual licensed or registered by the board, adjudication as mentally ill
39.20	or developmentally disabled, or as a chemically dependent person, a person dangerous
39.21	to the public, a sexually dangerous person, or a person who has a sexual psychopathic
39.22	personality, by a court of competent jurisdiction, within or without this state. Such
39.23	adjudication shall automatically suspend a license for the duration thereof unless the
39.24	board orders otherwise;
39.25	(12) for a pharmacist or pharmacy intern, engaging in unprofessional conduct as
39.26	specified in the board's rules. In the case of a pharmacy technician, engaging in conduct
39.27	specified in board rules that would be unprofessional if it were engaged in by a pharmacist
39.28	or pharmacist intern or performing duties specifically reserved for pharmacists under this
39.29	chapter or the rules of the board;
39.30	(13) for a pharmacy, operation of the pharmacy without a pharmacist present and on
39.31	duty except as allowed by a variance approved by the board;
39.32	(14) for a pharmacist, the inability to practice pharmacy with reasonable skill and
39.33	safety to patients by reason of illness, drunkenness, use of drugs, narcotics, chemicals, or
39.34	any other type of material or as a result of any mental or physical condition, including
39.35	deterioration through the aging process or loss of motor skills. In the case of registered

39.36

pharmacy technicians, pharmacist interns, or controlled substance researchers, the

40.1	inability to carry out duties allowed under this chapter or the rules of the board with
40.2	reasonable skill and safety to patients by reason of illness, drunkenness, use of drugs,
40.3	narcotics, chemicals, or any other type of material or as a result of any mental or physical
40.4	condition, including deterioration through the aging process or loss of motor skills;
40.5	(15) for a pharmacist, pharmacy, pharmacist intern, pharmacy technician, medical
40.6	gas distributor, or controlled substance researcher, revealing a privileged communication
40.7	from or relating to a patient except when otherwise required or permitted by law;
40.8	(16) for a pharmacist or pharmacy, improper management of patient records,
40.9	including failure to maintain adequate patient records, to comply with a patient's request
40.10	made pursuant to sections 144.291 to 144.298, or to furnish a patient record or report
40.11	required by law;
40.12	(17) paying, offering to pay, receiving, or agreeing to receive, a commission, rebate,
40.13	kickback, or other form of remuneration, directly or indirectly, for the referral of patients
40.14	or the dispensing of drugs or devices;
40.15	(18) engaging in abusive or fraudulent billing practices, including violations of the
40.16	federal Medicare and Medicaid laws or state medical assistance laws or rules;
40.17	(19) engaging in conduct with a patient that is sexual or may reasonably be
40.18	interpreted by the patient as sexual, or in any verbal behavior that is seductive or sexually
40.19	demeaning to a patient;
40.20	(20) failure to make reports as required by section 151.072 or to cooperate with an
40.21	investigation of the board as required by section 151.074;
40.22	(21) knowingly providing false or misleading information that is directly related
40.23	to the care of a patient unless done for an accepted therapeutic purpose such as the
40.24	dispensing and administration of a placebo;
40.25	(22) aiding suicide or aiding attempted suicide in violation of section 609.215 as
40.26	established by any of the following:
40.27	(i) a copy of the record of criminal conviction or plea of guilty for a felony in
40.28	violation of section 609.215, subdivision 1 or 2;
40.29	(ii) a copy of the record of a judgment of contempt of court for violating an
40.30	injunction issued under section 609.215, subdivision 4;
40.31	(iii) a copy of the record of a judgment assessing damages under section 609.215,
40.32	subdivision 5; or
40.33	(iv) a finding by the board that the person violated section 609.215, subdivision
40.34	1 or 2. The board shall investigate any complaint of a violation of section 609.215,
40.35	subdivision 1 or 2;

41.2

41.3

41.4

41.5

41.6

41.7

41.8

41.9

41.10

41.11

41.12

41.13

41.14

41.15

41.16

41.17

41.18

41.19

41.20

41.21

41.22

41.23

41.24

41.25

41.26

41.27

41.28

41.29

41.30

41.31

41.32

41.33

41.34

41.35

41.36

(23) for a pharmacist, practice of pharmacy under a lapsed or nonrenewed license. For a pharmacist intern, pharmacy technician, or controlled substance researcher, performing duties permitted to such individuals by this chapter or the rules of the board under a lapsed or nonrenewed registration. For a facility required to be licensed under this chapter, operation of the facility under a lapsed or nonrenewed license or registration; and (24) for a pharmacist, pharmacist intern, or pharmacy technician, termination or discharge from the health professional services program for reasons other than the satisfactory completion of the program.

- Subd. 3. Automatic suspension. (a) A license or registration issued under this chapter to a pharmacist, pharmacist intern, pharmacy technician, or controlled substance researcher is automatically suspended if: (1) a guardian of a licensee or registrant is appointed by order of a court pursuant to sections 524.5-101 to 524.5-502, for reasons other than the minority of the licensee or registrant; or (2) the licensee or registrant is committed by order of a court pursuant to chapter 253B. The license or registration remains suspended until the licensee is restored to capacity by a court and, upon petition by the licensee or registrant, the suspension is terminated by the board after a hearing.
- (b) For a pharmacist, pharmacy intern, or pharmacy technician, upon notice to the board of a judgment of, or a plea of guilty to, a felony reasonably related to the practice of pharmacy, the license or registration of the regulated person may be automatically suspended by the board. The license or registration will remain suspended until, upon petition by the regulated individual and after a hearing, the suspension is terminated by the board. The board may indefinitely suspend or revoke the license or registration of the regulated individual if, after a hearing before the board, the board finds that the felonious conduct would cause a serious risk of harm to the public.
- (c) For a facility that is licensed or registered by the board, upon notice to the board that an owner of the facility is subject to a judgment of, or a plea of guilty to, a felony reasonably related to the operation of the facility, the license or registration of the facility may be automatically suspended by the board. The license or registration will remain suspended until, upon petition by the facility and after a hearing, the suspension is terminated by the board. The board may indefinitely suspend or revoke the license or registration of the facility if, after a hearing before the board, the board finds that the felonious conduct would cause a serious risk of harm to the public.
- (d) For licenses and registrations that have been suspended or revoked pursuant to paragraphs (a) and (b), the regulated individual may have a license or registration reinstated, either with or without restrictions, by demonstrating clear and convincing evidence of rehabilitation, as provided in section 364.03. If the regulated individual has

42.2

42.3

42.4

42.5

42.6

42.7

42.8

42.9

42.10

42.11

42.12

42.13

42.14

42.15

42.16

42.17

42.18

42.19

42.20

42.21

42.22

42.23

42.24

42.25

42.26

42.27

42.28

42.29

42.30

42.31

42.32

42.33

42.34

42.35

42.36

the conviction subsequently overturned by court decision, the board shall conduct a hearing to review the suspension within 30 days after the receipt of the court decision. The regulated individual is not required to prove rehabilitation if the subsequent court decision overturns previous court findings of public risk.

- (e) For licenses and registrations that have been suspended or revoked pursuant to paragraph (c), the regulated facility may have a license or registration reinstated, either with or without restrictions, conditions, or limitations, by demonstrating clear and convincing evidence of rehabilitation of the convicted owner, as provided in section 364.03. If the convicted owner has the conviction subsequently overturned by court decision, the board shall conduct a hearing to review the suspension within 30 days after receipt of the court decision. The regulated facility is not required to prove rehabilitation of the convicted owner if the subsequent court decision overturns previous court findings of public risk.
- (f) The board may, upon majority vote of a quorum of its appointed members, suspend the license or registration of a regulated individual without a hearing if the regulated individual fails to maintain a current name and address with the board, as described in paragraphs (h) and (i), while the regulated individual is: (1) under board investigation, and a notice of conference has been issued by the board; (2) party to a contested case with the board; (3) party to an agreement for corrective action with the board; or (4) under a board order for disciplinary action. The suspension shall remain in effect until lifted by the board to the board's receipt of a petition from the regulated individual, along with the current name and address of the regulated individual.
- (g) The board may, upon majority vote of a quorum of its appointed members, suspend the license or registration of a regulated facility without a hearing if the regulated facility fails to maintain a current name and address of the owner of the facility with the board, as described in paragraphs (h) and (i), while the regulated facility is: (1) under board investigation, and a notice of conference has been issued by the board; (2) party to a contested case with the board; (3) party to an agreement for corrective action with the board; or (4) under a board order for disciplinary action. The suspension shall remain in effect until lifted by the board pursuant to the board's receipt of a petition from the regulated facility, along with the current name and address of the owner of the facility.
- (h) An individual licensed or registered by the board shall maintain a current name and home address with the board and shall notify the board in writing within 30 days of any change in name or home address. An individual regulated by the board shall also maintain a current business address with the board as required by section 214.073. For an individual, if a name change only is requested, the regulated individual must request a revised license or registration. The board may require the individual to substantiate

43.2

43.3

43.4

43.5

43.6

43.7

43.8

43.9

43.10

43.11

43.12

43.13

43.14

43.15

43.16

43.17

43.18

43.19

43.20

43.21

43.22

43.23

43.24

43.25

43.26

43.27

43.28

43.29

43.30

43.31

43.32

43.33

43.34

43.35

43.36

the name change by submitting official documentation from a court of law or agency authorized under law to receive and officially record a name change. In the case of an individual, if an address change only is requested, no request for a revised license or registration is required. If the current license or registration of an individual has been lost, stolen, or destroyed, the individual shall provide a written explanation to the board.

(i) A facility licensed or registered by the board shall maintain a current name and address with the board. A facility shall notify the board in writing within 30 days of any change in name. A facility licensed or registered by the board but located outside of the state must notify the board within 30 days of an address change. A facility licensed or registered by the board and located within the state must notify the board at least 60 days in advance of a change of address that will result from the move of the facility to a different location and must pass an inspection at the new location as required by the board. If the current license or registration of a facility has been lost, stolen, or destroyed, the facility shall provide a written explanation to the board.

Subd. 4. Effective dates. A suspension, revocation, condition, limitation, qualification, or restriction of a license or registration shall be in effect pending determination of an appeal. A revocation of a license pursuant to subdivision 1 is not appealable and shall remain in effect indefinitely.

Subd. 5. Conditions on reissued license. In its discretion, the board may restore and reissue a license or registration issued under this chapter, but as a condition thereof may impose any disciplinary or corrective measure that it might originally have imposed.

Subd. 6. Temporary suspension of license for pharmacists. In addition to any other remedy provided by law, the board may, without a hearing, temporarily suspend the license of a pharmacist if the board finds that the pharmacist has violated a statute or rule that the board is empowered to enforce and continued practice by the pharmacist would create a serious risk of harm to the public. The suspension shall take effect upon written notice to the pharmacist, specifying the statute or rule violated. The suspension shall remain in effect until the board issues a final order in the matter after a hearing. At the time it issues the suspension notice, the board shall schedule a disciplinary hearing to be held pursuant to the Administrative Procedure Act. The pharmacist shall be provided with at least 20 days' notice of any hearing held pursuant to this subdivision. The hearing shall be scheduled to begin no later than 30 days after the issuance of the suspension order.

Subd. 7. Temporary suspension of license for pharmacist interns, pharmacy technicians, and controlled substance researchers. In addition to any other remedy provided by law, the board may, without a hearing, temporarily suspend the registration of a pharmacist intern, pharmacy technician, or controlled substance researcher if the board

44.2

44.3

44.4

44.5

44.6

44.7

44.8

44.9

44.10

44.11

44.12

44.13

44.14

44.15

44.16

44.17

44.18

44.19

44.20

44.21

44.22

44.23

44.24

44.25

44.26

44.27

44.28

44.29

44.30

44.31

44.32

44.33

44.34

44.35

44.36

finds that the registrant has violated a statute or rule that the board is empowered to enforce and continued registration of the registrant would create a serious risk of harm to the public. The suspension shall take effect upon written notice to the registrant, specifying the statute or rule violated. The suspension shall remain in effect until the board issues a final order in the matter after a hearing. At the time it issues the suspension notice, the board shall schedule a disciplinary hearing to be held pursuant to the Administrative Procedure Act. The licensee or registrant shall be provided with at least 20 days' notice of any hearing held pursuant to this subdivision. The hearing shall be scheduled to begin no later than 30 days after the issuance of the suspension order. Subd. 8. Temporary suspension of license for pharmacies, drug wholesalers,

DM

drug manufacturers, medical gas manufacturers, and medical gas distributors. In addition to any other remedy provided by law, the board may, without a hearing, temporarily suspend the license or registration of a pharmacy, drug wholesaler, drug manufacturer, medical gas manufacturer, or medical gas distributor if the board finds that the licensee or registrant has violated a statute or rule that the board is empowered to enforce and continued operation of the licensed facility would create a serious risk of harm to the public. The suspension shall take effect upon written notice to the licensee or registrant, specifying the statute or rule violated. The suspension shall remain in effect until the board issues a final order in the matter after a hearing. At the time it issues the suspension notice, the board shall schedule a disciplinary hearing to be held pursuant to the Administrative Procedure Act. The licensee or registrant shall be provided with at least 20 days' notice of any hearing held pursuant to this subdivision. The hearing shall be scheduled to begin no later than 30 days after the issuance of the suspension order.

Subd. 9. Evidence. In disciplinary actions alleging a violation of subdivision 2, clause (4), (5), (6), or (7), a copy of the judgment or proceeding under the seal of the court administrator or of the administrative agency that entered the same shall be admissible into evidence without further authentication and shall constitute prima facie evidence of the contents thereof.

