

**SUBSTITUTE FOR
HOUSE BILL NO. 4217**

A bill to amend 1978 PA 368, entitled
"Public health code,"
by amending sections 7333, 16221, 16221b, 16226, 17744, 17751, and
17754 (MCL 333.7333, 333.16221, 333.16221b, 333.16226, 333.17744,
333.17751, and 333.17754), section 7333 as amended by 2018 PA 34,
sections 16221 and 16226 as amended by 2018 PA 463, section 16221b
as added by 2017 PA 249, section 17744 as added by 2012 PA 209,
section 17751 as amended by 2017 PA 165, and section 17754 as
amended by 2014 PA 525, and by adding section 17754a.

THE PEOPLE OF THE STATE OF MICHIGAN ENACT:

1 Sec. 7333. (1) As used in this section, "good faith" means the
2 prescribing or dispensing of a controlled substance by a
3 practitioner licensed under section 7303 in the regular course of



professional treatment to or for an individual who is under treatment by the practitioner for a pathology or condition other than that individual's physical or psychological dependence upon or addiction to a controlled substance, except as provided in this article. Application of good faith to a pharmacist means the dispensing of a controlled substance pursuant to a prescriber's order which, in the professional judgment of the pharmacist, is lawful. The pharmacist shall be guided by nationally accepted professional standards including, but not limited to, all of the following, in making the judgment:

(a) Lack of consistency in the doctor-patient relationship.

(b) Frequency of prescriptions for the same drug by 1 prescriber for larger numbers of patients.

(c) Quantities beyond those normally prescribed for the same drug.

(d) Unusual dosages.

(e) Unusual geographic distances between patient, pharmacist, and prescriber.

(2) Except as otherwise provided in this section, a practitioner, in good faith, may dispense a controlled substance included in schedule 2 **that is a prescription drug as determined under section 503(b) of the federal food, drug, and cosmetic act, 21 USC 353, or section 17708**, upon receipt of ~~a~~**either of the following:**

(a) **A** prescription of a practitioner licensed under section 7303 on a prescription form. ~~A practitioner may issue more~~ **More** than 1 prescription for a controlled substance included in schedule 2 **may be included** on a single prescription form.

(b) **A prescription that is electronically transmitted under**



section 17754a.

(3) In an emergency situation, as described in R 338.3165 of the Michigan Administrative Code, a controlled substance included in schedule 2 may be dispensed upon the oral prescription of a practitioner if the prescribing practitioner promptly fills out a prescription form and forwards the prescription form to the dispensing pharmacy within 7 days after the oral prescription is issued. A prescription for a controlled substance included in schedule 2 must not be filled more than 90 days after the date on which the prescription was issued. A pharmacist, consistent with federal law and regulations on the partial filling of a controlled substance included in schedule 2, may partially fill in increments a prescription for a controlled substance included in schedule 2.

(4) A practitioner, in good faith, may dispense a controlled substance included in schedule 3, 4, or 5 that is a prescription drug as determined under section 503(b) of the federal food, drug, and cosmetic act, 21 USC 353, or section 17708, upon receipt of ~~a~~ **any of the following:**

(a) A prescription on a prescription form. ~~or an~~

(b) An oral prescription of a practitioner.

(c) **A prescription that is electronically transmitted under section 17754a.**

(5) A prescription for a controlled substance included in schedule 3 or 4 must not be filled or refilled without specific refill instructions noted by the prescriber. A prescription for a controlled substance included in schedule 3 or 4 must not be filled or refilled later than 6 months after the date of the prescription or be refilled more than 5 times, unless renewed by the prescriber in accordance with rules promulgated by the administrator.



1 (6) ~~(5)~~—A controlled substance included in schedule 5 must not
2 be distributed or dispensed other than for a medical purpose, or in
3 any manner except in accordance with rules promulgated by the
4 administrator.

5 (7) ~~(6)~~—If a prescription is required under this section, the
6 prescription must contain the quantity of the controlled substance
7 prescribed in both written and numerical terms. A prescription is
8 in compliance with this subsection if, in addition to containing
9 the quantity of the controlled substance prescribed in written
10 terms, it contains preprinted numbers representative of the
11 quantity of the controlled substance prescribed next to which is a
12 box or line the prescriber may check.

13 (8) ~~(7)~~—A prescribing practitioner shall not use a
14 prescription form for a purpose other than prescribing. A
15 prescribing practitioner shall not postdate a prescription form
16 that contains a prescription for a controlled substance. ~~A—Until~~
17 **the date on which section 17754a applies, a** prescriber may transmit
18 a prescription by facsimile of a printed prescription form and by
19 electronic transmission of a printed prescription form, if not
20 prohibited by federal law. If, with the patient's consent, a
21 prescription is electronically transmitted **under this subsection,**
22 it must be transmitted directly to a pharmacy of the patient's
23 choice by the prescriber or the prescriber's authorized agent, and
24 the data must not be altered, modified, or extracted in the
25 transmission process.

26 (9) ~~(8)~~—Notwithstanding subsections (1) to ~~(5)~~, ~~(6)~~, a class B
27 dealer may acquire a limited permit only for the purpose of buying,
28 possessing, and administering a commercially prepared, premixed
29 solution of sodium pentobarbital to perform euthanasia on injured,



1 sick, homeless, or unwanted domestic pets and other animals, if the
2 class B dealer does all of the following:

3 (a) Applies to the administrator for a permit in accordance
4 with rules promulgated under this part. The application must
5 contain the name of the individual in charge of the day-to-day
6 operations of the class B dealer's facilities and the name of the
7 individual responsible for designating employees who will be
8 performing euthanasia on animals pursuant to this act.

9 (b) Complies with the rules promulgated by the administrator
10 for the storage, handling, and use of a commercially prepared,
11 premixed solution of sodium pentobarbital to perform euthanasia on
12 animals. The class B dealer shall maintain a record of use and
13 shall make the record available for inspection by the department of
14 licensing and regulatory affairs, the department of agriculture and
15 rural development, and the United States Department of Agriculture.

