ENROLLED HOUSE BILL No. 4352

AN ACT to amend 1978 PA 368, entitled “An act to protect and promote the public health; to codify, revise, consolidate, classify, and add to the laws relating to public health; to provide for the prevention and control of diseases and disabilities; to provide for the classification, administration, regulation, financing, and maintenance of personal, environmental, and other health services and activities; to create or continue, and prescribe the powers and duties of, departments, boards, commissions, councils, committees, task forces, and other agencies; to prescribe the powers and duties of governmental entities and officials; to regulate occupations, facilities, and agencies affecting the public health; to regulate health maintenance organizations and certain third party administrators and insurers; to provide for the imposition of a regulatory fee; to provide for the levy of taxes against certain health facilities or agencies; to promote the efficient and economical delivery of health care services, to provide for the appropriate utilization of health care facilities and services, and to provide for the collection and use of data and information; to provide for the transfer of property; to provide certain immunity from liability; to regulate and prohibit the sale and offering for sale of drug paraphernalia under certain circumstances; to provide for the implementation of federal law; to provide for penalties and remedies; to provide for sanctions for violations of this act and local ordinances; to provide for an appropriation and supplements; to repeal certain acts and parts of acts; to repeal certain parts of this act; and to repeal certain parts of this act on specific dates,” by amending sections 17745, 17751, 17754, and 17757 (MCL 333.17745, 333.17751, 333.17754, and 333.17757), sections 17745 and 17757 as amended by 2011 PA 210 and sections 17751 and 17754 as amended by 2012 PA 209, and by adding section 17744a.

The People of the State of Michigan enact:

Sec. 17744a. (1) Notwithstanding any provision of this act to the contrary, a prescriber may issue a prescription for and a dispensing prescriber or pharmacist may dispense auto-injectable epinephrine to a school board for the purpose of meeting the requirements of section 1179a of the revised school code, 1976 PA 451, MCL 380.1179a. When issuing a prescription for or dispensing auto-injectable epinephrine to a school board as authorized under this section, the prescriber, dispensing prescriber, or pharmacist, as appropriate, shall insert the name of the school board as the name of the patient. A school employee who is a licensed registered professional nurse or who is trained in the administration of an epinephrine auto-injector under section 1179a of the revised school code, 1976 PA 451, MCL 380.1179a, may possess and administer an epinephrine auto-injector dispensed to a school board under this section.

(172)
(2) A prescriber who issues a prescription for or a dispensing prescriber or pharmacist who dispenses auto-injectable epinephrine to a school board as authorized under this section is not liable in a civil action for a properly stored and dispensed epinephrine auto-injector that was a proximate cause of injury or death to an individual due to the administration of or failure to administer the epinephrine auto-injector.

Sec. 17745. (1) Except as otherwise provided in this subsection, a prescriber who wishes to dispense prescription drugs shall obtain from the board a drug control license for each location in which the storage and dispensing of prescription drugs occur. A drug control license is not necessary if the dispensing occurs in the emergency department, emergency room, or trauma center of a hospital licensed under article 17 or if the dispensing involves only the issuance of complimentary starter dose drugs.

(2) Except as otherwise provided in section 17744a, a dispensing prescriber shall dispense prescription drugs only to his or her own patients.

(3) A dispensing prescriber shall include in a patient’s chart or clinical record a complete record, including prescription drug names, dosages, and quantities, of all prescription drugs dispensed directly by the dispensing prescriber or indirectly under his or her delegatory authority. If prescription drugs are dispensed under the prescriber’s delegatory authority, the delegatee who dispenses the prescription drugs shall initial the patient’s chart, clinical record, or log of prescription drugs dispensed. In a patient’s chart or clinical record, a dispensing prescriber shall distinguish between prescription drugs dispensed to the patient, prescription drugs prescribed for the patient, and prescription drugs dispensed or prescribed as authorized under section 17744a. A dispensing prescriber shall retain information required under this subsection for not less than 5 years after the information is entered in the patient’s chart or clinical record.

(4) A dispensing prescriber shall store prescription drugs under conditions that will maintain their stability, integrity, and effectiveness and will assure that the prescription drugs are free of contamination, deterioration, and adulteration.