Subd. 10. Mental examination; access to medical data. (a) If the board has probable cause to believe that an individual licensed or registered by the board falls under subdivision 2, clause (14), it may direct the individual to submit to a mental or physical examination. For the purpose of this subdivision, every licensed or registered individual is deemed to have consented to submit to a mental or physical examination when directed in writing by the board and further to have waived all objections to the admissibility of the examining practitioner's testimony or examination reports on the grounds that the same constitute a privileged communication. Failure of a licensed or registered individual to

45.2

45.3

45.4

45.5

45.6

45.7

45.8

45.9

45.10

45.11

45.12

45.13

45.14

45.15

45.16

45.17

45.18

45.19

45.20

45.21

45.22

45.23

45.24

45.25

45.26

45.27

45.28

45.29

45.30

45.31

45.32

45.33

45.34

45.35

45.36

submit to an examination when directed constitutes an admission of the allegations against the individual, unless the failure was due to circumstances beyond the individual's control, in which case a default and final order may be entered without the taking of testimony or presentation of evidence. Pharmacists affected under this paragraph shall at reasonable intervals be given an opportunity to demonstrate that they can resume the competent practice of the profession of pharmacy with reasonable skill and safety to the public. Pharmacist interns, pharmacy technicians, or controlled substance researchers affected under this paragraph shall at reasonable intervals be given an opportunity to demonstrate that they can competently resume the duties that can be performed, under this chapter or the rules of the board, by similarly registered persons with reasonable skill and safety to the public. In any proceeding under this paragraph, neither the record of proceedings nor the orders entered by the board shall be used against a licensed or registered individual in any other proceeding.

(b) In addition to ordering a physical or mental examination, the board may, notwithstanding section 13.384, 144.651, or any other law limiting access to medical or other health data, obtain medical data and health records relating to an individual licensed or registered by the board, or to an applicant for licensure or registration, without the individual's consent, if the board has probable cause to believe that the individual falls under subdivision 2, clause (14). The medical data may be requested from a provider, as defined in section 144.291, subdivision 2, paragraph (h), an insurance company, or a government agency, including the Department of Human Services. A provider, insurance company, or government agency shall comply with any written request of the board under this subdivision and is not liable in any action for damages for releasing the data requested by the board if the data are released pursuant to a written request under this subdivision, unless the information is false and the provider giving the information knew, or had reason to believe, the information was false. Information obtained under this subdivision is classified as private under sections 13.01 to 13.87.

Subd. 11. Tax clearance certificate. (a) In addition to the provisions of subdivision 1, the board may not issue or renew a license or registration if the commissioner of revenue notifies the board and the licensee or applicant for a license that the licensee or applicant owes the state delinquent taxes in the amount of \$500 or more. The board may issue or renew the license or registration only if (1) the commissioner of revenue issues a tax clearance certificate, and (2) the commissioner of revenue or the licensee, registrant, or applicant forwards a copy of the clearance to the board. The commissioner of revenue may issue a clearance certificate only if the licensee, registrant, or applicant does not owe the state any uncontested delinquent taxes.

46.1	(b) For purposes of this subdivision, the following terms have the meanings given.
46.2	(1) "Taxes" are all taxes payable to the commissioner of revenue, including penalties
46.3	and interest due on those taxes.
46.4	(2) "Delinquent taxes" do not include a tax liability if (i) an administrative or court
46.5	action that contests the amount or validity of the liability has been filed or served, (ii) the
46.6	appeal period to contest the tax liability has not expired, or (iii) the licensee or applicant
46.7	has entered into a payment agreement to pay the liability and is current with the payments.
46.8	(c) In lieu of the notice and hearing requirements of subdivision 1, when a licensee,
46.9	registrant, or applicant is required to obtain a clearance certificate under this subdivision,
46.10	a contested case hearing must be held if the licensee or applicant requests a hearing in
46.11	writing to the commissioner of revenue within 30 days of the date of the notice provided
46.12	in paragraph (a). The hearing must be held within 45 days of the date the commissioner of
46.13	revenue refers the case to the Office of Administrative Hearings. Notwithstanding any law
46.14	to the contrary, the licensee or applicant must be served with 20 days' notice in writing
46.15	specifying the time and place of the hearing and the allegations against the licensee or
46.16	applicant. The notice may be served personally or by mail.
46.17	(d) A licensee or applicant must provide the licensee's or applicant's Social Security
46.18	number and Minnesota business identification number on all license applications. Upon
46.19	request of the commissioner of revenue, the board must provide to the commissioner of
46.20	revenue a list of all licensees and applicants that includes the licensee's or applicant's
46.21	name, address, Social Security number, and business identification number. The
46.22	commissioner of revenue may request a list of the licensees and applicants no more than
46.23	once each calendar year.
46.24	Subd. 12. Limitation. No board proceeding against a regulated person or facility
46.25	shall be instituted unless commenced within seven years from the date of the commission
46 26	of some portion of the offense or misconduct complained of except for alleged violations

Sec. 4. [151.072] REPORTING OBLIGATIONS.

of subdivision 2, clause (21).

46.27

46.28

46.29

46.30

46.31

46.32

46.33

46.34

46.35

Subdivision 1. Permission to report. A person who has knowledge of any conduct constituting grounds for discipline under the provisions of this chapter or the rules of the board may report the violation to the board.

Subd. 2. **Pharmacies.** A pharmacy located in this state must report to the board any discipline that is related to an incident involving conduct that would constitute grounds for discipline under the provisions of this chapter or the rules of the board, that is taken by the pharmacy or any of its administrators against a pharmacist, pharmacist intern, or

47.2

47.3

47.4

47.5

47.6

47.7

47.8

47.9

47.10

47.11

47.12

47.13

47.14

47.15

47.16

47.17

47.18

47.19

47.20

47.21

47.22

47.23

47.24

47.25

47.26

47.27

47.30

47.31

47.32

47.33

47.34

47.35

pharmacy technician, including the termination of employment of the individual or the revocation, suspension, restriction, limitation, or conditioning of an individual's ability to practice or work at or on behalf of the pharmacy. The pharmacy shall also report the resignation of any pharmacist, pharmacist intern, or technician prior to the conclusion of any disciplinary proceeding, or prior to the commencement of formal charges but after the individual had knowledge that formal charges were contemplated or in preparation. Each report made under this subdivision must state the nature of the action taken and state in detail the reasons for the action. Failure to report violations as required by this subdivision is a basis for discipline pursuant to section 151.071, subdivision 2, clause (8).

DM

Subd. 3. Licensees and registrants of the board. A licensee or registrant of the board shall report to the board personal knowledge of any conduct that the person reasonably believes constitutes grounds for disciplinary action under this chapter or the rules of the board by any pharmacist, pharmacist intern, pharmacy technician, or controlled substance researcher, including any conduct indicating that the person may be professionally incompetent, or may have engaged in unprofessional conduct or may be medically or physically unable to engage safely in the practice of pharmacy or to carry out the duties permitted to the person by this chapter or the rules of the board. Failure to report violations as required by this subdivision is a basis for discipline pursuant to section 151.071, subdivision 2, clause (20).

- Subd. 4. **Self-reporting.** A licensee or registrant of the board shall report to the board any personal action that would require that a report be filed with the board pursuant to subdivision 2.
- Subd. 5. **Deadlines; forms.** Reports required by subdivisions 2 to 4 must be submitted not later than 30 days after the occurrence of the reportable event or transaction. The board may provide forms for the submission of reports required by this section, may require that reports be submitted on the forms provided, and may adopt rules necessary to assure prompt and accurate reporting.
- Subd. 6. **Subpoenas.** The board may issue subpoenas for the production of any reports required by subdivisions 2 to 4 or any related documents.

Sec. 5. [151.073] IMMUNITY.

Subdivision 1. Reporting. Any person, health care facility, business, or organization is immune from civil liability or criminal prosecution for submitting in good faith a report to the board under section 151.072 or for otherwise reporting in good faith to the board violations or alleged violations of this chapter or the rules of the board. All such reports are investigative data as defined in chapter 13.

SF1484

48.1

48.2

48.3

48.4

48.5

48.6

48.7

48.8

48.9

48.10

48.11

48.12

48.13

48.14

48.15

48.16

48.17

48.18

48.19

48.20

48.21

48.22

48.23

48.24

48.25

48.26

48.27

48.28

48.29

48.30

48.31

48.32

48.33

Subd. 2. **Investigation.** (a) Members of the board and persons employed by the board or engaged on behalf of the board in the investigation of violations and in the preparation and management of charges or violations of this chapter of the rules of the board, or persons participating in the investigation or testifying regarding charges of violations, are immune from civil liability and criminal prosecution for any actions, transactions, or publications in the execution of, or relating to, their duties under this chapter or the rules of the board.

3rd Engrossment

(b) Members of the board and persons employed by the board or engaged in maintaining records and making reports regarding adverse health care events are immune from civil liability and criminal prosecution for any actions, transactions, or publications in the execution of, or relating to, their duties under section 151.301.

Sec. 6. [151.074] LICENSEE OR REGISTRANT COOPERATION.

An individual who is licensed or registered by the board, who is the subject of an investigation by or on behalf of the board, shall cooperate fully with the investigation.

An owner or employee of a facility that is licensed or registered by the board, when the facility is the subject of an investigation by or on behalf of the board, shall cooperate fully with the investigation. Cooperation includes responding fully and promptly to any question raised by, or on behalf of, the board relating to the subject of the investigation and providing copies of patient pharmacy records and other relevant records, as reasonably requested by the board, to assist the board in its investigation. The board shall maintain any records obtained pursuant to this section as investigative data pursuant to chapter 13.

Sec. 7. [151.075] DISCIPLINARY RECORD ON JUDICIAL REVIEW.

Upon judicial review of any board disciplinary action taken under this chapter, the reviewing court shall seal the administrative record, except for the board's final decision, and shall not make the administrative record available to the public.

Sec. 8. Minnesota Statutes 2012, section 151.211, is amended to read:

151.211 RECORDS OF PRESCRIPTIONS.

Subdivision 1. Retention of prescription drug orders. All prescriptions dispensed prescription drug orders shall be kept on file at the location in from which such dispensing occurred of the ordered drug occurs for a period of at least two years. Prescription drug orders that are electronically prescribed must be kept on file in the format in which they were originally received. Written or printed prescription drug orders and verbal prescription drug orders reduced to writing, must be kept on file as received or transcribed, except that such orders may be kept in an electronic format as allowed by the board.

DM

Electronic systems used to process and store prescription drug orders must be compliant 49.1 49.2 with the requirements of this chapter and the rules of the board. Prescription drug orders that are stored in an electronic format, as permitted by this subdivision, may be kept on 49.3 file at a remote location provided that they are readily and securely accessible from the 49.4 location at which dispensing of the ordered drug occurred. 49.5 49.6 Subd. 2. **Refill requirements.** No A prescription shall drug order may be refilled except only with the written, electronic, or verbal consent of the prescriber and in 49.7 accordance with the requirements of this chapter, the rules of the board, and where 49.8 49.9 applicable, section 152.11. The date of such refill must be recorded and initialed upon the original prescription drug order, or within the electronically maintained record of the 49.10 original prescription drug order, by the pharmacist, pharmacist intern, or practitioner 49.11 49.12 who refills the prescription. Sec. 9. [151.251] COMPOUNDING. 49.13 49.14 Subdivision 1. Exemption from manufacturing licensure requirement. Section 151.252 shall not apply to: 49.15 (1) a practitioner engaged in extemporaneous compounding, anticipatory 49.16 compounding, or compounding not done pursuant to a prescription drug order when 49.17 permitted by this chapter or the rules of the board; and 49.18 49.19 (2) a pharmacy in which a pharmacist is engaged in extemporaneous compounding, anticipatory compounding, or compounding not done pursuant to a prescription drug order 49.20 when permitted by this chapter or the rules of the board. 49.21 Subd. 2. Compounded drug. A drug product may be compounded under this 49.22 section if a pharmacist or practitioner: 49.23 (a) compounds the drug product using bulk drug substances, as defined in the federal 49.24 49.25 regulations published in Code of Federal Regulations, title 21, section 207.3(a)(4): 49.26 (1) that: (i) comply with the standards of an applicable United States Pharmacopoeia 49.27 or National Formulary monograph, if a monograph exists, and the United States 49.28 Pharmacopoeia chapter on pharmacy compounding; 49.29 (ii) if such a monograph does not exist, are drug substances that are components of 49.30 drugs approved for use in this country by the United States Food and Drug Administration; 49.31 49.32 or (iii) if such a monograph does not exist and the drug substance is not a component of 49.33 49.34 a drug approved for use in this country by the United States Food and Drug Administration, that appear on a list developed by the United States Food and Drug Administration through 49.35

50.1	regulations issued by the secretary of the federal Department of Health and Human
50.2	Services pursuant to section 503a of the Food, Drug and Cosmetic Act under paragraph (d);
50.3	(2) that are manufactured by an establishment that is registered under section 360
50.4	of the federal Food, Drug and Cosmetic Act, including a foreign establishment that is
50.5	registered under section 360(i) of that act; and
50.6	(3) that are accompanied by valid certificates of analysis for each bulk drug substance;
50.7	(b) compounds the drug product using ingredients, other than bulk drug substances,
50.8	that comply with the standards of an applicable United States Pharmacopoeia or National
50.9	Formulary monograph, if a monograph exists, and the United States Pharmacopoeia
50.10	chapters on pharmacy compounding;
50.11	(c) does not compound a drug product that appears on a list published by the secretary
50.12	of the federal Department of Health and Human Services in the Federal Register of drug
50.13	products that have been withdrawn or removed from the market because such drug products
50.14	or components of such drug products have been found to be unsafe or not effective;
50.15	(d) does not compound any drug products that are essentially copies of a
50.16	commercially available drug product; and
50.17	(e) does not compound any drug product that has been identified pursuant to
50.18	United States Code, title 21, section 353a, as a drug product that presents demonstrable
50.19	difficulties for compounding that reasonably demonstrate an adverse effect on the safety
50.20	or effectiveness of that drug product.
50.21	The term "essentially a copy of a commercially available drug product" does not
50.22	include a drug product in which there is a change, made for an identified individual
50.23	patient, that produces for that patient a significant difference, as determined by the
50.24	prescribing practitioner, between the compounded drug and the comparable commercially
50.25	available drug product.
50.26	Subd. 3. Exceptions. This section shall not apply to:
50.27	(1) compounded positron emission tomography drugs as defined in section 151.01,
50.28	subdivision 38; or
50.29	(2) radiopharmaceuticals.
50.30	Sec. 10. Minnesota Statutes 2013 Supplement, section 151.252, is amended by adding
50.31	a subdivision to read:
50.32	Subd. 1a. Outsourcing facility. (a) No person shall act as an outsourcing facility
50.33	without first obtaining a license from the board and paying any applicable manufacturer
50.34	licensing fee specified in section 151.065.