16 (c) Subject to subdivision (d), certifies that the class B
17 dealer or an employee of the class B dealer has received, and can
18 document completion of, a minimum of 16 hours of training,
19 including at least 12 hours of content training and at least 4
20 hours of practical training, in the use of a commercially prepared,
21 premixed solution of sodium pentobarbital and an animal
22 tranquilizer to perform euthanasia on animals from a training
23 program approved by the state veterinarian, in consultation with
24 the Michigan board of veterinary medicine, and given by a licensed
25 veterinarian pursuant to rules promulgated by the administrator.
26 The training described in this subdivision ~~shall~~**must** comply with
27 the American Veterinary Medical Association's guidelines for the
28 euthanasia of animals.

29 (d) Until December 31, 2021, ensures that the class B dealer



1 or an employee of the class B dealer who received, and can document
 2 the completion of, the 8 hours of training required immediately
 3 before ~~the effective date of the 2018 amendatory act that amended~~
 4 ~~this section~~ **May 22, 2018** only administers a commercially prepared,
 5 premixed solution of sodium pentobarbital to perform euthanasia on
 6 the animals described in this subsection. Beginning January 1,
 7 2022, the individuals described in this subdivision must have
 8 received, and be able to document the completion of, the training
 9 described in subdivision (c) to administer a commercially prepared,
 10 premixed solution of sodium pentobarbital or an animal tranquilizer
 11 to perform euthanasia on the animals described in this subsection.

12 (e) Certifies that only an individual described in subdivision
 13 (c) or (d) or an individual otherwise permitted to use a controlled
 14 substance pursuant to this article will administer the commercially
 15 prepared, premixed solution of sodium pentobarbital or an animal
 16 tranquilizer according to written procedures established by the
 17 class B dealer.

18 (f) Beginning January 1, 2022, certifies that the individual
 19 in charge of the day-to-day operations of the class B dealer's
 20 facilities has received, and can document the completion of, the
 21 training described in subdivision (c).

22 (g) Complies with all state and federal laws, rules, and
 23 regulations regarding the acquisition, use, and security of
 24 controlled substances.

25 **(10)** ~~(9)~~ Notwithstanding subsections (1) to ~~(5)~~, **(6)**, an
 26 animal control shelter or animal protection shelter registered with
 27 the department of agriculture and rural development pursuant to
 28 1969 PA 287, MCL 287.331 to 287.340, may acquire a limited permit
 29 only for the purpose of buying, possessing, and administering a



1 commercially prepared, premixed solution of sodium pentobarbital,
2 or an animal tranquilizer, to use exclusively as an adjunct in the
3 process of performing euthanasia on injured, sick, homeless, or
4 unwanted domestic pets and other animals, if the animal control
5 shelter or animal protection shelter does all of the following:

6 (a) Applies to the administrator for a permit in accordance
7 with rules promulgated under this part. The application must
8 contain the name of the individual in charge of the day-to-day
9 operations of the animal control shelter or animal protection
10 shelter and the name of the individual responsible for designating
11 employees who will be performing euthanasia on animals pursuant to
12 this act.

13 (b) Complies with the rules promulgated by the administrator
14 for the storage, handling, and use of a commercially prepared,
15 premixed solution of sodium pentobarbital or an animal tranquilizer
16 to perform euthanasia on animals. The animal control shelter or
17 animal protection shelter shall maintain a record of use and make
18 the record available for inspection by the department of licensing
19 and regulatory affairs and the department of agriculture and rural
20 development.

21 (c) Subject to subdivision (d), certifies that an employee of
22 the animal control shelter or animal protection shelter has
23 received, and can document completion of, a minimum of 16 hours of
24 training, including at least 12 hours of content training and at
25 least 4 hours of practical training, in the use of a commercially
26 prepared, premixed solution of sodium pentobarbital and an animal
27 tranquilizer to perform euthanasia on animals from a training
28 program approved by the state veterinarian, in consultation with
29 the Michigan board of veterinary medicine, and given by a licensed



1 veterinarian pursuant to rules promulgated by the administrator.
2 The training described in this subdivision must comply with the
3 American Veterinary Medical Association's guidelines for the
4 euthanasia of animals.

5 (d) Until December 31, 2021, ensures that an employee of the
6 animal control shelter or animal protection shelter who received,
7 and can document the completion of, the training required
8 immediately before ~~the effective date of the 2018 amendatory act~~
9 ~~that amended this section~~ **May 22, 2018** only administers a
10 commercially prepared solution of xylazine hydrochloride or a
11 commercially prepared, premixed solution of sodium pentobarbital to
12 perform euthanasia on the animals described in this subsection in
13 accordance with his or her training. Beginning January 1, 2022, the
14 employee described in this subdivision must have received, and be
15 able to document the completion of, the training described in
16 subdivision (c) to administer a commercially prepared, premixed
17 solution of sodium pentobarbital or an animal tranquilizer to
18 perform euthanasia on the animals described in this subsection.

19 (e) Certifies that only an individual described in subdivision
20 (c) or (d) or an individual otherwise permitted to use a controlled
21 substance pursuant to this article will administer a commercially
22 prepared, premixed solution of sodium pentobarbital or an animal
23 tranquilizer according to written procedures established by the
24 animal control shelter or animal protection shelter.

25 (f) Beginning January 1, 2022, certifies that the individual
26 in charge of the day-to-day operations of the animal control
27 shelter or animal protection shelter has received, and can document
28 the completion of, the training described in subdivision (c).

29 (g) Complies with all state and federal laws and regulations



1 regarding the acquisition, use, and security of controlled
2 substances.

3 **(11)** ~~(10)~~ The application described in subsection ~~(8) or (9)~~
4 **or (10)** must include the names and addresses of all individuals
5 employed by the animal control shelter or animal protection shelter
6 or class B dealer who have been trained as described in subsection
7 ~~(8)(e), (9)(c), (d), and (f) or (9)(e), (10)(c), (d), and (f)~~ and
8 the name of the veterinarian who trained them. The list of names
9 and addresses must be updated every 6 months.

