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STATE OF RHODE ISLAND

IN GENERAL ASSEMBLY

JANUARY SESSION, A.D. 2016

AN ACT

RELATING TO FOOD AND DRUGS - UNIFORMED CONTROLLED SUBSTANCE ACT

Introduced By: Representatives McNamara, Azzinaro, and Diaz

Date Introduced: March 03, 2016

Referred To: House Health, Education & Welfare

(Dept. of Health)

- (Dept. of Health) It is enacted by the General Assembly as follows: 1 SECTION 1. Section 21-28-3.32 of the General Laws in Chapter 21-28 entitled "Uniform 2 Controlled Substances Act" is hereby amended to read as follows: 3 21-28-3.32. Electronic prescription database. -- (a) The information contained in any 4 prescription drug monitoring database maintained by the department of health pursuant to § 21-5 28-3.18 of this chapter shall be disclosed only: (1) To a practitioner who certifies that the requested information is for the purpose of 6 evaluating the need for, or providing medical treatment to, a current patient to whom the 7 practitioner is prescribing or considering prescribing a controlled substance; 8 9 (2) To a pharmacist who certifies that the requested information is for a current client to 10 whom the pharmacist is dispensing, or considering dispensing, a controlled substance; 11 (3) To an authorized designee of the practitioner and/or pharmacist to consult the 12 prescription drug monitoring database on the practitioner's and/or pharmacist's behalf, provided 13 that: 14 (i) The designee so authorized is employed by the same professional practice or
 - pharmacy;
 - (ii) The practitioner or pharmacist takes reasonable steps to ensure that such designee is sufficiently competent in the use of the database;
- 18 (iii) The practitioner or pharmacist remains responsible for ensuring that access to the 19 database by the designee is limited to authorized purposes as provided for in subsections (a)(1)

2	(iv) The practitioner or pharmacist remains responsible for ensuring access to the
3	database by the designee occurs in a manner that protects the confidentiality of information
4	obtained from the database and remains responsible for any breach of confidentiality;
5	(v) The practitioner or pharmacist terminates the designee's access to the database at the
6	termination of the designee's employment; and
7	(vi) The ultimate decision as to whether or not to prescribe or dispense a controlled
8	substance remains with the practitioner or pharmacist and is reasonably informed by the relevant
9	controlled substance history information obtained from the database.
10	(4) Pursuant to a valid search warrant based on probable cause to believe a violation of
11	federal or state criminal law has occurred and that specified information contained in the database
12	would assist in the investigation of the crime;
13	(5) To a patient who requests his or her own prescription information, or the parent or
14	legal guardian of a minor child who requests the minor child's prescription information;
15	(6) To a health professional regulatory board that documents, in writing, that the
16	requested information is necessary for an investigation related to licensure, renewal, or
17	disciplinary action involving the applicant, licensee, or registrant to whom the requested
18	information pertains;
19	(7) To any vendor or contractor with whom the department has contracted to establish or
20	maintain the electronic system of the prescription drug monitoring database; or
21	(8) To public or private entities for statistical, research, or educational purposes, after
22	removing the patient and prescriber information that could be used to identify individual patients.
23	This shall not include entities receiving a waiver from the institutional review board.
24	(9) To any vendor, agent, contractor, or designee who operates an electronic health record
25	or clinical management system for the purpose of sharing data with practitioners, pharmacists, or
26	licensed health care facilities or designees.
27	(b) Information stored in the prescription drug monitoring database shall include only the
28	following:
29	(1) Patient's first and last name, and/or patient identification number; provided, however,
30	the patient's social security number shall not be recorded in whole or in part, patient sex, patient
31	date of birth, and patient address;
32	(2) Prescribing practitioner's name and drug enforcement administration prescriber
33	information number;
34	(3) Prescribing practitioner's office or hospital contact information;

and (a)(2) of this section;

1 (4) Prescription name, prescription number, prescription species code, national drug code 2 number, prescription dosage, prescription quantity, days' supply, new-refill code, number of 3 refills authorized, date the prescription was written, date the prescription was filled, payment 4 type; provided, however, no credit card number shall be recorded in whole or in part; and 5 (5) The drug enforcement administration pharmacy number of the pharmacy filling the prescription. 6 7 (c) The department shall disclose any information relating to a patient maintained in the 8 prescription drug monitoring database to that patient, at no cost to the patient, within thirty (30) 9 business days after the department receives a written request from the patient for the information. 10 This information shall include the records maintained by the department pursuant to subsection 11 (e). Notwithstanding the above, the department may, at the request of the law enforcement 12 agency, withhold for up to sixty (60) days following the conclusion of a law enforcement 13 investigation, the disclosure to the patient that information has been obtained pursuant to 14 subdivision (a)(3). 15 (d) A patient may request, from the dispensing pharmacy, correction of any inaccurate 16 information contained within the prescription drug monitoring database in accordance with the 17 procedure specified by § 5-37.3-5(c). 18 (e) The department shall, for the period of time that prescription information is 19 maintained, maintain records of the information disclosed through the prescription drug 20 monitoring database, including, but not limited to: 21 (1) The identity of each person who requests or receives information from the 22 prescription drug monitoring database and the organization, if any, the person represents; (2) The information released to each person or organization and the basis for its release 23 24 under subsection (a); and 25 (3) The dates the information was requested and provided. 26 (f) Prescription information contained within the prescription drug monitoring database 27 shall be removed no later than five (5) years from the date the information is entered into the 28 database. Records in existence prior to the enactment of this section shall be removed no later 29 than ten (10) years from the date the information is entered into the database. 30 (g) The department shall promptly notify any affected individual of an improper 31 disclosure of information from the prescription drug monitoring database or a breach in the 32 security of the prescription drug monitoring database that poses a significant risk of disclosure of 33 patient information to an unauthorized individual.

(h) At the time of signing a prescription that is required by the department to be entered

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- 1 into the prescription drug monitoring database, the prescribing practitioner shall inform the
- 2 patient in writing of the existence of the prescription drug monitoring database, the patient's right
- 3 to access their own prescription information, and the name and contact information of the agency
- 4 operating the program.

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- (i) No person shall access information in the prescription monitoring database except to the extent and for the purposes authorized by subsection (a).
- 7 (j) In any civil action allowing a violation of this chapter, the court may award damages,
- 8 including punitive damages, and reasonable attorneys' fees and costs to a prevailing plaintiff, and
- 9 injunctive and any other appropriate relief.
- 10 (k) Any pharmacist who, in his or her professional judgment, refuses to fill a prescription
- based on information contained within the prescription drug monitoring database shall inform the
- 12 prescribing physician within twenty-four (24) hours.
- 13 (l) All practitioners shall, as a condition of the initial registration or renewal of the
- 14 practitioner's authority to prescribe controlled substances, register with the prescription drug
- monitoring database maintained by the department of health.
- SECTION 2. This act shall take effect upon passage.

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EXPLANATION

BY THE LEGISLATIVE COUNCIL

OF

AN ACT

RELATING TO FOOD AND DRUGS - UNIFORMED CONTROLLED SUBSTANCE ACT

This act would authorize any vendor, agent, contractor, or designee who operates an electronic medical health record (EMR) or clinical management system to have access to the prescription drug monitoring program (PDMP).

This act would take effect upon passage.

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