51.2

51.3

51.4

51.5

51.6

51.7

51.8

51.9

51.10

51.11

51.12

51.13

51.14

51.15

51.16

51.17

51.18

51.19

51.20

51.21

51.22

51.23

51.24

51.25

51.26

51.27

51.28

51.29

51.30

51.31

51.32

51.33

51.34

51.35

(b) Application for an outsourcing facility license under this section shall be made
in a manner specified by the board and may differ from the application required of other
drug manufacturers.

- (c) No license shall be issued or renewed for an outsourcing facility unless the applicant agrees to operate in a manner prescribed for outsourcing facilities by federal and state law and according to Minnesota Rules.
- (d) No license shall be issued or renewed for an outsourcing facility unless the applicant supplies the board with proof of such registration by the United States Food and Drug Administration as required by United States Code, title 21, section 353b.
- (e) No license shall be issued or renewed for an outsourcing facility that is required to be licensed or registered by the state in which it is physically located unless the applicant supplies the board with proof of such licensure or registration. The board may establish, by rule, standards for the licensure of an outsourcing facility that is not required to be licensed or registered by the state in which it is physically located.
- (f) The board shall require a separate license for each outsourcing facility located within the state and for each outsourcing facility located outside of the state at which drugs that are shipped into the state are prepared.
- (g) The board shall not issue an initial or renewed license for an outsourcing facility unless the facility passes an inspection conducted by an authorized representative of the board. In the case of an outsourcing facility located outside of the state, the board may require the applicant to pay the cost of the inspection, in addition to the license fee in section 151.065, unless the applicant furnishes the board with a report, issued by the appropriate regulatory agency of the state in which the facility is located or by the United States Food and Drug Administration, of an inspection that has occurred within the 24 months immediately preceding receipt of the license application by the board. The board may deny licensure unless the applicant submits documentation satisfactory to the board that any deficiencies noted in an inspection report have been corrected.
 - Sec. 11. Minnesota Statutes 2012, section 151.26, is amended to read:

151.26 EXCEPTIONS.

Subdivision 1. **Generally.** Nothing in this chapter shall subject a person duly licensed in this state to practice medicine, dentistry, or veterinary medicine, to inspection by the State Board of Pharmacy, nor prevent the person from administering drugs, medicines, chemicals, or poisons in the person's practice, nor prevent a duly licensed practitioner from furnishing to a patient properly packaged and labeled drugs, medicines, chemicals, or poisons as may be considered appropriate in the treatment of such patient;

unless the person is engaged in the dispensing, sale, or distribution of drugs and the board provides reasonable notice of an inspection.

Except for the provisions of section 151.37, nothing in this chapter applies to or interferes with the dispensing, in its original package and at no charge to the patient, of a legend drug, other than a controlled substance, that was packaged by a manufacturer and provided to the dispenser for distribution dispensing as a professional sample, so long as the sample is prepared and distributed pursuant to Code of Federal Regulations, title 21, section 203, subpart D.

Nothing in this chapter shall prevent the sale of drugs, medicines, chemicals, or poisons at wholesale to licensed physicians, dentists and veterinarians for use in their practice, nor to hospitals for use therein.

Nothing in this chapter shall prevent the sale of drugs, chemicals, or poisons either at wholesale or retail for use for commercial purposes, or in the arts, nor interfere with the sale of insecticides, as defined in Minnesota Statutes 1974, section 24.069, and nothing in this chapter shall prevent the sale of common household preparations and other drugs, chemicals, and poisons sold exclusively for use for nonmedicinal purposes—; provided that this exception does not apply to any compound, substance, or derivative that is not approved for human consumption by the United States Food and Drug Administration or specifically permitted for human consumption under Minnesota law and, when introduced into the body, induces an effect similar to that of a Schedule I or Schedule II controlled substance listed in section 152.02, subdivisions 2 and 3, or Minnesota Rules, parts 6800.4210 and 6800.4220, regardless of whether the substance is marketed for the purpose of human consumption.

Nothing in this chapter shall apply to or interfere with the vending or retailing of any nonprescription medicine or drug not otherwise prohibited by statute which that is prepackaged, fully prepared by the manufacturer or producer for use by the consumer, and labeled in accordance with the requirements of the state or federal Food and Drug Act; nor to the manufacture, wholesaling, vending, or retailing of flavoring extracts, toilet articles, cosmetics, perfumes, spices, and other commonly used household articles of a chemical nature, for use for nonmedicinal purposes; provided that this exception does not apply to any compound, substance, or derivative that is not approved for human consumption by the United States Food and Drug Administration or specifically permitted for human consumption under Minnesota law that, when introduced into the body, induces an effect similar to that of a Schedule I or Schedule II controlled substance listed in section 152.02, subdivisions 2 and 3, or Minnesota Rules, parts 6800.4210 and 6800.4220, regardless of whether the substance is marketed for the purpose of human consumption. Nothing in

52.1

52.2

52.3

52.4

52.5

52.6

52.7

52.8

52.9

52.10

52.11

52.12

52.13

52.14

52.15

52.16

52.17

52.18

52.19

52.20

52.21

52.22

52.23

52.24

52.25

52.26

52.27

52.28

52.29

52.30

52.31

52.32

52.33

52.34

52.35

52.36

53.4

53.5

53.6

53.7

53.8

53.9

53.10

53.11

53.12

53.13

53.14

53.15

53.16

53.17

53.18

53.19

53.20

53.21

53.22

53.23

53.24

53.25

53.26

53.27

53.28

53.29

53.30

53.31

53.32

53.33

DM 3rd Engrossment this chapter shall prevent the sale of drugs or medicines by licensed pharmacists at a 53.1 discount to persons over 65 years of age. 53.2

Sec. 12. Minnesota Statutes 2012, section 151.34, is amended to read:

151.34 PROHIBITED ACTS.

It shall be unlawful to:

- (1) manufacture, sell or deliver, hold or offer for sale any drug that is adulterated or misbranded;
 - (2) adulterate or misbrand any drug;
- (3) receive in commerce any drug that is adulterated or misbranded, and to deliver or proffer delivery thereof for pay or otherwise;
- (4) refuse to permit entry or inspection, or to permit the taking of a sample, or to permit access to or copying of any record as authorized by this chapter;
 - (5) remove or dispose of a detained or embargoed article in violation of this chapter;
- (6) alter, mutilate, destroy, obliterate, or remove the whole or any part of the labeling of, or to do any other act with respect to a drug, if such act is done while such drug is held for sale and results in such drug being adulterated or misbranded;
- (7) use for a person's own advantage or to reveal other than to the board or its authorized representative or to the courts when required in any judicial proceeding under this chapter any information acquired under authority of this chapter concerning any method or process which that is a trade secret and entitled to protection;
- (8) use on the labeling of any drug any representation or suggestion that an application with respect to such drug is effective under the federal act or that such drug complies with such provisions;
- (9) in the case of a manufacturer, packer, or distributor offering legend drugs for sale within this state, fail to maintain for transmittal or to transmit, to any practitioner licensed by applicable law to administer such drug who makes written request for information as to such drug, true and correct copies of all printed matter which that is required to be included in any package in which that drug is distributed or sold, or such other printed matter as is approved under the federal act. Nothing in this paragraph shall be construed to exempt any person from any labeling requirement imposed by or under provisions of this chapter;
 - (10) conduct a pharmacy without a pharmacist in charge;
 - (11) dispense a legend drug without first obtaining a valid prescription for that drug;
- (12) conduct a pharmacy without proper registration with the board;
- (13) practice pharmacy without being licensed to do so by the board; or 53.34

54.2

54.3

54.4

54.5

54.6

54.7

54.8

54.9

54.10

54.11

54.12

54.13

54.14

54.15

54.16

54.17

54.18

54.19

54.20

54.21

54.22

54.23

54.24

54.25

54.26

54.27

54.28

54.29

54.30

54.31

54.32

54.33

54.34

S1484-3

(14) sell at retail federally restricted medical gases without proper registration with the board except as provided in this chapter-; or

(15) sell any compound, substance, or derivative that is not approved for human consumption by the United States Food and Drug Administration or specifically permitted for human consumption under Minnesota law and, when introduced into the body, induces an effect similar to that of a Schedule I or Schedule II controlled substance listed in section 152.02, subdivisions 2 and 3, or Minnesota Rules, parts 6800.4210 and 6800.4220, regardless of whether the substance is marketed for the purpose of human consumption.

EFFECTIVE DATE. This section is effective August 1, 2014, and applies to sales on or after that date.

Sec. 13. Minnesota Statutes 2012, section 151.35, is amended to read:

151.35 DRUGS, ADULTERATION.

A drug shall be deemed to be adulterated:

- (1) if it consists in whole or in part of any filthy, putrid or decomposed substance; or if it has been produced, prepared, packed, or held under unsanitary conditions whereby it may have been rendered injurious to health, or whereby it may have been contaminated with filth; or if the methods used in, or the facilities or controls used for, its manufacture, processing, packing, or holding do not conform to or are not operated or administered in conformity with current good manufacturing practice as required under the federal act to assure that such drug is safe and has the identity, strength, quality, and purity characteristics, which it purports or is represented to possess; or the facility in which it was produced was not registered by the United States Food and Drug Administration or licensed by the board; or, its container is composed, in whole or in part, of any poisonous or deleterious substance which may render the contents injurious to health; or it bears or contains, for purposes of coloring only, a color additive which is unsafe within the meaning of the federal act, or it is a color additive, the intended use of which in or on drugs is for the purposes of coloring only, and is unsafe within the meaning of the federal act;
- (2) if it purports to be or is represented as a drug the name of which is recognized in the United States Pharmacopoeia or the National Formulary, and its strength differs from, or its quality or purity falls below, the standard set forth therein. Such determination as to strength, quality, or purity shall be made in accordance with the tests or methods of assay set forth in such compendium, or in the absence of or inadequacy of such tests or methods of assay, those prescribed under authority of the federal act. No drug defined in the United States Pharmacopoeia or the National Formulary shall be deemed to be

DM

55.1	adulterated under this paragraph because it differs from the standard of strength, quality,
55.2	or purity therefor set forth in such compendium, if its difference in strength, quality, or
55.3	purity from such standard is plainly stated on its label;
55.4	(3) if it is not subject to the provisions of paragraph (2) of this section and its
55.5	strength differs from, or its purity or quality differs from that which it purports or is
55.6	represented to possess;
55.7	(4) if any substance has been mixed or packed therewith so as to reduce its quality or
55.8	strength, or substituted wholly or in part therefor.
55.9	Sec. 14. Minnesota Statutes 2012, section 151.361, subdivision 2, is amended to read:
55.10	Subd. 2. After January 1, 1983. (a) No legend drug in solid oral dosage form
55.11	may be manufactured, packaged or distributed for sale in this state after January 1, 1983
55.12	unless it is clearly marked or imprinted with a symbol, number, company name, words,
55.13	letters, national drug code or other mark uniquely identifiable to that drug product. An
55.14	identifying mark or imprint made as required by federal law or by the federal Food and
55.15	Drug Administration shall be deemed to be in compliance with this section.
55.16	(b) The Board of Pharmacy may grant exemptions from the requirements of this
55.17	section on its own initiative or upon application of a manufacturer, packager, or distributor
55.18	indicating size or other characteristics which that render the product impractical for the
55.19	imprinting required by this section.
55.20	(c) The provisions of clauses (a) and (b) shall not apply to any of the following:
55.21	(1) Drugs purchased by a pharmacy, pharmacist, or licensed wholesaler prior to
55.22	January 1, 1983, and held in stock for resale.
55.23	(2) Drugs which are manufactured by or upon the order of a practitioner licensed by
55.24	law to prescribe or administer drugs and which are to be used solely by the patient for
55.25	whom prescribed.
55.26	Sec. 15. Minnesota Statutes 2012, section 151.37, as amended by Laws 2013, chapter
55.27	43, section 30, Laws 2013, chapter 55, section 2, and Laws 2013, chapter 108, article
55.28	10, section 5, is amended to read:
55.29	151.37 LEGEND DRUGS, WHO MAY PRESCRIBE, POSSESS.
55.30	Subdivision 1. Prohibition. Except as otherwise provided in this chapter, it shall be
55.31	unlawful for any person to have in possession, or to sell, give away, barter, exchange, or

55.32

55.33

55.34

distribute a legend drug.