10 **(12)** ~~(11)~~ If an animal control shelter or animal protection
11 shelter or class B dealer issued a permit pursuant to subsection
12 ~~(8) or (9)~~ **or (10)** does not have in its employ an individual
13 trained as described in subsection ~~(8)(e) (9)(c) or (d) and (8)(f),~~
14 **(9)(f), or (9)(e) (10)(c) or (d) and (9)(f), (10)(f),** the animal
15 control shelter or animal protection shelter or class B dealer
16 shall immediately notify the administrator and shall cease to
17 administer a commercially prepared, premixed solution of sodium
18 pentobarbital or an animal tranquilizer for the purposes described
19 in subsection ~~(8) or (9)~~ **or (10)** until the administrator is
20 notified that 1 of the following has occurred:

21 (a) An individual trained as described in subsection ~~(8)(e),~~
22 **(9)(c), (d), or (f) or (9)(e), (10)(c), (d), or (f)** has been hired
23 by the animal control shelter or animal protection shelter or class
24 B dealer.

25 (b) An individual employed by the animal control shelter or
26 animal protection shelter or class B dealer has been trained as
27 described in subsection ~~(8)(e) (9)(c) or (f) or (9)(e) (10)(c) or~~
28 ~~(f).~~

29 **(13)** ~~(12)~~ A veterinarian, including a veterinarian who trains



1 individuals as described in subsection ~~(8)(e)~~, **(9)(c)**, (d), or (f),
 2 or ~~(9)(e)~~, **(10)(c)**, (d), or (f), is not civilly or criminally
 3 liable for the use of a commercially prepared, premixed solution of
 4 sodium pentobarbital or an animal tranquilizer by an animal control
 5 shelter or animal protection shelter or a class B dealer, unless
 6 the veterinarian is employed by or under contract with the animal
 7 control shelter or animal protection shelter or class B dealer and
 8 the terms of the veterinarian's employment or the contract require
 9 the veterinarian to be responsible for the use or administration of
 10 the commercially prepared, premixed solution of sodium
 11 pentobarbital or animal tranquilizer.

12 **(14)** ~~(13)~~—A person shall not knowingly use or permit the use
 13 of a commercially prepared, premixed solution of sodium
 14 pentobarbital or an animal tranquilizer in violation of this
 15 section.

16 **(15)** ~~(14)~~—This section does not require that a veterinarian be
 17 employed by or under contract with an animal control shelter or
 18 animal protection shelter or class B dealer to obtain, possess, or
 19 administer a commercially prepared, premixed solution of sodium
 20 pentobarbital or an animal tranquilizer pursuant to this section.

21 **(16)** ~~(15)~~—Notwithstanding subsections (1) to ~~(5)~~, **(6)**, an
 22 animal control shelter registered with the department of
 23 agriculture and rural development pursuant to 1969 PA 287, MCL
 24 287.331 to 287.340, may acquire a limited permit only for the
 25 purpose of buying, possessing, and administering an animal
 26 tranquilizer to sedate or immobilize an animal running at large
 27 that is dangerous or difficult to capture, if the animal control
 28 shelter does all of the following:

29 (a) Applies to the administrator for a permit in accordance



1 with the rules promulgated under this part. The application must
2 contain the name of the individual in charge of the day-to-day
3 operations of the animal control shelter and the name of the
4 individual responsible for designating employees who will be
5 administering an animal tranquilizer pursuant to this act.

6 (b) Complies with the rules promulgated by the administrator
7 for the storage, handling, and use of an animal tranquilizer. The
8 animal control shelter shall maintain a record of use and shall
9 make the record available for inspection by the department of
10 licensing and regulatory affairs and the department of agriculture
11 and rural development.

12 (c) Subject to subdivision (d), certifies that an employee of
13 the animal control shelter has received, and can document
14 completion of, both of the following in the following order:

15 (i) The training described in subsection ~~(9)(e)~~ **(10)(c)**.

16 (ii) A minimum of 16 hours of training, including at least 12
17 hours of content training and at least 4 hours of practical
18 training, in the use of animal tranquilizers to sedate or
19 immobilize the animals described in this subsection from a training
20 program approved by the state veterinarian, in consultation with
21 the Michigan board of veterinary medicine, and given by a licensed
22 veterinarian pursuant to rules promulgated by the administrator.

23 (d) Until December 31, 2021, ensures that an employee of the
24 animal control shelter who received, and can document the
25 completion of, the training required immediately before ~~the~~
26 ~~effective date of the 2018 amendatory act that amended this section~~
27 **May 22, 2018** only administers a commercially prepared solution of
28 xylazine hydrochloride to sedate or immobilize the animals
29 described in this subsection. Beginning January 1, 2022, the



1 employee described in this subdivision must have received, and be
 2 able to document the completion of, the training described in
 3 subdivision (c) to administer an animal tranquilizer to perform
 4 euthanasia on the animals described in this subsection.

5 (e) Certifies that only an individual described in subdivision
 6 (c) or (d) or an individual otherwise permitted to use a controlled
 7 substance pursuant to this article will administer an animal
 8 tranquilizer according to written procedures established by the
 9 animal control shelter.

10 (f) Beginning January 1, 2022, certifies that the individual
 11 in charge of the day-to-day operations of the animal control
 12 shelter has received, and can document the completion of, the
 13 training described in subdivision (c).

14 (g) Complies with all state and federal laws, rules, and
 15 regulations regarding the acquisition, use, and security of
 16 controlled substances.

17 **(17)** ~~(16)~~—The application described in subsection ~~(15)~~—**(16)**
 18 must include the names and business addresses of all individuals
 19 employed by the animal control shelter who have been trained as
 20 described in subsection ~~(15)(e)~~, **(16)(c)**, (d), and (f) and must
 21 include documented proof of the training. The list of names and
 22 business addresses must be updated every 6 months.