(5) A dispensing prescriber shall store prescription drugs in a substantially constructed, securely lockable cabinet. Access to the cabinet shall be limited to individuals authorized to dispense prescription drugs in compliance with this part and article 7.

(6) Unless otherwise requested by a patient, a dispensing prescriber shall dispense a prescription drug in a safety closure container that complies with the poison prevention packaging act of 1970, 15 USC 1471 to 1477.

(7) A dispensing prescriber shall dispense a drug in a container that bears a label containing all of the following information:
   (a) The name and address of the location from which the prescription drug is dispensed.
   (b) Except as otherwise authorized under section 17744a, the patient’s name and record number.
   (c) The date the prescription drug was dispensed.
   (d) The prescriber’s name or, if dispensed under the prescriber’s delegatory authority, the name of the delegatee.
   (e) The directions for use.
   (f) The name and strength of the prescription drug.
   (g) The quantity dispensed.
   (h) The expiration date of the prescription drug or the statement required under section 17756.

(8) A dispensing prescriber who dispenses a complimentary starter dose drug to a patient shall give the patient at least all of the following information, either by dispensing the complimentary starter dose drug to the patient in a container that bears a label containing the information or by giving the patient a written document that may include, but is not limited to, a preprinted insert that comes with the complimentary starter dose drug, that contains all of the following information:
   (a) The name and strength of the complimentary starter dose drug.
   (b) Directions for the patient’s use of the complimentary starter dose drug.
   (c) The expiration date of the complimentary starter dose drug or the statement required under section 17756.

(9) The information required under subsection (8) is in addition to, and does not supersede or modify, other state or federal law regulating the labeling of prescription drugs.

(10) In addition to meeting the requirements of this part, a dispensing prescriber who dispenses controlled substances shall comply with section 7303a.

(11) The board may periodically inspect locations from which prescription drugs are dispensed.

(12) The act, task, or function of dispensing prescription drugs shall be delegated only as provided in this part and sections 16215, 17048, 17076, 17212, and 17548.

(13) A supervising physician may delegate in writing to a pharmacist practicing in a hospital pharmacy within a hospital licensed under article 17 the receipt of complimentary starter dose drugs other than controlled substances as defined by article 7 or federal law. When the delegated receipt of complimentary starter dose drugs occurs, both the...
The pharmacist’s name and the supervising physician’s name shall be used, recorded, or otherwise indicated in connection with each receipt. A pharmacist described in this subsection may dispense a prescription for complimentary starter dose drugs written or transmitted by facsimile, electronic transmission, or other means of communication by a prescriber.

(14) As used in this section, “complimentary starter dose” means a prescription drug packaged, dispensed, and distributed in accordance with state and federal law that is provided to a dispensing prescriber free of charge by a manufacturer or distributor and dispensed free of charge by the dispensing prescriber to his or her patients.

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

(2) Subject to subsection (5), a pharmacist may dispense a prescription written and signed; written or created in an electronic format, signed, and transmitted by facsimile; or transmitted electronically or by other means of communication by a physician prescriber or dentist prescriber in a state other than Michigan, but not including a prescription for a controlled substance as defined in section 7104 except under circumstances described in section 17763(e), only if the pharmacist in the exercise of his or her professional judgment determines all of the following:

(a) Except as otherwise authorized under section 17744a, that the prescription was issued pursuant to an existing physician-patient or dentist-patient relationship.

(b) That the prescription is authentic.

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

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

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

(5) A pharmacist shall not dispense a drug or device under a prescription transmitted by facsimile or created in electronic format and printed out for use by the patient unless the document is manually signed by the prescriber. This subsection does not apply to a prescription that is transmitted by a computer to a facsimile machine if that prescription complies with section 17754.

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

(7) A prescription that is contained within a patient’s chart in a health facility or agency licensed under article 17 or other medical institution and that is transmitted to a pharmacy under section 17744 is the original prescription. If all other requirements of this part are met, a pharmacist shall dispense a drug or device under a prescription described in this subsection. A pharmacist may dispense a drug or device under a prescription described in this subsection even if the prescription does not contain the quantity ordered. If a prescription described in this subsection does not contain the quantity ordered, the pharmacist shall consult with the prescriber to determine an agreed-upon quantity. The pharmacist shall record the quantity dispensed on the prescription and shall maintain that documentation with the prescription as required in section 17752.