Subd. 2. Prescribing and filing. (a) A licensed practitioner in the course of

professional practice only, may prescribe, administer, and dispense a legend drug, and

56.2

56.3

56.4

56.5

56.6

56.7

56.8

56.9

56.10

56.11

56.12

56.13

56.14

56.15

56.16

56.17

56.18

56.19

56.20

56.21

56.22

56.23

56.24

56.25

56.26

56.27

56.28

56.29

56.30

56.31

56.32

56.33

56.34

56.35

56.36

may cause the same to be administered by a nurse, a physician assistant, or medical student or resident under the practitioner's direction and supervision, and may cause a person who is an appropriately certified, registered, or licensed health care professional to prescribe, dispense, and administer the same within the expressed legal scope of the person's practice as defined in Minnesota Statutes. A licensed practitioner may prescribe a legend drug, without reference to a specific patient, by directing a licensed dietitian or licensed nutritionist, pursuant to section 148.634; a nurse, pursuant to section 148.235, subdivisions 8 and 9; physician assistant; medical student or resident; or pharmacist according to section 151.01, subdivision 27, to adhere to a particular practice guideline or protocol when treating patients whose condition falls within such guideline or protocol, and when such guideline or protocol specifies the circumstances under which the legend drug is to be prescribed and administered. An individual who verbally, electronically, or otherwise transmits a written, oral, or electronic order, as an agent of a prescriber, shall not be deemed to have prescribed the legend drug. This paragraph applies to a physician assistant only if the physician assistant meets the requirements of section 147A.18.

- (b) The commissioner of health, if a licensed practitioner, or a person designated by the commissioner who is a licensed practitioner, may prescribe a legend drug to an individual or by protocol for mass dispensing purposes where the commissioner finds that the conditions triggering section 144.4197 or 144.4198, subdivision 2, paragraph (b), exist. The commissioner, if a licensed practitioner, or a designated licensed practitioner, may prescribe, dispense, or administer a legend drug or other substance listed in subdivision 10 to control tuberculosis and other communicable diseases. The commissioner may modify state drug labeling requirements, and medical screening criteria and documentation, where time is critical and limited labeling and screening are most likely to ensure legend drugs reach the maximum number of persons in a timely fashion so as to reduce morbidity and mortality.
- (c) A licensed practitioner that dispenses for profit a legend drug that is to be administered orally, is ordinarily dispensed by a pharmacist, and is not a vaccine, must file with the practitioner's licensing board a statement indicating that the practitioner dispenses legend drugs for profit, the general circumstances under which the practitioner dispenses for profit, and the types of legend drugs generally dispensed. It is unlawful to dispense legend drugs for profit after July 31, 1990, unless the statement has been filed with the appropriate licensing board. For purposes of this paragraph, "profit" means (1) any amount received by the practitioner in excess of the acquisition cost of a legend drug for legend drugs that are purchased in prepackaged form, or (2) any amount received by the practitioner in excess of the acquisition cost of a legend drug plus the cost of

57.2

57.3

57.4

57.5

57.6

57.7

57.8

57.9

57.10

57.11

57.12

57.13

57.14

57.16

57.18

57.19

57.20

57.21

57.22

57.23

57.24

57.25

57.26

57.27

57.28

57.29

57.30

57.31

57.32

making the drug available if the legend drug requires compounding, packaging, or other
treatment. The statement filed under this paragraph is public data under section 13.03.
This paragraph does not apply to a licensed doctor of veterinary medicine or a registered
pharmacist. Any person other than a licensed practitioner with the authority to prescribe,
dispense, and administer a legend drug under paragraph (a) shall not dispense for profit.
To dispense for profit does not include dispensing by a community health clinic when the
profit from dispensing is used to meet operating expenses.

- (d) A prescription of drug order for the following drugs is not valid, unless it can be established that the prescription of drug order was based on a documented patient evaluation, including an examination, adequate to establish a diagnosis and identify underlying conditions and contraindications to treatment:
 - (1) controlled substance drugs listed in section 152.02, subdivisions 3 to 5;
- (2) drugs defined by the Board of Pharmacy as controlled substances under section 152.02, subdivisions 7, 8, and 12;
- 57.15 (3) muscle relaxants;
 - (4) centrally acting analgesics with opioid activity;
- 57.17 (5) drugs containing butalbital; or
 - (6) phoshodiesterase type 5 inhibitors when used to treat erectile dysfunction.
 - (e) For the purposes of paragraph (d), the requirement for an examination shall be met if an in-person examination has been completed in any of the following circumstances:
 - (1) the prescribing practitioner examines the patient at the time the prescription or drug order is issued;
 - (2) the prescribing practitioner has performed a prior examination of the patient;
 - (3) another prescribing practitioner practicing within the same group or clinic as the prescribing practitioner has examined the patient;
 - (4) a consulting practitioner to whom the prescribing practitioner has referred the patient has examined the patient; or
 - (5) the referring practitioner has performed an examination in the case of a consultant practitioner issuing a prescription or drug order when providing services by means of telemedicine.
 - (f) Nothing in paragraph (d) or (e) prohibits a licensed practitioner from prescribing a drug through the use of a guideline or protocol pursuant to paragraph (a).
- 57.33 (g) Nothing in this chapter prohibits a licensed practitioner from issuing a 57.34 prescription or dispensing a legend drug in accordance with the Expedited Partner Therapy 57.35 in the Management of Sexually Transmitted Diseases guidance document issued by the 57.36 United States Centers for Disease Control.

S1484-3

58.5

58.6

58.7

58.8

58.9

58.10

58.11

58.12

58.13

58.14

58.15

58.16

58.17

58.18

58.19

58.20

58.21

58.22

58.23

58.24

58.25

58.26

58.27

58.28

58.29

58.30

58.31

58.32

58.33

58.34

58.35

(h) Nothing in paragraph (d) or (e) limits prescription, administration, or dispensing of legend drugs through a public health clinic or other distribution mechanism approved by the commissioner of health or a board of health in order to prevent, mitigate, or treat a pandemic illness, infectious disease outbreak, or intentional or accidental release of a biological, chemical, or radiological agent.

- (i) No pharmacist employed by, under contract to, or working for a pharmacy licensed under section 151.19, subdivision 1, may dispense a legend drug based on a prescription that the pharmacist knows, or would reasonably be expected to know, is not valid under paragraph (d).
- (j) No pharmacist employed by, under contract to, or working for a pharmacy licensed under section 151.19, subdivision 2, may dispense a legend drug to a resident of this state based on a prescription that the pharmacist knows, or would reasonably be expected to know, is not valid under paragraph (d).
- (k) Nothing in this chapter prohibits the commissioner of health, if a licensed practitioner, or, if not a licensed practitioner, a designee of the commissioner who is a licensed practitioner, from prescribing legend drugs for field-delivered therapy in the treatment of a communicable disease according to the Centers For Disease Control and Prevention Partner Services Guidelines.
- Subd. 2a. **Delegation.** A supervising physician may delegate to a physician assistant who is registered with the Board of Medical Practice and certified by the National Commission on Certification of Physician Assistants and who is under the supervising physician's supervision, the authority to prescribe, dispense, and administer legend drugs and medical devices, subject to the requirements in chapter 147A and other requirements established by the Board of Medical Practice in rules.
- Subd. 3. **Veterinarians.** A licensed doctor of veterinary medicine, in the course of professional practice only and not for use by a human being, may personally prescribe, administer, and dispense a legend drug, and may cause the same to be administered or dispensed by an assistant under the doctor's direction and supervision.
- Subd. 4. **Research.** (a) Any qualified person may use legend drugs in the course of a bona fide research project, but cannot administer or dispense such drugs to human beings unless such drugs are prescribed, dispensed, and administered by a person lawfully authorized to do so.
- (b) Drugs may be dispensed or distributed by a pharmacy licensed by the board for use by, or administration to, patients enrolled in a bona fide research study that is being conducted pursuant to either an investigational new drug application approved by the

S1484-3

59.1

59.2

59.3

59.4

59.5

59.6

59.7

59.8

59.9

59.10

59.11

59.12

59.13

59.14

59.15

59.16

59.17

59.18

59.19

59.20

59.21

59.22

59.23

59.24

59.25

59.26

59.27

59.28

59.29

59.30

59.31

59.32

59.33

59.34

United States Food and Drug Administration or that has been approved by an institutional review board. For the purposes of this subdivision only:

- (1) a prescription drug order is not required for a pharmacy to dispense a research drug, unless the study protocol requires the pharmacy to receive such an order;
- (2) notwithstanding the prescription labeling requirements found in this chapter or the rules promulgated by the board, a research drug may be labeled as required by the study protocol; and
- (3) dispensing and distribution of research drugs by pharmacies shall not be considered eompounding, manufacturing, or wholesaling under this chapter-; and
- (4) a pharmacy may compound drugs for research studies as provided in this subdivision but must follow applicable standards established by United States Pharmacopeia, chapter 795 or 797, for nonsterile and sterile compounding, respectively.
- (c) An entity that is under contract to a federal agency for the purpose of distributing drugs for bona fide research studies is exempt from the drug wholesaler licensing requirements of this chapter. Any other entity is exempt from the drug wholesaler licensing requirements of this chapter if the board finds that the entity is licensed or registered according to the laws of the state in which it is physically located and it is distributing drugs for use by, or administration to, patients enrolled in a bona fide research study that is being conducted pursuant to either an investigational new drug application approved by the United States Food and Drug Administration or that has been approved by an institutional review board.
- Subd. 5. Exclusion for course of practice. Nothing in this chapter shall prohibit the sale to, or the possession of, a legend drug by licensed drug wholesalers, licensed manufacturers, registered pharmacies, local detoxification centers, licensed hospitals, bona fide hospitals wherein animals are treated, or licensed pharmacists and licensed practitioners while acting within the course of their practice only.
- Subd. 6. Exclusion for course of employment. (a) Nothing in this chapter shall prohibit the possession of a legend drug by an employee, agent, or sales representative of a registered drug manufacturer, or an employee or agent of a registered drug wholesaler, or registered pharmacy, while acting in the course of employment.
- (b) Nothing in this chapter shall prohibit the following entities from possessing a legend drug for the purpose of disposing of the legend drug as pharmaceutical waste:
 - (1) a law enforcement officer;
 - (2) a hazardous waste transporter licensed by the Department of Transportation;
- (3) a facility permitted by the Pollution Control Agency to treat, store, or dispose of 59.35 hazardous waste, including household hazardous waste; 59.36

60.2

60.3

60.4

60.5

60.6

60.7

60.8

60.9

60.10

60.11

60.12

60.13

60.14

60.15

60.16

60.17

60.18

60.19

60.20

60.21

60.22

60.23

60.24

60.25

60.26

60.27

60.28

60.29

60.30

60.31

60.32

60.33

60.34

60.35

(4) a facility licensed by the Pollution Control Agency or a metropolitan county as a very small quantity generator collection program or a minimal generator;

- (5) a county that collects, stores, transports, or disposes of a legend drug pursuant to a program in compliance with applicable federal law or a person authorized by the county to conduct one or more of these activities; or
 - (6) a sanitary district organized under chapter 115, or a special law.
- Subd. 7. Exclusion for prescriptions. (a) Nothing in this chapter shall prohibit the possession of a legend drug by a person for that person's use when it has been dispensed to the person in accordance with a valid prescription issued by a practitioner.
- (b) Nothing in this chapter shall prohibit a person, for whom a legend drug has been dispensed in accordance with a written or oral prescription by a practitioner, from designating a family member, caregiver, or other individual to handle the legend drug for the purpose of assisting the person in obtaining or administering the drug or sending the drug for destruction.
- (c) Nothing in this chapter shall prohibit a person for whom a prescription drug has been dispensed in accordance with a valid prescription issued by a practitioner from transferring the legend drug to a county that collects, stores, transports, or disposes of a legend drug pursuant to a program in compliance with applicable federal law or to a person authorized by the county to conduct one or more of these activities.
- Subd. 8. Misrepresentation. It is unlawful for a person to procure, attempt to procure, possess, or control a legend drug by any of the following means:
 - (1) deceit, misrepresentation, or subterfuge;
 - (2) using a false name; or
- (3) falsely assuming the title of, or falsely representing a person to be a manufacturer, wholesaler, pharmacist, practitioner, or other authorized person for the purpose of obtaining a legend drug.
- Subd. 9. Exclusion for course of laboratory employment. Nothing in this chapter shall prohibit the possession of a legend drug by an employee or agent of a registered analytical laboratory while acting in the course of laboratory employment.
- Subd. 10. Purchase of drugs and other agents by commissioner of health. The commissioner of health, in preparation for and in carrying out the duties of sections 144.05, 144.4197, and 144.4198, may purchase, store, and distribute antituberculosis drugs, biologics, vaccines, antitoxins, serums, immunizing agents, antibiotics, antivirals, antidotes, other pharmaceutical agents, and medical supplies to treat and prevent communicable disease.