23 **(18)** ~~(17)~~—If an animal control shelter issued a permit
 24 pursuant to subsection ~~(15)~~—**(16)** does not have in its employ an
 25 individual trained as described in subsection ~~(15)(e)~~, **(16)(c)** or
 26 (d) and ~~(15)(f)~~, **(16)(f)**, the animal control shelter shall
 27 immediately notify the administrator and shall cease to administer
 28 an animal tranquilizer for the purposes described in subsection
 29 ~~(15)~~—**(16)** until the administrator is notified that 1 of the



1 following has occurred:

2 (a) An individual trained as described in subsection ~~(15)(e)~~,
3 **(16)(c)**, (d), or (f) has been hired by the animal control shelter.

4 (b) An individual employed by the animal control shelter has
5 been trained as described in subsection ~~(15)(e)~~ **(16)(c)** or (f).

6 **(19)** ~~(18)~~ A veterinarian, including a veterinarian who trains
7 individuals as described in subsection ~~(15)(e)~~, **(16)(c)**, (d), or
8 (f), is not civilly or criminally liable for the use of an animal
9 tranquilizer by an animal control shelter unless the veterinarian
10 is employed by or under contract with the animal control shelter
11 and the terms of the veterinarian's employment or the contract
12 require the veterinarian to be responsible for the use or
13 administration of an animal tranquilizer.

14 **(20)** ~~(19)~~ As used in this section:

15 (a) "Animal tranquilizer" means a commercially prepared
16 solution of xylazine hydrochloride, a commercially prepared
17 solution of ketamine, or a commercially prepared compound
18 containing tiletamine and zolazepam.

19 (b) "Class B dealer" means a class B dealer licensed by the
20 United States Department of Agriculture pursuant to the animal
21 welfare act, 7 USC 2131 to ~~2159~~ **2160** and the department of
22 agriculture and rural development pursuant to 1969 PA 224, MCL
23 287.381 to 287.395.

24 Sec. 16221. Subject to section 16221b, the department shall
25 investigate any allegation that 1 or more of the grounds for
26 disciplinary subcommittee action under this section exist, and may
27 investigate activities related to the practice of a health
28 profession by a licensee, a registrant, or an applicant for
29 licensure or registration. The department may hold hearings,



1 administer oaths, and order the taking of relevant testimony. After
2 its investigation, the department shall provide a copy of the
3 administrative complaint to the appropriate disciplinary
4 subcommittee. The disciplinary subcommittee shall proceed under
5 section 16226 if it finds that 1 or more of the following grounds
6 exist:

7 (a) Except as otherwise specifically provided in this section,
8 a violation of general duty, consisting of negligence or failure to
9 exercise due care, including negligent delegation to or supervision
10 of employees or other individuals, whether or not injury results,
11 or any conduct, practice, or condition that impairs, or may impair,
12 the ability to safely and skillfully engage in the practice of the
13 health profession.

14 (b) Personal disqualifications, consisting of 1 or more of the
15 following:

16 (i) Incompetence.

17 (ii) Subject to sections 16165 to 16170a, substance use
18 disorder as defined in section 100d of the mental health code, 1974
19 PA 258, MCL 330.1100d.

20 (iii) Mental or physical inability reasonably related to and
21 adversely affecting the licensee's or registrant's ability to
22 practice in a safe and competent manner.

23 (iv) Declaration of mental incompetence by a court of competent
24 jurisdiction.

25 (v) Conviction of a misdemeanor punishable by imprisonment for
26 a maximum term of 2 years; conviction of a misdemeanor involving
27 the illegal delivery, possession, or use of a controlled substance;
28 or conviction of any felony other than a felony listed or described
29 in another subparagraph of this subdivision. A certified copy of



1 the court record is conclusive evidence of the conviction.

2 (vi) Lack of good moral character.

3 (vii) Conviction of a criminal offense under section 520e or
4 520g of the Michigan penal code, 1931 PA 328, MCL 750.520e and
5 750.520g. A certified copy of the court record is conclusive
6 evidence of the conviction.

7 (viii) Conviction of a violation of section 492a of the Michigan
8 penal code, 1931 PA 328, MCL 750.492a. A certified copy of the
9 court record is conclusive evidence of the conviction.

10 (ix) Conviction of a misdemeanor or felony involving fraud in
11 obtaining or attempting to obtain fees related to the practice of a
12 health profession. A certified copy of the court record is
13 conclusive evidence of the conviction.

14 (x) Final adverse administrative action by a licensure,
15 registration, disciplinary, or certification board involving the
16 holder of, or an applicant for, a license or registration regulated
17 by another state or a territory of the United States, by the United
18 States military, by the federal government, or by another country.
19 A certified copy of the record of the board is conclusive evidence
20 of the final action.

21 (xi) Conviction of a misdemeanor that is reasonably related to
22 or that adversely affects the licensee's or registrant's ability to
23 practice in a safe and competent manner. A certified copy of the
24 court record is conclusive evidence of the conviction.

25 (xii) Conviction of a violation of section 430 of the Michigan
26 penal code, 1931 PA 328, MCL 750.430. A certified copy of the court
27 record is conclusive evidence of the conviction.

28 (xiii) Conviction of a criminal offense under section 83, 84,
29 316, 317, 321, 520b, 520c, 520d, or 520f of the Michigan penal



code, 1931 PA 328, MCL 750.83, 750.84, 750.316, 750.317, 750.321, 750.520b, 750.520c, 750.520d, and 750.520f. A certified copy of the court record is conclusive evidence of the conviction.

(xiv) Conviction of a violation of section 136 or 136a of the Michigan penal code, 1931 PA 328, MCL 750.136 and 750.136a. A certified copy of the court record is conclusive evidence of the conviction.

(c) Prohibited acts, consisting of 1 or more of the following:

(i) Fraud or deceit in obtaining or renewing a license or registration.

(ii) Permitting a license or registration to be used by an unauthorized person.

(iii) Practice outside the scope of a license.

(iv) Obtaining, possessing, or attempting to obtain or possess a controlled substance ~~as defined in section 7104~~ or a drug as defined in section 7105 without lawful authority; or selling, prescribing, giving away, or administering drugs for other than lawful diagnostic or therapeutic purposes.