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

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

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

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

(d) The time and date of the transmission.

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

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

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

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

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

Sec. 17757. (1) Upon a request made in person or by telephone, a pharmacist engaged in the business of selling drugs
at retail shall provide the current selling price of a drug dispensed by that pharmacy or comparative current selling
prices of generic and brand name drugs dispensed by that pharmacy. The information shall be provided to the person
making the request before a drug is dispensed to the person. A person who makes a request for price information under
this subsection is not obligated to purchase the drug for which the price or comparative prices are requested.

(2) A pharmacist engaged in the business of selling drugs at retail shall conspicuously display the notice described
in subsection (3) at each counter over which prescription drugs are dispensed.

(3) The notice required under subsection (2) shall be in substantially the following form:

NOTICE TO CONSUMERS
ABOUT PRESCRIPTION DRUGS

Under Michigan law, you have the right to find out the price of a prescription drug before the pharmacist fills the
prescription. You are under no obligation to have the prescription filled here and may use this price information to shop
around at other pharmacies. You may request price information in person or by telephone.

Every pharmacy has the current selling prices of both generic and brand name drugs dispensed by the pharmacy.

Ask your pharmacist if a lower-cost generic drug is available to fill your prescription. A generic drug contains the
same medicine as a brand name drug and is a suitable substitute in most instances.

A generic drug may not be dispensed by your pharmacist if your doctor has written “dispense as written” or the
initials “d.a.w.” on the prescription.

If you have questions about the drugs that have been prescribed for you, ask your doctor or pharmacist for more
information.

To avoid dangerous drug interactions, let your doctor and pharmacist know about any other medications you are
taking. This is especially important if you have more than 1 doctor or have prescriptions filled at more than 1 pharmacy.

(4) The notice required under subsection (2) shall also contain the address and phone number of the board and the
department. The text of the notice shall be in at least 32-point bold type and shall be printed on paper at least 11 inches
by 17 inches in size. The notice may be printed on multiple pages.

(5) A copy of the notice required under subsection (2) shall be provided to each licensee by the department. The
department shall provide additional copies if needed. A person may duplicate or reproduce the notice if the duplication
or reproduction is a true copy of the notice as produced by the department, without any additions or deletions.

(6) The pharmacist shall furnish to the purchaser of a prescription drug at the time the drug is delivered to the
purchaser a receipt evidencing the transactions, which contains all of the following:

(a) The brand name of the drug, if applicable.
(b) The name of the manufacturer or the supplier of the drug, if the drug does not have a brand name.
(c) The strength of the drug, if significant.
(d) The quantity dispensed, if applicable.
(e) The name and address of the pharmacy.
(f) The serial number of the prescription.
(g) The date the prescription was originally dispensed.
(h) The name of the prescriber or, if prescribed under the prescriber’s delegatory authority, the name of the delegatee.
(i) Except as otherwise authorized under section 17744a, the name of the patient for whom the drug was prescribed.
(j) The price for which the drug was sold to the purchaser.

(7) The items required under subsection (6)(a), (b), and (c) may be omitted by a pharmacist only if the omission is
expressly required by the prescriber. The pharmacist shall retain a copy of each receipt for 90 days. The inclusion of the
items required under subsection (6) on the prescription container label is a valid receipt to the purchaser. Including the
items required under subsection (6) on the written prescription form and retaining the form constitutes retention of a
copy of the receipt.

(8) The board may promulgate rules to implement this section.
Enacting section 1. This amendatory act does not take effect unless House Bill No. 4353 of the 97th Legislature is enacted into law.

[Signature]
Clerk of the House of Representatives

[Signature]
Carol Monezy Viventi
Secretary of the Senate

Approved .................................................................