61.2

61.3

61.4

61.5

61.6

61.7

61.8

61.9

61.10

61.11

61.12

61.13

61.14

61.15

61.16

61.17

61.18

61.19

61.20

61.21

61.22

61.23

61.24

61.25

61.26

61.27

61.28

61.29

61.30

61.31

61.32

61.33

61.34

DM

3rd Engrossment

or from other hospitals or health care entities that are members of such organizations;

62.2

62.3

62.4

62.5

62.6

62.7

62.8

62.9

62.10

62.11

62.12

62.13

62.14

62.15

62.16

62.17

62.18

62.19

62.20

62.21

62.22

62.23

62.24

62.25

62.26

62.27

62.28

62.29

62.30

62.31

62.32

62.33

62.34

62.35

62.36

(3) the sale, purchase, or trade of a drug or an offer to sell, purchase, or trade a
drug by a charitable organization described in section 501(c)(3) of the Internal Revenue
Code of 1986, as amended through December 31, 1988, to a nonprofit affiliate of the
organization to the extent otherwise permitted by law;

- (4) the sale, purchase, or trade of a drug or offer to sell, purchase, or trade a drug among hospitals or other health care entities that are under common control;
- (5) the sale, purchase, or trade of a drug or offer to sell, purchase, or trade a drug for emergency medical reasons;
- (6) the sale, purchase, or trade of a drug, an offer to sell, purchase, or trade a drug, or the dispensing of a drug pursuant to a prescription;
- (7) the transfer of prescription or nonprescription drugs by a retail pharmacy to another retail pharmacy to alleviate a temporary shortage;
- (8) the distribution of prescription or nonprescription drug samples by manufacturers representatives; or
 - (9) the sale, purchase, or trade of blood and blood components.
- (b) "Wholesale drug distributor" means anyone engaged in wholesale drug distribution including, but not limited to, manufacturers; repackers repackagers; own-label distributors; jobbers; brokers; warehouses, including manufacturers' and distributors' warehouses, chain drug warehouses, and wholesale drug warehouses; independent wholesale drug traders; and pharmacies that conduct wholesale drug distribution. A wholesale drug distributor does not include a common carrier or individual hired primarily to transport prescription or nonprescription drugs.
- (c) "Manufacturer" means anyone who is engaged in the manufacturing, preparing, propagating, compounding, processing, packaging, repackaging, or labeling of a prescription drug has the meaning provided in section 151.01, subdivision 14b.
- (d) "Prescription drug" means a drug required by federal or state law or regulation to be dispensed only by a prescription, including finished dosage forms and active ingredients subject to United States Code, title 21, sections 811 and 812.
- (e) "Blood" means whole blood collected from a single donor and processed either for transfusion or further manufacturing.
- (f) "Blood components" means that part of blood separated by physical or mechanical means.
- (g) "Reverse distribution" means the receipt of prescription or nonprescription drugs received from or shipped to Minnesota locations for the purpose of returning the drugs to their producers or distributors.
 - (h) "Reverse distributor" means a person engaged in the reverse distribution of drugs.

63.2

63.3

63.4

63.5

63.6

63.7

63.8

63.9

63.10

63.11

63.12

63.13

63.14

63.16

63.17

63.18

63.19

63.20

63.21

63.22

63.23

63.24

63.25

63.26

63.27

63.28

63.29

63.30

63.31

63.32

63.33

63.34

- Sec. 17. Minnesota Statutes 2012, section 151.58, subdivision 2, is amended to read:
- Subd. 2. **Definitions.** For purposes of this section only, the terms defined in this subdivision have the meanings given.
- (a) "Automated drug distribution system" or "system" means a mechanical system approved by the board that performs operations or activities, other than compounding or administration, related to the storage, packaging, or dispensing of drugs, and collects, controls, and maintains all required transaction information and records.
- (b) "Health care facility" means a nursing home licensed under section 144A.02; a housing with services establishment registered under section 144D.01, subdivision 4, in which a home provider licensed under chapter 144A is providing centralized storage of medications; or a community behavioral health hospital or Minnesota sex offender program facility operated by the Department of Human Services.
- (c) "Managing pharmacy" means a pharmacy licensed by the board that controls and is responsible for the operation of an automated drug distribution system.
- Sec. 18. Minnesota Statutes 2012, section 151.58, subdivision 3, is amended to read: 63.15
 - Subd. 3. Authorization. A pharmacy may use an automated drug distribution system to fill prescription drug orders for patients of a health care facility provided that the policies and procedures required by this section have been approved by the board. The automated drug distribution system may be located in a health care facility that is not at the same location as the managing pharmacy. When located within a health care facility, the system is considered to be an extension of the managing pharmacy.
 - Sec. 19. Minnesota Statutes 2012, section 151.58, subdivision 5, is amended to read:
 - Subd. 5. Operation of automated drug distribution systems. (a) The managing pharmacy and the pharmacist in charge are responsible for the operation of an automated drug distribution system.
 - (b) Access to an automated drug distribution system must be limited to pharmacy and nonpharmacy personnel authorized to procure drugs from the system, except that field service technicians may access a system located in a health care facility for the purposes of servicing and maintaining it while being monitored either by the managing pharmacy, or a licensed nurse within the health care facility. In the case of an automated drug distribution system that is not physically located within a licensed pharmacy, access for the purpose of procuring drugs shall be limited to licensed nurses. Each person authorized to access the system must be assigned an individual specific access code. Alternatively, access to the system may be controlled through the use of biometric identification procedures. A

64.2

64.3

64.4

64.5

64.6

64.7

648

64.9

64.10

64.11

64.12

64.13

64.14

64.15

64.16

64.17

64.18

64.19

64.20

64.21

64.22

64.23

64.24

64.25

64.26

64.27

64.28

64.29

64.30

64.31

64.32

64.33

64.34

64.35

64.36

policy specifying time access parameters, including time-outs, logoffs, and lockouts, must be in place.

- (c) For the purposes of this section only, the requirements of section 151.215 are met if the following clauses are met:
- (1) a pharmacist employed by and working at the managing pharmacy, or at a pharmacy that is acting as a central services pharmacy for the managing pharmacy, pursuant to Minnesota Rules, part 6800.4075, must review, interpret, and approve all prescription drug orders before any drug is distributed from the system to be administered to a patient. A pharmacy technician may perform data entry of prescription drug orders provided that a pharmacist certifies the accuracy of the data entry before the drug can be released from the automated drug distribution system. A pharmacist employed by and working at the managing pharmacy must certify the accuracy of the filling of any cassettes, canisters, or other containers that contain drugs that will be loaded into the automated drug distribution system; and
- (2) when the automated drug dispensing system is located and used within the managing pharmacy, a pharmacist must personally supervise and take responsibility for all packaging and labeling associated with the use of an automated drug distribution system.
- (d) Access to drugs when a pharmacist has not reviewed and approved the prescription drug order is permitted only when a formal and written decision to allow such access is issued by the pharmacy and the therapeutics committee or its equivalent. The committee must specify the patient care circumstances in which such access is allowed, the drugs that can be accessed, and the staff that are allowed to access the drugs.
- (e) In the case of an automated drug distribution system that does not utilize bar coding in the loading process, the loading of a system located in a health care facility may be performed by a pharmacy technician, so long as the activity is continuously supervised, through a two-way audiovisual system by a pharmacist on duty within the managing pharmacy. In the case of an automated drug distribution system that utilizes bar coding in the loading process, the loading of a system located in a health care facility may be performed by a pharmacy technician or a licensed nurse, provided that the managing pharmacy retains an electronic record of loading activities.
- (f) The automated drug distribution system must be under the supervision of a pharmacist. The pharmacist is not required to be physically present at the site of the automated drug distribution system if the system is continuously monitored electronically by the managing pharmacy. A pharmacist on duty within a pharmacy licensed by the board must be continuously available to address any problems detected by the monitoring or to answer questions from the staff of the health care facility. The licensed pharmacy

may be the managing pharmacy or a pharmacy which is acting as a central services pharmacy, pursuant to Minnesota Rules, part 6800.4075, for the managing pharmacy.

- Sec. 20. Minnesota Statutes 2013 Supplement, section 152.02, subdivision 2, is amended to read:
- Subd. 2. **Schedule I.** (a) Schedule I consists of the substances listed in this subdivision.
- (b) Opiates. Unless specifically excepted or unless listed in another schedule, any of the following substances, including their analogs, isomers, esters, ethers, salts, and salts of isomers, esters, and ethers, whenever the existence of the analogs, isomers, esters, ethers, and salts is possible:
- 65.11 (1) acetylmethadol;
- 65.12 (2) allylprodine;

65.3

65.4

65.5

65.6

65.7

65.8

65.9

65.10

- (3) alphacetylmethadol (except levo-alphacetylmethadol, also known as
- 65.14 levomethadyl acetate);
- 65.15 (4) alphameprodine;
- 65.16 (5) alphamethadol;
- 65.17 (6) alpha-methylfentanyl benzethidine;
- 65.18 (7) betacetylmethadol;
- 65.19 (8) betameprodine;
- 65.20 (9) betamethadol;
- 65.21 (10) betaprodine;
- 65.22 (11) clonitazene;
- 65.23 (12) dextromoramide;
- 65.24 (13) diampromide;
- 65.25 (14) diethyliambutene;
- 65.26 (15) difenoxin;
- 65.27 (16) dimenoxadol;
- 65.28 (17) dimepheptanol;
- 65.29 (18) dimethyliambutene;
- 65.30 (19) dioxaphetyl butyrate;
- 65.31 (20) dipipanone;
- 65.32 (21) ethylmethylthiambutene;
- 65.33 (22) etonitazene;
- 65.34 (23) etoxeridine;
- 65.35 (24) furethidine;

66.1	(25) hydroxypethidine;
66.2	(26) ketobemidone;
66.3	(27) levomoramide;
66.4	(28) levophenacylmorphan;
66.5	(29) 3-methylfentanyl;
66.6	(30) acetyl-alpha-methylfentanyl;
66.7	(31) alpha-methylthiofentanyl;
66.8	(32) benzylfentanyl beta-hydroxyfentanyl;
66.9	(33) beta-hydroxy-3-methylfentanyl;
66.10	(34) 3-methylthiofentanyl;
66.11	(35) thenylfentanyl;
66.12	(36) thiofentanyl;
66.13	(37) para-fluorofentanyl;
66.14	(38) morpheridine;
66.15	(39) 1-methyl-4-phenyl-4-propionoxypiperidine;
66.16	(40) noracymethadol;
66.17	(41) norlevorphanol;
66.18	(42) normethadone;
66.19	(43) norpipanone;
66.20	(44) 1-(2-phenylethyl)-4-phenyl-4-acetoxypiperidine (PEPAP);
66.21	(45) phenadoxone;
66.22	(46) phenampromide;
66.23	(47) phenomorphan;
66.24	(48) phenoperidine;
66.25	(49) piritramide;
66.26	(50) proheptazine;
66.27	(51) properidine;
66.28	(52) propiram;
66.29	(53) racemoramide;
66.30	(54) tilidine;
66.31	(55) trimeperidine-;
66.32	(56) N-(1-Phenethylpiperidin-4-yl)-N-phenylacetamide (acetyl fentanyl).
66.33	(c) Opium derivatives. Any of the following substances, their analogs, salts, isomers,
66.34	and salts of isomers, unless specifically excepted or unless listed in another schedule,
66.35	whenever the existence of the analogs, salts, isomers, and salts of isomers is possible:
66.36	(1) acetorphine;

(2) acetyldihydrocodeine; 67.1 (3) benzylmorphine; 67.2 (4) codeine methylbromide; 67.3 (5) codeine-n-oxide; 67.4 (6) cyprenorphine; 67.5 (7) desomorphine; 67.6 (8) dihydromorphine; 67.7 (9) drotebanol; 67.8 (10) etorphine; 67.9 (11) heroin; 67.10 (12) hydromorphinol; 67.11 (13) methyldesorphine; 67.12 (14) methyldihydromorphine; 67.13 (15) morphine methylbromide; 67.14 67.15 (16) morphine methylsulfonate; (17) morphine-n-oxide; 67.16 (18) myrophine; 67.17 (19) nicocodeine; 67.18 (20) nicomorphine; 67.19 (21) normorphine; 67.20 (22) pholcodine; 67.21 (23) thebacon. 67.22 67.23 (d) Hallucinogens. Any material, compound, mixture or preparation which contains any quantity of the following substances, their analogs, salts, isomers (whether optical, 67.24 positional, or geometric), and salts of isomers, unless specifically excepted or unless listed 67.25 67.26 in another schedule, whenever the existence of the analogs, salts, isomers, and salts of isomers is possible: 67.27 (1) methylenedioxy amphetamine; 67.28 (2) methylenedioxymethamphetamine; 67.29 (3) methylenedioxy-N-ethylamphetamine (MDEA); 67.30 (4) n-hydroxy-methylenedioxyamphetamine; 67.31 (5) 4-bromo-2,5-dimethoxyamphetamine (DOB); 67.32 (6) 2,5-dimethoxyamphetamine (2,5-DMA); 67.33 (7) 4-methoxyamphetamine; 67.34 (8) 5-methoxy-3, 4-methylenedioxy amphetamine; 67.35