(d) Except as otherwise specifically provided in this section, unethical business practices, consisting of 1 or more of the following:

(i) False or misleading advertising.

(ii) Dividing fees for referral of patients or accepting kickbacks on medical or surgical services, appliances, or medications purchased by or in behalf of patients.

(iii) Fraud or deceit in obtaining or attempting to obtain third party reimbursement.

(e) Except as otherwise specifically provided in this section, unprofessional conduct, consisting of 1 or more of the following:



1 (i) Misrepresentation to a consumer or patient or in obtaining
2 or attempting to obtain third party reimbursement in the course of
3 professional practice.

4 (ii) Betrayal of a professional confidence.

5 (iii) Promotion for personal gain of an unnecessary drug,
6 device, treatment, procedure, or service.

7 (iv) Either of the following:

8 (A) A requirement by a licensee other than a physician or a
9 registrant that an individual purchase or secure a drug, device,
10 treatment, procedure, or service from another person, place,
11 facility, or business in which the licensee or registrant has a
12 financial interest.

13 (B) A referral by a physician for a designated health service
14 that violates 42 USC 1395nn or a regulation promulgated under that
15 section. For purposes of this subdivision, 42 USC 1395nn and the
16 regulations promulgated under that section as they exist on June 3,
17 2002 are incorporated by reference. A disciplinary subcommittee
18 shall apply 42 USC 1395nn and the regulations promulgated under
19 that section regardless of the source of payment for the designated
20 health service referred and rendered. If 42 USC 1395nn or a
21 regulation promulgated under that section is revised after June 3,
22 2002, the department shall officially take notice of the revision.
23 Within 30 days after taking notice of the revision, the department
24 shall decide whether or not the revision pertains to referral by
25 physicians for designated health services and continues to protect
26 the public from inappropriate referrals by physicians. If the
27 department decides that the revision does both of those things, the
28 department may promulgate rules to incorporate the revision by
29 reference. If the department does promulgate rules to incorporate



1 the revision by reference, the department shall not make any
2 changes to the revision. As used in this sub-subparagraph,
3 "designated health service" means that term as defined in 42 USC
4 1395nn and the regulations promulgated under that section and
5 "physician" means that term as defined in sections 17001 and 17501.

6 (v) For a physician who makes referrals under 42 USC 1395nn or
7 a regulation promulgated under that section, refusing to accept a
8 reasonable proportion of patients eligible for Medicaid and
9 refusing to accept payment from Medicaid or Medicare as payment in
10 full for a treatment, procedure, or service for which the physician
11 refers the individual and in which the physician has a financial
12 interest. A physician who owns all or part of a facility in which
13 he or she provides surgical services is not subject to this
14 subparagraph if a referred surgical procedure he or she performs in
15 the facility is not reimbursed at a minimum of the appropriate
16 Medicaid or Medicare outpatient fee schedule, including the
17 combined technical and professional components.

18 (vi) Any conduct by a health professional with a patient while
19 he or she is acting within the health profession for which he or
20 she is licensed or registered, including conduct initiated by a
21 patient or to which the patient consents, that is sexual or may
22 reasonably be interpreted as sexual, including, but not limited to,
23 sexual intercourse, kissing in a sexual manner, or touching of a
24 body part for any purpose other than appropriate examination,
25 treatment, or comfort.

26 (vii) Offering to provide practice-related services, such as
27 drugs, in exchange for sexual favors.

28 (viii) A violation of section 16655(4) by a dental therapist.

29 (f) Failure to notify under section 16222(3) or (4).



1 (g) Failure to report a change of name or mailing address as
2 required in section 16192.

3 (h) A violation, or aiding or abetting in a violation, of this
4 article or of a rule promulgated under this article.

5 (i) Failure to comply with a subpoena issued pursuant to this
6 part, failure to respond to a complaint issued under this article,
7 article 7, or article 8, failure to appear at a compliance
8 conference or an administrative hearing, or failure to report under
9 section 16222(1) or 16223.

10 (j) Failure to pay an installment of an assessment levied
11 under the insurance code of 1956, 1956 PA 218, MCL 500.100 to
12 500.8302, within 60 days after notice by the appropriate board.

13 (k) A violation of section 17013 or 17513.

14 (l) Failure to meet 1 or more of the requirements for licensure
15 or registration under section 16174.

16 (m) A violation of section 17015, 17015a, 17017, 17515, or
17 17517.

18 (n) A violation of section 17016 or 17516.

19 (o) Failure to comply with section 9206(3).

20 (p) A violation of section 5654 or 5655.

21 (q) A violation of section 16274.

22 (r) A violation of section 17020 or 17520.

23 (s) A violation of the medical records access act, 2004 PA 47,
24 MCL 333.26261 to 333.26271.

25 (t) A violation of section 17764(2).

26 (u) Failure to comply with the terms of a practice agreement
27 described in section 17047(2)(a) or (b), 17547(2)(a) or (b), or
28 18047(2)(a) or (b).

29 (v) A violation of section 7303a(2).



(w) A violation of section 7303a(4) or (5).

(x) A violation of section 7303b.

(y) A violation of section 17754a.

Sec. 16221b. **(1)** If the department has a reasonable basis to believe that a licensee has violated ~~section 7303a(4) or (5), any~~ **of the following**, the department is not required to investigate under section 16221 or 16231 and may issue a letter to the licensee notifying the licensee that he or she may be in violation of ~~section 7303a(4) or (5).~~ **the applicable section:**

(a) Section 7303a(4).

(b) Section 7303a(5).

(c) Section 17754a.

(2) A letter that is issued under this section is not considered discipline.

Sec. 16226. (1) After finding the existence of 1 or more of the grounds for disciplinary subcommittee action listed in section 16221, a disciplinary subcommittee shall impose 1 or more of the following sanctions for each violation:

<u>Violations of Section 16221</u>	<u>Sanctions</u>
Subdivision (a), (b) (i),	Probation, limitation, denial,
(b) (ii), (b) (iii), (b) (iv),	suspension, revocation,
(b) (v), (b) (vi), (b) (vii),	permanent revocation,
(b) (ix), (b) (x), (b) (xi),	restitution, or fine.
or (b) (xii)	
Subdivision (b) (viii)	Revocation, permanent revocation,
	or denial.