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Governor
AN ACT to amend 1976 PA 451, entitled “An act to provide a system of public instruction and elementary and secondary schools; to revise, consolidate, and clarify the laws relating to elementary and secondary education; to provide for the organization, regulation, and maintenance of schools, school districts, public school academies, intermediate school districts, and other public school entities; to prescribe rights, powers, duties, and privileges of schools, school districts, public school academies, intermediate school districts, and other public school entities; to provide for the regulation of school teachers and certain other school employees; to provide for school elections and to prescribe powers and duties with respect thereto; to provide for the levy and collection of taxes; to provide for the borrowing of money and issuance of bonds and other evidences of indebtedness; to establish a fund and provide for expenditures from that fund; to provide for and prescribe the powers and duties of certain state departments, the state board of education, and certain other boards and officials; to provide for licensure of boarding schools; to prescribe penalties; and to repeal acts and parts of acts,” by amending sections 1178 and 1179 (MCL 380.1178 and 380.1179), section 1178 as amended by 2006 PA 48 and section 1179 as amended by 2004 PA 73, and by adding section 1179a.

The People of the State of Michigan enact:

Sec. 1178. (1) Subject to subsection (2), a school administrator, teacher, or other school employee designated by the school administrator, who in good faith administers medication to a pupil in the presence of another adult or in an emergency that threatens the life or health of the pupil, pursuant to written permission of the pupil’s parent or guardian, and in compliance with the instructions of a physician, physician’s assistant, or certified nurse practitioner, or a school employee who in good faith administers an epinephrine auto-injector to an individual consistent with the policies under section 1179a, is not liable in a criminal action or for civil damages as a result of an act or omission in the administration of the medication or epinephrine auto-injector, except for an act or omission amounting to gross negligence or willful and wanton misconduct.

(2) If a school employee is a licensed registered professional nurse, subsection (1) applies to that school employee regardless of whether the medication or epinephrine auto-injector is administered in the presence of another adult.

(3) A school district, nonpublic school, member of a school board, or director or officer of a nonpublic school is not liable for damages in a civil action for injury, death, or loss to person or property allegedly arising from a person acting under this section.

Sec. 1179. (1) If the conditions prescribed in subsection (2) are met, notwithstanding any school or school district policy to the contrary, a pupil of a public school or nonpublic school may possess and use 1 or more of the following at school, on school-sponsored transportation, or at any activity, event, or program sponsored by or in which the pupil’s school is participating:

(a) A metered dose inhaler or a dry powder inhaler to alleviate asthmatic symptoms or for use before exercise to prevent the onset of asthmatic symptoms.

(b) An epinephrine auto-injector or epinephrine inhaler to treat anaphylaxis.
(2) Subsection (1) applies to a pupil if all of the following conditions are met:

(a) The pupil has written approval to possess and use the inhaler or epinephrine auto-injector as described in subsection (1) from the pupil’s physician or other health care provider authorized by law to prescribe an inhaler or epinephrine auto-injector and, if the pupil is a minor, from the pupil’s parent or legal guardian.

(b) The principal or other chief administrator of the pupil’s school has received a copy of each written approval required under subdivision (a) for the pupil.

(c) There is on file at the pupil’s school a written emergency care plan that contains specific instructions for the pupil’s needs, that is prepared by a physician licensed in this state in collaboration with the pupil and the pupil’s parent or legal guardian, and that is updated as necessary for changing circumstances.

(3) A school district, nonpublic school, member of a school board, director or officer of a nonpublic school, or employee of a school district or nonpublic school is not liable for damages in a civil action for injury, death, or loss to person or property allegedly arising from a pupil being prohibited by an employee of the school or school district from using an inhaler or epinephrine auto-injector because of the employee’s reasonable belief formed after a reasonable and ordinary inquiry that the conditions prescribed in subsection (2) had not been satisfied. A school district, nonpublic school, member of a school board, director or officer of a nonpublic school, or employee of a school district or nonpublic school is not liable for damages in a civil action for injury, death, or loss to person or property allegedly arising from a pupil being permitted by an employee of the school or school district to use or possess an inhaler or epinephrine auto-injector because of the employee’s reasonable belief formed after a reasonable and ordinary inquiry that the conditions prescribed in subsection (2) had been satisfied. This subsection does not eliminate, limit, or reduce any other immunity or defense that a school district, nonpublic school, member of a school board, director or officer of a nonpublic school, or employee of a school district or nonpublic school may have under section 1178 or other state law.