67.36

(9) alpha-ethyltryptamine;

```
(10) bufotenine;
68.1
             (11) diethyltryptamine;
68.2
             (12) dimethyltryptamine;
68.3
             (13) 3,4,5-trimethoxy amphetamine;
68.4
             (14) 4-methyl-2, 5-dimethoxyamphetamine (DOM);
68.5
             (15) ibogaine;
68.6
             (16) lysergic acid diethylamide (LSD);
68.7
             (17) mescaline;
68.8
             (18) parahexyl;
68.9
             (19) N-ethyl-3-piperidyl benzilate;
68.10
             (20) N-methyl-3-piperidyl benzilate;
68.11
             (21) psilocybin;
68.12
             (22) psilocyn;
68.13
             (23) tenocyclidine (TPCP or TCP);
68.14
68.15
             (24) N-ethyl-1-phenyl-cyclohexylamine (PCE);
             (25) 1-(1-phenylcyclohexyl) pyrrolidine (PCPy);
68.16
             (26) 1-[1-(2-thienyl)cyclohexyl]-pyrrolidine (TCPy);
68.17
             (27) 4-chloro-2,5-dimethoxyamphetamine (DOC);
68.18
             (28) 4-ethyl-2,5-dimethoxyamphetamine (DOET);
68.19
             (29) 4-iodo-2,5-dimethoxyamphetamine (DOI);
68.20
             (30) 4-bromo-2,5-dimethoxyphenethylamine (2C-B);
68.21
             (31) 4-chloro-2,5-dimethoxyphenethylamine (2C-C);
68.22
68.23
             (32) 4-methyl-2,5-dimethoxyphenethylamine (2-CD);
             (33) 4-ethyl-2,5-dimethoxyphenethylamine (2C-E);
68.24
             (34) 4-iodo-2,5-dimethoxyphenethylamine (2C-I);
68.25
68.26
             (35) 4-propyl-2,5-dimethoxyphenethylamine (2C-P);
             (36) 4-isopropylthio-2,5-dimethoxyphenethylamine (2C-T-4);
68.27
             (37) 4-propylthio-2,5-dimethoxyphenethylamine (2C-T-7);
68.28
             (38) 2-(8-bromo-2,3,6,7-tetrahydrofuro [2,3-f][1]benzofuran-4-yl)ethanamine
68.29
       (2-CB-FLY);
68.30
             (39) bromo-benzodifuranyl-isopropylamine (Bromo-DragonFLY);
68.31
             (40) alpha-methyltryptamine (AMT);
68.32
             (41) N,N-diisopropyltryptamine (DiPT);
68.33
             (42) 4-acetoxy-N,N-dimethyltryptamine (4-AcO-DMT);
68.34
             (43) 4-acetoxy-N,N-diethyltryptamine (4-AcO-DET);
68.35
             (44) 4-hydroxy-N-methyl-N-propyltryptamine (4-HO-MPT);
68.36
```

```
(45) 4-hydroxy-N,N-dipropyltryptamine (4-HO-DPT);
69.1
             (46) 4-hydroxy-N,N-diallyltryptamine (4-HO-DALT);
69.2
             (47) 4-hydroxy-N,N-diisopropyltryptamine (4-HO-DiPT);
69.3
             (48) 5-methoxy-N,N-diisopropyltryptamine (5-MeO-DiPT);
69.4
             (49) 5-methoxy-α-methyltryptamine (5-MeO-AMT);
69.5
             (50) 5-methoxy-N,N-dimethyltryptamine (5-MeO-DMT);
69.6
             (51) 5-methylthio-N,N-dimethyltryptamine (5-MeS-DMT);
69.7
             (52) 5-methoxy-N-methyl-N-propyltryptamine (5-MeO-MiPT);
69.8
             (53) 5-methoxy-α-ethyltryptamine (5-MeO-AET);
69.9
             (54) 5-methoxy-N,N-dipropyltryptamine (5-MeO-DPT);
69.10
             (55) 5-methoxy-N,N-diethyltryptamine (5-MeO-DET);
69.11
             (56) 5-methoxy-N,N-diallytryptamine (5-MeO-DALT);
69.12
             (57) methoxetamine (MXE);
69.13
             (58) 5-iodo-2-aminoindane (5-IAI);
69.14
69.15
             (59) 5,6-methylenedioxy-2-aminoindane (MDAI);
             (60) 2-(4-iodo-2,5-dimethoxyphenyl)-N-[(2-methoxyphenyl)methyl]ethanamine
69.16
       (25I-NBOMe).
69.17
             (e) Peyote. All parts of the plant presently classified botanically as Lophophora
69.18
       williamsii Lemaire, whether growing or not, the seeds thereof, any extract from any part
69.19
       of the plant, and every compound, manufacture, salts, derivative, mixture, or preparation
69.20
       of the plant, its seeds or extracts. The listing of peyote as a controlled substance in
69.21
       Schedule I does not apply to the nondrug use of peyote in bona fide religious ceremonies
69.22
69.23
        of the American Indian Church, and members of the American Indian Church are exempt
69.24
       from registration. Any person who manufactures peyote for or distributes peyote to the
       American Indian Church, however, is required to obtain federal registration annually and
69.25
69.26
       to comply with all other requirements of law.
             (f) Central nervous system depressants. Unless specifically excepted or unless listed
69.27
       in another schedule, any material compound, mixture, or preparation which contains any
69.28
       quantity of the following substances, their analogs, salts, isomers, and salts of isomers
69.29
       whenever the existence of the analogs, salts, isomers, and salts of isomers is possible:
69.30
             (1) mecloqualone;
69.31
             (2) methagualone;
69.32
             (3) gamma-hydroxybutyric acid (GHB), including its esters and ethers;
69.33
             (4) flunitrazepam.
69.34
             (g) Stimulants. Unless specifically excepted or unless listed in another schedule, any
69.35
       material compound, mixture, or preparation which contains any quantity of the following
69.36
```

substances, their analogs, salts, isomers, and salts of isomers whenever the existence of 70.1 70.2 the analogs, salts, isomers, and salts of isomers is possible: (1) aminorex; 70.3 (2) cathinone; 70.4 (3) fenethylline; 70.5 (4) methcathinone; 70.6 (5) methylaminorex; 70.7 (6) N,N-dimethylamphetamine; 70.8 (7) N-benzylpiperazine (BZP); 70.9 (8) methylmethcathinone (mephedrone); 70.10 (9) 3,4-methylenedioxy-N-methylcathinone (methylone); 70.11 (10) methoxymethcathinone (methedrone); 70.12 (11) methylenedioxypyrovalerone (MDPV); 70.13 (12) fluoromethcathinone; 70.14 70.15 (13) methylethcathinone (MEC); (14) 1-benzofuran-6-ylpropan-2-amine (6-APB); 70.16 (15) dimethylmethcathinone (DMMC); 70.17 (16) fluoroamphetamine; 70.18 (17) fluoromethamphetamine; 70.19 (18) α-methylaminobutyrophenone (MABP or buphedrone); 70.20 (19) β-keto-N-methylbenzodioxolylpropylamine (bk-MBDB or butylone); 70.21 (20) 2-(methylamino)-1-(4-methylphenyl)butan-1-one (4-MEMABP or BZ-6378); 70.22 70.23 (21) naphthylpyrovalerone (naphyrone); and (22) (RS)-1-phenyl-2-(1-pyrrolidinyl)-1-pentanone (alpha-PVP or 70.24 alpha-pyrrolidinovalerophenone; 70.25 70.26 (23) (RS)-1-(4-methylphenyl)-2-(1-pyrrolidinyl)-1-hexanone (4-Me-PHP oe MPHP); and 70.27 (22) (24) any other substance, except bupropion or compounds listed under a 70.28 different schedule, that is structurally derived from 2-aminopropan-1-one by substitution 70.29 at the 1-position with either phenyl, naphthyl, or thiophene ring systems, whether or not 70.30 the compound is further modified in any of the following ways: 70.31 (i) by substitution in the ring system to any extent with alkyl, alkylenedioxy, alkoxy, 70.32 70.33

- haloalkyl, hydroxyl, or halide substituents, whether or not further substituted in the ring system by one or more other univalent substituents; 70.34
- (ii) by substitution at the 3-position with an acyclic alkyl substituent; 70.35

- (iii) by substitution at the 2-amino nitrogen atom with alkyl, dialkyl, benzyl, or methoxybenzyl groups; or
 - (iv) by inclusion of the 2-amino nitrogen atom in a cyclic structure.
- (h) Marijuana, tetrahydrocannabinols, and synthetic cannabinoids. Unless specifically excepted or unless listed in another schedule, any natural or synthetic material, compound, mixture, or preparation that contains any quantity of the following substances, their analogs, isomers, esters, ethers, salts, and salts of isomers, esters, and ethers, whenever the existence of the isomers, esters, ethers, or salts is possible:
- (1) marijuana; 71.9

71.2

71.3

71.4

71.5

71.6

71.7

71.8

71.10

71.11

71.12

71.13

71.14

71.15

- (2) tetrahydrocannabinols naturally contained in a plant of the genus Cannabis, synthetic equivalents of the substances contained in the cannabis plant or in the resinous extractives of the plant, or synthetic substances with similar chemical structure and pharmacological activity to those substances contained in the plant or resinous extract, including, but not limited to, 1 cis or trans tetrahydrocannabinol, 6 cis or trans tetrahydrocannabinol, and 3,4 cis or trans tetrahydrocannabinol;
- (3) synthetic cannabinoids, including the following substances: 71.16
- (i) Naphthoylindoles, which are any compounds containing a 3-(1-napthoyl)indole 71.17 structure with substitution at the nitrogen atom of the indole ring by an alkyl, haloalkyl, 71.18 alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-piperidinyl)methyl or 71.19 2-(4-morpholinyl)ethyl group, whether or not further substituted in the indole ring to any 71.20 extent and whether or not substituted in the naphthyl ring to any extent. Examples of 71.21 naphthoylindoles include, but are not limited to: 71.22
- 71.23 (A) 1-Pentyl-3-(1-naphthoyl)indole (JWH-018 and AM-678);
- (B) 1-Butul-3-(1-naphthoyl)indole (JWH-073); 71.24
- (C) 1-Pentyl-3-(4-methoxy-1-naphthoyl)indole (JWH-081); 71.25
- 71.26 (D) 1-[2-(4-morpholinyl)ethyl]-3-(1-naphthoyl)indole (JWH-200);
- (E) 1-Propyl-2-methyl-3-(1-naphthoyl)indole (JWH-015); 71.27
- (F) 1-Hexyl-3-(1-naphthoyl)indole (JWH-019); 71.28
- (G) 1-Pentyl-3-(4-methyl-1-naphthoyl)indole (JWH-122); 71.29
- (H) 1-Pentyl-3-(4-ethyl-1-naphthoyl)indole (JWH-210); 71.30
- (I) 1-Pentyl-3-(4-chloro-1-naphthoyl)indole (JWH-398); 71.31
- (J) 1-(5-fluoropentyl)-3-(1-naphthoyl)indole (AM-2201). 71.32
- (ii) Napthylmethylindoles, which are any compounds containing a 71.33 1H-indol-3-yl-(1-naphthyl)methane structure with substitution at the nitrogen atom 71.34 of the indole ring by an alkyl, haloalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 71.35
- 1-(N-methyl-2-piperidinyl)methyl or 2-(4-morpholinyl)ethyl group, whether or not further 71.36

DM

substituted in the indole ring to any extent and whether or not substituted in the naphthyl 72.1 ring to any extent. Examples of naphthylmethylindoles include, but are not limited to: 72.2 (A) 1-Pentyl-1H-indol-3-yl-(1-naphthyl)methane (JWH-175); 72.3 (B) 1-Pentyl-1H-indol-3-yl-(4-methyl-1-naphthyl)methan (JWH-184). 72.4 (iii) Naphthoylpyrroles, which are any compounds containing a 72.5 3-(1-naphthoyl)pyrrole structure with substitution at the nitrogen atom of the 72.6 pyrrole ring by an alkyl, haloalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 72.7 1-(N-methyl-2-piperidinyl)methyl or 2-(4-morpholinyl)ethyl group whether or not 72.8 further substituted in the pyrrole ring to any extent, whether or not substituted in the 72.9 naphthyl ring to any extent. Examples of naphthoylpyrroles include, but are not limited to, 72.10 (5-(2-fluorophenyl)-1-pentylpyrrol-3-yl)-naphthalen-1-ylmethanone (JWH-307). 72.11 (iv) Naphthylmethylindenes, which are any compounds containing a 72.12 naphthylideneindene structure with substitution at the 3-position of the indene 72.13 ring by an allkyl, haloalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 72.14 72.15 1-(N-methyl-2-piperidinyl)methyl or 2-(4-morpholinyl)ethyl group whether or not further substituted in the indene ring to any extent, whether or not substituted in the naphthyl 72.16 ring to any extent. Examples of naphthylemethylindenes include, but are not limited to, 72.17 72.18 E-1-[1-(1-naphthalenylmethylene)-1H-inden-3-yl]pentane (JWH-176). (v) Phenylacetylindoles, which are any compounds containing a 3-phenylacetylindole 72.19 structure with substitution at the nitrogen atom of the indole ring by an alkyl, haloalkyl, 72.20 alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-piperidinyl)methyl or 72.21 2-(4-morpholinyl)ethyl group whether or not further substituted in the indole ring to 72.22 72.23 any extent, whether or not substituted in the phenyl ring to any extent. Examples of phenylacetylindoles include, but are not limited to: 72.24 (A) 1-(2-cyclohexylethyl)-3-(2-methoxyphenylacetyl)indole (RCS-8); 72.25 72.26 (B) 1-pentyl-3-(2-methoxyphenylacetyl)indole (JWH-250); (C) 1-pentyl-3-(2-methylphenylacetyl)indole (JWH-251); 72.27 (D) 1-pentyl-3-(2-chlorophenylacetyl)indole (JWH-203). 72.28 (vi) Cyclohexylphenols, which are compounds containing a 72.29 2-(3-hydroxycyclohexyl)phenol structure with substitution at the 5-position 72.30 of the phenolic ring by an alkyl, haloalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 72.31 1-(N-methyl-2-piperidinyl)methyl or 2-(4-morpholinyl)ethyl group whether or not 72.32 substituted in the cyclohexyl ring to any extent. Examples of cyclohexylphenols include, 72.33 but are not limited to: 72.34 (A) 5-(1,1-dimethylheptyl)-2-[(1R,3S)-3-hydroxycyclohexyl]-phenol (CP 47,497); 72.35