1	Subdivision (b) (<i>xiii</i>)	Permanent revocation
2		for a violation described in
3		subsection (5); otherwise,
4		probation, limitation, denial,
5		suspension, revocation,
6		restitution, or fine.
7		
8	Subdivision (b) (<i>xiv</i>)	Permanent revocation.
9		
10	Subdivision (c) (<i>i</i>)	Denial, revocation, suspension,
11		probation, limitation, or fine.
12		
13	Subdivision (c) (<i>ii</i>)	Denial, suspension, revocation,
14		restitution, or fine.
15		
16	Subdivision (c) (<i>iii</i>)	Probation, denial, suspension,
17		revocation, restitution, or fine.
18		
19	Subdivision (c) (<i>iv</i>)	Fine, probation, denial,
20	or (d) (<i>iii</i>)	suspension, revocation, permanent
21		revocation, or restitution.
22		
23	Subdivision (d) (<i>i</i>)	Reprimand, fine, probation,
24	or (d) (<i>ii</i>)	denial, or restitution.
25		
26	Subdivision (e) (<i>i</i>),	Reprimand, fine, probation,
27	(e) (<i>iii</i>), (e) (<i>iv</i>), (e) (<i>v</i>),	limitation, suspension,
28	(h), or (s)	revocation, permanent revocation,

1		denial, or restitution.
2		
3	Subdivision (e) (ii)	Reprimand, probation, suspension,
4	or (i) (i)	revocation, permanent
5		revocation, restitution,
6		denial, or fine.
7		
8	Subdivision (e) (vi) ,	Probation, suspension, revocation
9	(e) (vii) , or (e) (viii)	limitation, denial,
10		restitution, or fine.
11		
12	Subdivision (f)	Reprimand, denial, limitation,
13		probation, or fine.
14		
15	Subdivision (g)	Reprimand or fine.
16		
17	Subdivision (j)	Suspension or fine.
18		
19	Subdivision (k) , (p) ,	Reprimand, probation, suspension,
20	or (r)	revocation, permanent revocation,
21		or fine.
22		
23	Subdivision (l)	Reprimand, denial, or
24		limitation.
25		
26	Subdivision (m) or (o)	Denial, revocation, restitution,
27		probation, suspension,
28		limitation, reprimand, or fine.

1
2 Subdivision (n) Revocation or denial.
3
4 Subdivision (q) Revocation.
5
6 Subdivision (t) Revocation, permanent revocation,
7 fine, or restitution.
8
9 Subdivision (u) Denial, revocation, probation,
10 suspension, limitation,
11 reprimand,
12 or fine.
13
14 Subdivision (v) or (x) Probation, limitation, denial,
15 fine, suspension, revocation, or
16 permanent revocation.
17
18 Subdivision (w) Denial, fine, reprimand,
19 probation, limitation,
20 suspension, revocation, or
21 permanent revocation.
22 **Subdivision (y) Subject to subsection (7),**
23 **fine.**

24 (2) Determination of sanctions for violations under this
25 section shall be made by a disciplinary subcommittee. If, during
26 judicial review, the court of appeals determines that a final
27 decision or order of a disciplinary subcommittee prejudices
28 substantial rights of the petitioner for 1 or more of the grounds
29 listed in section 106 of the administrative procedures act of 1969,

1 ~~1969 PA 306,~~ MCL 24.306, and holds that the final decision or order
2 is unlawful and is to be set aside, the court shall state on the
3 record the reasons for the holding and may remand the case to the
4 disciplinary subcommittee for further consideration.

5 (3) A disciplinary subcommittee may impose a fine in an amount
6 that does not exceed \$250,000.00 for a violation of section
7 16221(a) or (b). A disciplinary subcommittee shall impose a fine of
8 at least \$25,000.00 if the violation of section 16221(a) or (b)
9 results in the death of 1 or more patients.

10 (4) A disciplinary subcommittee may require a licensee or
11 registrant or an applicant for licensure or registration who has
12 violated this article, article 7, or article 8 or a rule
13 promulgated under this article, article 7, or article 8 to
14 satisfactorily complete an educational program, a training program,
15 or a treatment program, a mental, physical, or professional
16 competence examination, or a combination of those programs and
17 examinations.

18 (5) A disciplinary subcommittee shall impose the sanction of
19 permanent revocation for a violation of section 16221(b) *(xiii)* if the
20 violation occurred while the licensee or registrant was acting
21 within the health profession for which he or she was licensed or
22 registered.

23 (6) Except as otherwise provided in subsection (5) and this
24 subsection, a disciplinary subcommittee shall not impose the
25 sanction of permanent revocation under this section without a
26 finding that the licensee or registrant engaged in a pattern of
27 intentional acts of fraud or deceit resulting in personal financial
28 gain to the licensee or registrant and harm to the health of
29 patients under the licensee's or registrant's care. This subsection



1 does not apply if a disciplinary subcommittee finds that a licensee
2 or registrant has violated section 16221(b) (xiv) .

3 **(7) A disciplinary subcommittee shall impose a fine of \$250.00**
4 **for each violation of section 16221(y) .**

5 Sec. 17744. (1) A prescriber may designate an agent to act on
6 behalf of or at the discretion of that prescriber. A designation of
7 an agent by a prescriber under this section is not required to be
8 in writing to be a valid designation. If a designation of an agent
9 by a prescriber under this section is contained in a written
10 document, the prescriber or the agent may transmit that document to
11 a pharmacy that will dispense a prescription issued by that
12 prescriber.

13 (2) Only a prescriber acting within the scope of his or her
14 practice may issue a prescription. An agent may prepare and
15 transmit a prescription that has been signed by the prescriber,
16 including a signature that meets the requirements of section 17754
17 **or 17754a.** The prescriber issuing a prescription and the pharmacist
18 dispensing a drug or device under a prescription is responsible for
19 all of the requirements of state and federal law, rules, and
20 regulations regarding the issuance of prescriptions and dispensing
21 of drugs or devices under prescriptions.