(4) As part of its general powers, a school district may request a pupil’s parent or legal guardian to provide an extra inhaler or epinephrine auto-injector to designated school personnel for use in case of emergency. A parent or legal guardian is not required to provide an extra inhaler or epinephrine auto-injector to school personnel.

(5) A principal or other chief administrator who is aware that a pupil is in possession of an inhaler or epinephrine auto-injector pursuant to this section shall notify each of the pupil’s classroom teachers of that fact and of the provisions of this section.

(6) As used in this section and in section 1179a:

(a) “School board” includes a school board, intermediate school board, or the board of directors of a public school academy.

(b) “School district” includes a school district, intermediate school district, or public school academy.

Sec. 1179a. (1) Beginning with the 2014-2015 school year, a school board shall ensure that, in each school it operates with an instructional and administrative staff of at least 10, there are at least 2 employees at the school who have been trained in the appropriate use and administration of an epinephrine auto-injector and that, in each school it operates with an instructional and administrative staff of fewer than 10, there is at least 1 employee at the school who has been trained in the appropriate use and administration of an epinephrine auto-injector. The training required under this subsection shall be conducted under the supervision of, and shall include evaluation by, a licensed registered professional nurse.

(2) Not later than the beginning of the 2014-2015 school year, a school board shall develop and implement policies that are consistent with the department’s medication administration guidelines, as revised under subsection (4), and that provide for the possession of at least 2 epinephrine auto-injectors in each school operated by the school board to be used for administration by a licensed registered professional nurse who is employed or contracted by the school district or by a school employee who is trained in the administration of an epinephrine auto-injector and that, in each school it operates with an instructional and administrative staff of fewer than 10, there is at least 1 employee at the school who has been trained in the appropriate use and administration of an epinephrine auto-injector. The policies shall authorize a licensed registered professional nurse who is employed or contracted by the school district or a school employee who is trained in the administration of an epinephrine auto-injector to administer an epinephrine auto-injector to a pupil who has a prescription on file at the school. The policies also shall authorize a licensed registered professional nurse who is employed or contracted by the school district or a school employee who is trained in the administration of an epinephrine auto-injector to administer an epinephrine auto-injector to any other individual on school grounds who is believed to have an anaphylactic reaction. The policies also shall require notification to the parent or legal guardian of a pupil to whom an epinephrine auto-injector has been administered.

(3) A licensed registered professional nurse who is employed or contracted by the school district or a school employee who is trained in the administration of an epinephrine auto-injector under subsection (1) may possess and administer an epinephrine auto-injector.

(4) The department, in conjunction with the department of community health and with input from the Michigan association of school nurses, the Michigan nurses association, the Michigan parent teacher association, the American college of allergy, asthma, and immunology, the Michigan chapter of the American academy of pediatrics, the
school-community health alliance of Michigan, and other school health organizations and entities, shall identify, develop, and adopt appropriate revisions to the medication administration guidelines issued by the department, including, but not limited to, those relating to the specification of training needs and requirements for the administration and maintenance of stock epinephrine auto-injectors, including stocking of both junior and regular dose epinephrine auto-injectors, as necessary, and storage requirements.

(5) At least annually, a school district shall report to the department, in the form and manner prescribed by the department, all instances of administration of an epinephrine auto-injector to a pupil at school. The reporting shall include at least all of the following:

(a) The number of instances of administration of an epinephrine auto-injector to a pupil at school in a school year.

(b) The number of pupils who were administered an epinephrine auto-injector at school who were not previously known to be severely allergic.

(c) The number of pupils who were administered an epinephrine auto-injector at school using the school’s stock of epinephrine auto-injectors.

(6) A school board shall attempt to obtain funding or resources from private sources, or from another source other than this state, for fulfilling the requirements of this section. If a school board is unable to obtain this alternative funding for all or part of its costs of complying with this section, the school board may apply to the department for reimbursement for the unfunded costs of complying with this section, in the form and manner prescribed by the department. The legislature shall appropriate funds for making this reimbursement. The department shall make the reimbursement according to the appropriation that is made for this purpose. The department annually shall submit a report to the legislature detailing the number of school boards that apply for reimbursement and the number of school boards that are able to secure alternative funding.

[Signatures]

Clerk of the House of Representatives

Secretary of the Senate

Governor