- 3rd Engrossment (B) 5-(1,1-dimethyloctyl)-2-[(1R,3S)-3-hydroxycyclohexyl]-phenol 73.1 (Cannabicyclohexanol or CP 47,497 C8 homologue); 73.2 (C) 5-(1,1-dimethylheptyl)-2-[(1R,2R)-5-hydroxy-2-(3-hydroxypropyl)cyclohexyl] 73.3 -phenol (CP 55,940). 73.4 (vii) Benzoylindoles, which are any compounds containing a 3-(benzoyl)indole 73.5 structure with substitution at the nitrogen atom of the indole ring by an alkyl, haloalkyl, 73.6 alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-piperidinyl)methyl or 73.7 2-(4-morpholinyl)ethyl group whether or not further substituted in the indole ring to 73.8 any extent and whether or not substituted in the phenyl ring to any extent. Examples of 73.9 benzoylindoles include, but are not limited to: 73.10 (A) 1-Pentyl-3-(4-methoxybenzoyl)indole (RCS-4); 73.11 (B) 1-(5-fluoropentyl)-3-(2-iodobenzoyl)indole (AM-694); 73.12 (C) (4-methoxyphenyl-[2-methyl-1-(2-(4-morpholinyl)ethyl)indol-3-yl]methanone 73.13 (WIN 48,098 or Pravadoline). 73.14 73.15 (viii) Others specifically named: (A) (6aR,10aR)-9-(hydroxymethyl)-6,6-dimethyl-3-(2-methyloctan-2-yl) 73.16 -6a,7,10,10a-tetrahydrobenzo[c]chromen-1-ol (HU-210); 73.17 (B) (6aS,10aS)-9-(hydroxymethyl)-6,6-dimethyl-3-(2-methyloctan-2-yl) 73.18 -6a,7,10,10a-tetrahydrobenzo[c]chromen-1-ol (Dexanabinol or HU-211); 73.19 (C) 2,3-dihydro-5-methyl-3-(4-morpholinylmethyl)pyrrolo[1,2,3-de] 73.20 -1,4-benzoxazin-6-yl-1-naphthalenylmethanone (WIN 55,212-2); 73.21 (D) (1-pentylindol-3-yl)-(2,2,3,3-tetramethylcyclopropyl)methanone (UR-144); 73.22 (E) (1-(5-fluoropentyl)-1H-indol-3-yl)(2,2,3,3-tetramethylcyclopropyl)methanone 73.23 (XLR-11); 73.24 (F) 1-pentyl-N-tricyclo[3.3.1.13,7]dec-1-yl-1H-indazole-3-carboxamide 73.25 73.26 (AKB-48(APINACA)); (G) N-((3s,5s,7s)-adamantan-1-yl)-1-(5-fluoropentyl)-1H-indazole-3-carboxamide 73.27 (5-Fluoro-AKB-48); 73.28 (H) 1-pentyl-8-quinolinyl ester-1H-indole-3-carboxylic acid (PB-22); 73.29 (I) 8-quinolinyl ester-1-(5-fluoropentyl)-1H-indole-3-carboxylic acid (5-Fluoro 73.30 PB-22)-; 73.31

1H-indazole-3-carboxamide (AB-FUBINACA).

3-carboxamide (AB-PINACA);

73.32

73.33

73.34

73.35

(K) N-[(1S)-1-(aminocarbonyl)-2-methylpropyl]-1-[(4-fluorophenyl)methyl]-

(J) N-[(1S)-1-(aminocarbonyl)-2-methylpropyl]-1-pentyl-1H-indazole-

DM

74.1	(i) A controlled substance analog, to the extent that it is implicitly or explicitly
74.2	intended for human consumption.
74.3	Sec. 21. Minnesota Statutes 2012, section 152.02, subdivision 8b, is amended to read:
74.4	Subd. 8b. Board of Pharmacy; expedited scheduling of additional substances.
74.5	(a) The state Board of Pharmacy may, by rule, add a substance to Schedule I provided that
74.6	it finds that the substance has a high potential for abuse, has no currently accepted medical
74.7	use in the United States, has a lack of accepted safety for use under medical supervision,
74.8	has known adverse health effects, and is currently available for use within the state. For
74.9	the purposes of this subdivision only, the board may use the expedited rulemaking process
74.10	under section 14.389. The scheduling of a substance under this subdivision expires the
74.11	day after the adjournment of the legislative session immediately following the substance's
74.12	scheduling unless the legislature by law ratifies the action.
74.13	(b) If the board schedules a substance under this subdivision, the board shall notify
74.14	in a timely manner the chairs and ranking minority members of the senate and house of
74.15	representatives committees having jurisdiction over criminal justice and health policy
74.16	and finance of the action and the reasons for it. The notice must include a copy of the
74.17	administrative law judge's decision on the matter.
74.18	(e) This subdivision expires August 1, 2014.
74.19	Sec. 22. Minnesota Statutes 2012, section 152.126, as amended by Laws 2013, chapter
74.20	113, article 3, section 3, is amended to read:
74.21	152.126 CONTROLLED SUBSTANCES PRESCRIPTION ELECTRONIC
74.22	REPORTING SYSTEM PRESCRIPTION MONITORING PROGRAM.
74.23	Subdivision 1. Definitions. (a) For purposes of this section, the terms defined in
74.24	this subdivision have the meanings given.
74.25	(a) (b) "Board" means the Minnesota State Board of Pharmacy established under
74.26	chapter 151.
74.27	(b) (c) "Controlled substances" means those substances listed in section 152.02,
74.28	subdivisions 3 to 5 6, and those substances defined by the board pursuant to section
74.29	152.02, subdivisions 7, 8, and 12. For the purposes of this section, controlled substances
74.30	includes tramadol and butalbital.
74.31	(e) (d) "Dispense" or "dispensing" has the meaning given in section 151.01,
74.32	subdivision 30. Dispensing does not include the direct administering of a controlled
74.33	substance to a patient by a licensed health care professional.

75.2

75.3

75.4

75.5

75.6

75.7

75.8

75.9

75.10

75.11

75.12

75.13

75.14

75.15

75.16

75.17

75.18

75.19

75.20

75.21

75.22

75.23

75.24

75.25

75.26

75.27

75.28

75.29

75.30

75.31

75.32

75.33

75.34

DM

(d) (e) "Dispenser" means a person authorized by law to dispense a controlled
substance, pursuant to a valid prescription. For the purposes of this section, a dispenser
does not include a licensed hospital pharmacy that distributes controlled substances for
inpatient hospital care, a licensed pharmacy, located on the same premises as a residentia
hospice, when the licensed pharmacy is dispensing controlled substances to be used
by an individual who is a resident of the hospice or a veterinarian who is dispensing
prescriptions under section 156.18.
(e) (f) "Prescriber" means a licensed health care professional who is authorized to
prescribe a controlled substance under section 152.12, subdivision 1 or 2.
(f) (g) "Prescription" has the meaning given in section 151.01, subdivision 16.
Subd. 1a. Treatment of intractable pain. This section is not intended to limit or
interfere with the legitimate prescribing of controlled substances for pain. No prescriber
shall be subject to disciplinary action by a health-related licensing board for prescribing a
controlled substance according to the provisions of section 152.125.
Subd. 2. Prescription electronic reporting system. (a) The board shall establish
by January 1, 2010, an electronic system for reporting the information required under
subdivision 4 for all controlled substances dispensed within the state.
(b) The board may contract with a vendor for the purpose of obtaining technical
assistance in the design, implementation, operation, and maintenance of the electronic
reporting system.
Subd. 3. Prescription Electronic Reporting Monitoring Program Advisory
Committee Task Force. (a) The board shall convene may appoint an advisory committee
The committee must include task force consisting of at least one representative of:
(1) the Department of Health;
(2) the Department of Human Services;
(3) each health-related licensing board that licenses prescribers;
(4) a professional medical association, which may include an association of pain
management and chemical dependency specialists;
(5) a professional pharmacy association;
(6) a professional nursing association;
(7) a professional dental association;
(8) a consumer privacy or security advocate; and
(9) a consumer or patient rights organization: and
(10) an association of medical examiners and coroners.

DM

6.1	(b) The advisory eommittee task force shall advise the board on the development and
76.2	operation of the electronic reporting system prescription monitoring program, including,
76.3	but not limited to:
6.4	(1) technical standards for electronic prescription drug reporting;
6.5	(2) proper analysis and interpretation of prescription monitoring data; and
6.6	(3) an evaluation process for the program; and
6.7	(4) criteria for the unsolicited provision of prescription monitoring data by the
6.8	board to prescribers and dispensers.
6.9	(c) The task force is governed by section 15.059. Notwithstanding section 15.059,
6.10	subdivision 5, the task force shall not expire.
6.11	Subd. 4. Reporting requirements; notice. (a) Each dispenser must submit the
6.12	following data to the board or its designated vendor, subject to the notice required under
76.13	paragraph (d) :
76.14	(1) name of the prescriber;
6.15	(2) national provider identifier of the prescriber;
76.16	(3) name of the dispenser;
6.17	(4) national provider identifier of the dispenser;
76.18	(5) prescription number;
6.19	(6) name of the patient for whom the prescription was written;
6.20	(7) address of the patient for whom the prescription was written;
6.21	(8) date of birth of the patient for whom the prescription was written;
6.22	(9) date the prescription was written;
76.23	(10) date the prescription was filled;
6.24	(11) name and strength of the controlled substance;
6.25	(12) quantity of controlled substance prescribed;
76.26	(13) quantity of controlled substance dispensed; and
76.27	(14) number of days supply.
76.28	(b) The dispenser must submit the required information by a procedure and in a
76.29	format established by the board. The board may allow dispensers to omit data listed in this
76.30	subdivision or may require the submission of data not listed in this subdivision provided
76.31	the omission or submission is necessary for the purpose of complying with the electronic
76.32	reporting or data transmission standards of the American Society for Automation in
76.33	Pharmacy, the National Council on Prescription Drug Programs, or other relevant national
76.34	standard-setting body.
6.35	(c) A dispenser is not required to submit this data for those controlled substance

prescriptions dispensed for:

76.35

	SF1484	REVISOR	DM	S1484-3	3rd Engrossment
77.1	(1) inc	lividuals residing in l i	eensed skilled	l nursing or intermedia	ate care facilities;
77.2	(2) inc	dividuals receiving as	sisted living s	ervices under chapter	144G or through a
77.3	medical ass	istance home and con	munity-based	l waiver;	
77.4	(3) inc	dividuals receiving me	edication intra	venously;	
77.5	(4) inc	lividuals receiving ho	spice and other	er palliative or end-of-	life care; and
77.6	(5) inc	lividuals receiving sea	rvices from a	nome care provider reg	gulated under chapter
77.7	144A.				
77.8	(1) inc	dividuals residing in a	health care f	acility as defined in se	ection 151.58,
77.9	subdivision	2, paragraph (b), whe	en a drug is di	stributed through the u	use of an automated
77.10	drug distrib	ution system accordin	g to section 1	51.58; and	
77.11	(2) inc	dividuals receiving a c	drug sample th	nat was packaged by a	manufacturer and
77.12	provided to	the dispenser for disp	ensing as a p	rofessional sample pur	rsuant to Code of
77.13	Federal Reg	gulations, title 21, sect	tion 203, subp	eart D.	
77.14	(d) A	dispenser must not su	bmit data und	er this subdivision un	less provide to the
77.15	patient for v	whom the prescription	was written	a conspicuous notice of	of the reporting
77.16	requirement	s of this section is giv	en to the pati	ent for whom the pres	eription was written
77.17	and notice	that the information m	nay be used for	r program administrat	ion purposes.
77.18	Subd.	5. Use of data by bo	ard. (a) The b	ooard shall develop and	d maintain a database
77.19	of the data r	eported under subdivi	ision 4. The b	oard shall maintain da	ta that could identify
77.20	an individua	al prescriber or disper	nser in encryp	ted form. Except as of	therwise allowed
77.21	under subdi	vision 6, the database	may be used	by permissible users	identified under
77.22	subdivision	6 for the identificatio	n of:		
77.23	(1) inc	lividuals receiving pro	escriptions for	r controlled substances	s from prescribers
77.24	who subseq	uently obtain controll	ed substances	from dispensers in qu	nantities or with a
77.25	frequency in	nconsistent with gener	rally recogniz	ed standards of use for	r those controlled
77.26	substances,	including standards a	ccepted by na	tional and internationa	al pain management
77.27	associations	; and			
77.28	(2) inc	lividuals presenting for	orged or other	wise false or altered p	prescriptions for
77.29	controlled s	ubstances to dispense	rs.		