22 (3) A prescriber or his or her agent may transmit to a
23 pharmacy a prescription that is contained within a patient's chart
24 in a health facility or agency licensed under article 17 or other
25 medical institution. A prescription that is contained within a
26 patient's chart in a health facility or agency licensed under
27 article 17 or other medical institution and that is created in an
28 electronic format may contain more than 6 prescriptions and may
29 contain prescriptions for schedule 3 through 5 controlled



1 substances and noncontrolled substances on the same form.

2 Sec. 17751. (1) A pharmacist shall not dispense a drug
3 requiring a prescription under the federal act or a law of this
4 state except under authority of an original prescription or an
5 equivalent record of an original prescription approved by the
6 board.

7 (2) Subject to subsection (5), a pharmacist may dispense a
8 prescription written and signed; written or created in an
9 electronic format, signed, and transmitted by facsimile; or
10 transmitted electronically or by other means of communication by a
11 physician prescriber, dentist prescriber, or veterinarian
12 prescriber in another state, but not including a prescription for a
13 controlled substance except under circumstances described in
14 section 17763(e), only if the pharmacist in the exercise of his or
15 her professional judgment determines all of the following:

16 (a) Except as otherwise authorized under section 5110, 17744a,
17 or 17744b, if the prescriber is a physician or dentist, that the
18 prescription was issued pursuant to an existing physician-patient
19 or dentist-patient relationship.

20 (b) That the prescription is authentic.

21 (c) That the prescribed drug is appropriate and necessary for
22 the treatment of an acute, chronic, or recurrent condition.

23 (3) A pharmacist or a prescriber shall dispense a prescription
24 only if the prescription falls within the scope of practice of the
25 prescriber.

26 (4) A pharmacist shall not knowingly dispense a prescription
27 after the death of the prescriber or patient.

28 (5) A pharmacist shall not dispense a drug or device under a
29 prescription transmitted by facsimile or created in electronic



1 format and printed out for use by the patient unless the document
2 is manually signed by the prescriber. This subsection does not
3 apply to a prescription that is transmitted by a computer to a
4 facsimile machine if that prescription complies with section 17754
5 **or 17754a.**

6 (6) After consultation with and agreement from the prescriber,
7 a pharmacist may add or change a patient's address, a dosage form,
8 a drug strength, a drug quantity, a direction for use, or an issue
9 date with regard to a prescription. A pharmacist shall note the
10 details of the consultation and agreement required under this
11 subsection on the prescription and shall maintain that
12 documentation with the prescription as required in section 17752. A
13 pharmacist shall not change the patient's name, controlled
14 substance prescribed unless authorized to dispense a lower cost
15 generically equivalent drug product under section 17755, or the
16 prescriber's signature with regard to a prescription.

17 (7) A prescription that is contained within a patient's chart
18 in a health facility or agency licensed under article 17 or other
19 medical institution and that is transmitted to a pharmacy under
20 section 17744 is the original prescription. If all other
21 requirements of this part are met, a pharmacist shall dispense a
22 drug or device under a prescription described in this subsection. A
23 pharmacist may dispense a drug or device under a prescription
24 described in this subsection even if the prescription does not
25 contain the quantity ordered. If a prescription described in this
26 subsection does not contain the quantity ordered, the pharmacist
27 shall consult with the prescriber to determine an agreed-upon
28 quantity. The pharmacist shall record the quantity dispensed on the
29 prescription and shall maintain that documentation with the



1 prescription as required in section 17752.

2 (8) If, after consulting with a patient, a pharmacist
3 determines in the exercise of his or her professional judgment that
4 dispensing additional quantities of a prescription drug is
5 appropriate for the patient, the pharmacist may dispense, at one
6 time, additional quantities of the prescription drug up to the
7 total number of dosage units authorized by the prescriber on the
8 original prescription for the patient and any refills of the
9 prescription. Except for a controlled substance included in
10 schedule 5 that does not contain an opioid, this subsection does
11 not apply to a prescription for a controlled substance.

12 Sec. 17754. (1) Except as otherwise provided under article 7,
13 article 8, and the federal act, a prescription may be transmitted
14 electronically if the prescription is transmitted in compliance
15 with the health insurance portability and accountability act of
16 1996, Public Law 104-191, or regulations promulgated under that
17 act, 45 CFR parts 160 and 164, by a prescriber or his or her agent
18 and the data are not altered or modified in the transmission
19 process. The electronically transmitted prescription shall include
20 all of the following information:

21 (a) The name, address, and telephone number of the prescriber.

22 (b) Except as otherwise authorized under section 5110, 17744a,
23 or 17744b, the full name of the patient for whom the prescription
24 is issued.

25 (c) An electronic signature or other identifier that
26 specifically identifies and authenticates the prescriber or his or
27 her agent.

28 (d) The time and date of the transmission.

29 (e) The identity of the pharmacy intended to receive the



1 transmission.

2 (f) Any other information required by the federal act or state
3 law.

4 (2) The electronic equipment or system utilized in the
5 transmission and communication of prescriptions shall provide
6 adequate confidentiality safeguards and be maintained to protect
7 patient confidentiality as required under any applicable federal
8 and state law and to ensure against unauthorized access. The
9 electronic transmission of a prescription shall be communicated in
10 a retrievable, recognizable form acceptable to the intended
11 recipient. The electronic form utilized in the transmission of a
12 prescription shall not include "dispense as written" or "d.a.w." as
13 the default setting.

14 (3) Before dispensing a prescription that is electronically
15 transmitted, the pharmacist shall exercise professional judgment
16 regarding the accuracy, validity, and authenticity of the
17 transmitted prescription.

18 (4) An electronically transmitted prescription that meets the
19 requirements of this section is the original prescription.