- (b) No permissible user identified under subdivision 6 may access the database for the sole purpose of identifying prescribers of controlled substances for unusual or excessive prescribing patterns without a valid search warrant or court order.
- (c) No personnel of a state or federal occupational licensing board or agency may access the database for the purpose of obtaining information to be used to initiate or substantiate a disciplinary action against a prescriber when the disciplinary action relates

77.31

77.32

77.33

77.34

78.2

78.3

78.4

78.5

78.6

78.7

78.8

78.9

78.10

78.11

78.12

78.13

78.14

78.15

78.16

78.17

78.18

78.19

78.20

78.21

78.22

78.23

78.24

78.25

78.26

78.27

78.28

78.29

78.30

78.31

78.32

78.33

78.34

78.35

78.36

to allegations involving unusual or excessive prescribing of the drugs for which data is collected under subdivision 4.

DM

- (d) Data reported under subdivision 4 shall be retained by the board in the database for a 12-month period, and shall be removed from the database no later than 12 months from the last day of the month during which the data was received. made available to permissible users for a 12-month period beginning the day the data was received and ending 12 months from the last day of the month in which the data was received, except that permissible users defined in subdivision 6, paragraph (b), clauses (6) and (7), may use all data collected under this section for the purposes of administering, operating, and maintaining the prescription monitoring program and conducting trend analyses and other studies necessary to evaluate the effectiveness of the program.
- (e) The board shall not retain data reported under subdivision 4 for a period longer than five years from the date the data was received.
- Subd. 6. Access to reporting system data. (a) Except as indicated in this subdivision, the data submitted to the board under subdivision 4 is private data on individuals as defined in section 13.02, subdivision 12, and not subject to public disclosure.
- (b) Except as specified in subdivision 5, the following persons shall be considered permissible users and may access the data submitted under subdivision 4 in the same or similar manner, and for the same or similar purposes, as those persons who are authorized to access similar private data on individuals under federal and state law:
- (1) a prescriber or an agent or employee of the prescriber to whom the prescriber has delegated the task of accessing the data, to the extent the information relates specifically to a current patient, to whom the prescriber is prescribing or considering prescribing any controlled substance or to whom the prescriber is providing other medical treatment for which access to the data may be necessary and with the provision that the prescriber remains responsible for the use or misuse of data accessed by a delegated agent or employee;
- (2) a dispenser or an agent or employee of the dispenser to whom the dispenser has delegated the task of accessing the data, to the extent the information relates specifically to a current patient to whom that dispenser is dispensing or considering dispensing any controlled substance and with the provision that the dispenser remains responsible for the use or misuse of data accessed by a delegated agent or employee;
- (3) a licensed pharmacist who is providing pharmaceutical care for which access to the data may be necessary to the extent that the information relates specifically to a current patient for whom the pharmacist is providing pharmaceutical care;
- (3) (4) an individual who is the recipient of a controlled substance prescription for which data was submitted under subdivision 4, or a guardian of the individual, parent or

DM guardian of a minor, or health care agent of the individual acting under a health care 79.1 79.2 directive under chapter 145C; (4) (5) personnel of the a health-related licensing board specifically listed in section 79.3 214.01, subdivision 2, or the Emergency Medical Services Regulatory Board, assigned to 79.4 conduct a bona fide investigation of a complaint received by that board alleging that a 79.5 specific licensee is impaired by use of a drug for which data is collected under subdivision 79.6 4, has engaged in activity that would constitute a crime as defined in section 152.025, or 79.7 has engaged in the behavior specified in section 152.126, subdivision 5, paragraph (a); 79.8 (5) (6) personnel of the board engaged in the collection, review, and analysis 79.9 of controlled substance prescription information as part of the assigned duties and 79.10 responsibilities under this section; 79.11 (6) (7) authorized personnel of a vendor under contract with the board state of 79.12 Minnesota who are engaged in the design, implementation, operation, and maintenance of 79.13 the electronic reporting system prescription monitoring program as part of the assigned 79.14 79.15 duties and responsibilities of their employment, provided that access to data is limited to the minimum amount necessary to carry out such duties and responsibilities; 79.16 (7) (8) federal, state, and local law enforcement authorities acting pursuant to a 79.17 valid search warrant; 79.18 (8) (9) personnel of the medical assistance program Minnesota health care programs 79.19 assigned to use the data collected under this section to identify and manage recipients 79.20 whose usage of controlled substances may warrant restriction to a single primary care 79.21 physician provider, a single outpatient pharmacy, or and a single hospital; and 79.22 79.23 (9) (10) personnel of the Department of Human Services assigned to access the data pursuant to paragraph (h)-; 79.24 79.25

(11) a coroner or medical examiner, or an agent or employee of the coroner or medical examiner to whom the coroner or medical examiner has delegated the task of accessing the data, conducting an investigation pursuant to section 390.11, and with the provision that the coroner or medical examiner remains responsible for the use or misuse of data accessed by a delegated agent or employee; and

(12) personnel of the health professionals services program established under section 214.31, to the extent that the information relates specifically to an individual who is currently enrolled in and being monitored by the program. The health professionals services program personnel shall not provide this data to a health-related licensing board or the Emergency Medical Services Regulatory Board, except as permitted under section 214.33, subdivision 3.

79.26

79.27

79.28

79.29

79.30

79.31

79.32

79.33

79.34

80.2

80.3

80.4

80.5

80.6

80.7

80.8

80.9

80.10

80.11

80.12

80.13

80.14

80.15

80.16

80.17

80.18

80.19

80.20

80.21

80.22

80.23

80.24

80.25

80.26

80.27

80.28

80.29

80.30

80.31

80.32

80.33

80.34

80.35

80.36

For purposes of clause (3) (4), access by an individual includes persons in the definition of an individual under section 13.02.

- (c) Any A permissible user identified in paragraph (b), who clauses (1), (2), (3), (6), (7), (9), (10), and (11) may directly accesses access the data electronically. If the data is directly accessed electronically, the permissible user shall implement and maintain a comprehensive information security program that contains administrative, technical, and physical safeguards that are appropriate to the user's size and complexity, and the sensitivity of the personal information obtained. The permissible user shall identify reasonably foreseeable internal and external risks to the security, confidentiality, and integrity of personal information that could result in the unauthorized disclosure, misuse, or other compromise of the information and assess the sufficiency of any safeguards in place to control the risks.
- (d) The board shall not release data submitted under this section subdivision 4 unless it is provided with evidence, satisfactory to the board, that the person requesting the information is entitled to receive the data.
- (e) The board shall not release the name of a prescriber without the written consent of the prescriber or a valid search warrant or court order. The board shall provide a mechanism for a prescriber to submit to the board a signed consent authorizing the release of the prescriber's name when data containing the prescriber's name is requested.
- (f) (e) The board shall maintain a log of all persons who access the data for a period of at least three years and shall ensure that any permissible user complies with paragraph (c) prior to attaining direct access to the data.
- (g) (f) Section 13.05, subdivision 6, shall apply to any contract the board enters into pursuant to subdivision 2. A vendor shall not use data collected under this section for any purpose not specified in this section.
- (g) The board may participate in an interstate prescription monitoring program data exchange system provided that permissible users in other states have access to the data only as allowed under this section, and that section 13.05, subdivision 6, applies to any contract or memorandum of understanding that the board enters into under this paragraph.
- (h) With available appropriations, the commissioner of human services shall establish and implement a system through which the Department of Human Services shall routinely access the data for the purpose of determining whether any client enrolled in an opioid treatment program licensed according to chapter 245A has been prescribed or dispensed a controlled substance in addition to that administered or dispensed by the opioid treatment program. When the commissioner determines there have been multiple prescribers or multiple prescriptions of controlled substances, the commissioner shall:

DM

81.1	(1) inform the medical director of the opioid treatment program only that the
81.2	commissioner determined the existence of multiple prescribers or multiple prescriptions of
81.3	controlled substances; and
81.4	(2) direct the medical director of the opioid treatment program to access the data
81.5	directly, review the effect of the multiple prescribers or multiple prescriptions, and
81.6	document the review.
81.7	If determined necessary, the commissioner of human services shall seek a federal waiver
81.8	of, or exception to, any applicable provision of Code of Federal Regulations, title 42, part
81.9	2.34, item (c), prior to implementing this paragraph.
81.10	(i) The board may provide data submitted under subdivision 4 for public research,
81.11	policy, or education purposes, but only after the removal of any information that is likely
81.12	to reveal the identity of the patient, prescriber, or dispenser who is the subject of the data.
81.13	(j) The board shall review the data submitted under subdivision 4 on at least a
81.14	quarterly basis and shall establish criteria, in consultation with the advisory task force,
81.15	for referring information about a patient to prescribers and dispensers who prescribed or
81.16	dispensed the prescriptions in question if the criteria are met.
81.17	Subd. 7. Disciplinary action. (a) A dispenser who knowingly fails to submit data to
81.18	the board as required under this section is subject to disciplinary action by the appropriate
81.19	health-related licensing board.
81.20	(b) A prescriber or dispenser authorized to access the data who knowingly discloses
81.21	the data in violation of state or federal laws relating to the privacy of health care data
81.22	shall be subject to disciplinary action by the appropriate health-related licensing board,
81.23	and appropriate civil penalties.
81.24	Subd. 8. Evaluation and reporting. (a) The board shall evaluate the prescription
81.25	electronic reporting system to determine if the system is negatively impacting appropriate
81.26	prescribing practices of controlled substances. The board may contract with a vendor to
81.27	design and conduct the evaluation.
81.28	(b) The board shall submit the evaluation of the system to the legislature by July
81.29	15, 2011.
81.30	Subd. 9. Immunity from liability; no requirement to obtain information. (a) A
81.31	pharmacist, prescriber, or other dispenser making a report to the program in good faith
81.32	under this section is immune from any civil, criminal, or administrative liability, which
81 33	might otherwise be incurred or imposed as a result of the report, or on the basis that the

81.34

81.35

81.36

pharmacist or prescriber did or did not seek or obtain or use information from the program.

to obtain information about a patient from the program, and the pharmacist, prescriber,

(b) Nothing in this section shall require a pharmacist, prescriber, or other dispenser

82.2

82.3

82.4

82.5

82.6

82.7

82.8

82.9

82.10

82.11

82.12

82.13

82.14

82.15

82.16

82.17

82.18

82.19

82.20

82.21

82.22

82.23

82.24

82.25

82.26

82.27

82.28

82.29

82.30

82.31

82.32

82.33

82.34

82.35

or other dispenser, if acting in good faith, is immune from any civil, criminal, or administrative liability that might otherwise be incurred or imposed for requesting, receiving, or using information from the program.

DM

Subd. 10. **Funding.** (a) The board may seek grants and private funds from nonprofit charitable foundations, the federal government, and other sources to fund the enhancement and ongoing operations of the prescription electronic reporting system monitoring program established under this section. Any funds received shall be appropriated to the board for this purpose. The board may not expend funds to enhance the program in a way that conflicts with this section without seeking approval from the legislature.

(b) Notwithstanding any other section, the administrative services unit for the health-related licensing boards shall apportion between the Board of Medical Practice, the Board of Nursing, the Board of Dentistry, the Board of Podiatric Medicine, the Board of Optometry, the Board of Veterinary Medicine, and the Board of Pharmacy an amount to be paid through fees by each respective board. The amount apportioned to each board shall equal each board's share of the annual appropriation to the Board of Pharmacy from the state government special revenue fund for operating the prescription electronic reporting system monitoring program under this section. Each board's apportioned share shall be based on the number of prescribers or dispensers that each board identified in this paragraph licenses as a percentage of the total number of prescribers and dispensers licensed collectively by these boards. Each respective board may adjust the fees that the boards are required to collect to compensate for the amount apportioned to each board by the administrative services unit.

EFFECTIVE DATE. This section is effective the day following final enactment.

Sec. 23. STUDY REQUIRED; PRESCRIPTION MONITORING PROGRAM DATABASE.

The Board of Pharmacy, in collaboration with the Prescription Monitoring Program Advisory Task Force, shall study program database and report to the chairs and ranking minority members of the senate health and human services policy and finance division and the house of representatives health and human services policy and finance committees by December 15, 2014, with recommendations on whether or not to (1) require the use of the prescription monitoring by prescribers when prescribing or considering prescribing, and pharmacists when dispensing or considering dispensing, a controlled substance as defined in Minnesota Statutes, section 152.126, subdivision 1, paragraph (c); and (2) allow for the use of the prescription monitoring program database to identify potentially inappropriate prescribing of controlled substances.

Sec. 24. APPROPRIATION.
(a) \$210,000 in fiscal year 2015 is appropriated from the state government special
revenue fund to the Board of Pharmacy to implement changes to the prescription monitoring
program. The base for this appropriation is \$171,000 in fiscal years 2016 and 2017.
(b) \$5,000 in fiscal year 2015 is appropriated from the state government special

S1484-3

3rd Engrossment

DM

revenue fund to the Board of Pharmacy for costs attributable to the board's cease and desist authority.

REVISOR

SF1484

83.1

83.2

83.3

83.4

APPENDIX Article locations in S1484-3

ARTICLE 1	HEALTH-RELATED LICENSING BOARDS	Page.Ln 1.27
ARTICLE 2	BOARD OF PHARMACY	Page.Ln 24.1