20 (5) **This section does not apply beginning on the date on which**
21 **section 17754a applies.**

22 **Sec. 17754a. (1) Except as otherwise provided under article 8,**
23 **the federal act, or subsection (5), and subject to subsection (10),**
24 **beginning January 1, 2021, a prescriber or his or her agent shall**
25 **electronically transmit a prescription, including a prescription**
26 **for a controlled substance, directly to a pharmacy of the patient's**
27 **choice. A prescription that is transmitted electronically under**
28 **this section must be in compliance with the health insurance**
29 **portability and accountability act of 1996, Public Law 104-191, or**



1 regulations promulgated under that act, 45 CFR parts 160 and 164,
2 and the data must not be altered or modified in the transmission
3 process. The electronically transmitted prescription must include
4 all of the following information:

5 (a) The name, address, and telephone number of the prescriber.

6 (b) Except as otherwise authorized under section 5110, 17744a,
7 or 17744b, the full name of the patient for whom the prescription
8 is issued.

9 (c) An electronic signature or other identifier that
10 specifically identifies and authenticates the prescriber or his or
11 her agent.

12 (d) The time and date of the transmission.

13 (e) The identity of the pharmacy intended to receive the
14 transmission.

15 (f) Any other information required by the federal act or state
16 law.

17 (2) The electronic equipment or system utilized in the
18 transmission and communication of prescriptions under this section
19 must provide adequate confidentiality safeguards and be maintained
20 to protect patient confidentiality as required under any applicable
21 federal and state law and to ensure against unauthorized access.
22 The electronic transmission of a prescription under this section
23 must be communicated in a retrievable, recognizable form acceptable
24 to the intended recipient. The electronic form utilized in the
25 transmission of a prescription must not include "dispense as
26 written" or "d.a.w." as the default setting.

27 (3) Before dispensing a prescription that is electronically
28 transmitted under this section, the pharmacist shall exercise
29 professional judgment regarding the accuracy, validity, and



1 authenticity of the transmitted prescription.

2 (4) An electronically transmitted prescription that meets the
3 requirements of this section is the original prescription.

4 (5) The requirement to transmit a prescription electronically
5 under subsection (1) does not apply under any of the following
6 circumstances:

7 (a) If the prescription is issued by a prescriber who is a
8 veterinarian licensed under this article.

9 (b) Subject to subsection (6), if the prescription is issued
10 under a circumstance in which electronic transmission is not
11 available due to a temporary technological or electrical failure.

12 (c) If the prescription is issued by a prescriber who has
13 received a waiver from the department under subsection (7).

14 (d) Subject to subsection (6), if the prescription is issued
15 by a prescriber who reasonably believes that electronically
16 transmitting the prescription would make it impractical for the
17 patient who is the subject of the prescription to obtain the
18 prescription drug in a timely manner and that the delay would
19 adversely affect the patient's medical condition.

20 (e) If the prescription is orally prescribed under section
21 7333(3) or (4).

22 (f) Subject to subsection (6), if the prescription is issued
23 by a prescriber to be dispensed outside of this state.

24 (g) If the prescription is issued by a prescriber who is
25 located outside of this state to be dispensed by a pharmacy located
26 inside of this state.

27 (h) If the prescription is issued and dispensed in the same
28 health care facility and the individual for whom the prescription
29 is issued uses the drug exclusively in the health care facility. As



1 used in this subdivision, "health care facility" includes, but is
2 not limited to, any of the following:

3 (i) A hospital.

4 (ii) A hospice.

5 (iii) A dialysis treatment clinic.

6 (iv) A freestanding surgical outpatient facility.

7 (v) A nursing home.

8 (vi) A long-term care facility that provides rehabilitative,
9 restorative, or ongoing skilled nursing care to an individual who
10 is in need of assistance with activities of daily living.

11 (i) Subject to subsection (6), if the prescription contains
12 content that is not supported by the National Council for
13 Prescription Drug Programs Prescriber/Pharmacist Interface SCRIPT
14 Standard.

15 (j) Subject to subsection (6), if the prescription is for a
16 drug for which the FDA requires the prescription to contain content
17 that cannot be transmitted electronically.

18 (k) If the prescription is issued under circumstances in which
19 the prescriber is not required to include on the prescription a
20 name of a patient for whom the prescription is issued including,
21 but not limited to, a prescription issued under section 5110.

22 (l) Subject to subsection (6), if the prescription is issued by
23 a prescriber who is prescribing the drug under a research protocol.

24 (6) If a prescription for a controlled substance is not
25 electronically transmitted under an exception described in
26 subsection (5)(b), (d), (f), (i), (j), or (l), the prescriber shall
27 document the applicable exception at the time the prescriber issues
28 the prescription. If the prescription is not electronically
29 transmitted under an exception described in subsection (5)(d), the



1 prescriber shall also document the specific reason for not
2 electronically transmitting the prescription. A prescriber shall
3 provide the documentation required under this subsection to the
4 department on request.

5 (7) If a prescriber cannot meet the requirements of subsection
6 (1) or (2), the prescriber may apply to the department for a
7 waiver. The department shall grant a waiver to a prescriber if the
8 department determines that the prescriber cannot meet the
9 requirements of subsection (1) or (2) due to an economic hardship,
10 a technological limitation that is not reasonably within the
11 control of the prescriber, such as insufficient internet
12 connectivity or the use of a health record technology certified by
13 the federal Centers for Medicare and Medicaid Services that does
14 not allow for the electronic transmission of a prescription for a
15 controlled substance, or another exceptional circumstance. A waiver
16 that is granted under this subsection is valid for a period not to
17 exceed 1 year and is renewable.

18 (8) A pharmacist who receives a prescription that was not
19 transmitted electronically to the pharmacy may dispense the
20 prescription without determining whether an exception under
21 subsection (5) applies.

22 (9) The department, in consultation with the board, shall
23 promulgate rules to implement this section.

24 (10) If the federal Centers for Medicare and Medicaid Services
25 delays the Medicare requirement for the electronic transmission of
26 prescriptions for controlled substances beyond January 1, 2021,
27 then the department shall, by rule, delay the implementation date
28 of subsection (1) to the date established by the federal Centers
29 for Medicare and Medicaid Services for the Medicare requirement.

