

# NAMSDL



**National Alliance for Model State Drug Laws**

## **STATES THAT ALLOW PRACTITIONERS AND/OR PHARMACISTS TO DESIGNATE AN AUTHORIZED AGENT TO ACCESS THE PMP DATABASE**

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## Introduction

Each state determines by statute or regulation the persons or entities entitled to access or receive information in the prescription monitoring program database in that particular state. This memorandum sets out those states that allow access to or receipt of database information by agents or delegates of certain authorized users. This does not mean that if a particular state is not listed in this memorandum or the accompanying map that the state does not allow access to agents or delegates. If such persons fall within the definition of “practitioner” or “health care provider” in the state, he or she may qualify for access to the prescription monitoring program database. The following states either specifically include agents or delegates in the list of persons or entities entitled to access or NAMSDDL was informed by the administrator of the state prescription monitoring program that such persons are allowed access.

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Alabama  
§ 20-2-214

Code of Alabama (2013)  
Title 20. Food, Drugs, and Cosmetics.  
Chapter 2. Controlled Substances.  
Article 10. . Controlled Substances Prescription Database.

§ 20-2-214. Limited access to database permitted for certain persons or entities.

**The following persons or entities shall be permitted access to the information in the controlled substances database, subject to the limitations indicated below:**

(1) Authorized representatives of the certifying boards, provided, however, that access shall be limited to information concerning the licensees of the certifying board, however, authorized representatives from the Board of Medical Examiners may access the database to inquire about certified registered nurse practitioners (CRNPs), or certified nurse midwives (CNMs) that hold a Qualified Alabama Controlled Substances Registration Certificate (QACSC).

(2) A licensed practitioner approved by the department who has authority to prescribe, dispense, or administer controlled substances. The licensed practitioner's access shall be limited to information concerning himself or herself, registrants who possess a Qualified Alabama Controlled Substances Registration Certificate over whom the practitioner exercises physician supervision or with whom they have a joint practice agreement, a certified registered nurse practitioner and a certified nurse midwife with a Qualified Alabama Controlled Substances Registration Certificate over whom the practitioner exercises professional oversight and direction pursuant to an approved collaborative practice agreement, a current patient of the practitioner, and individuals seeking treatment from the practitioner. Practitioners shall have no requirement or obligation, under this article, to access or check the information in the controlled substances database prior to prescribing, dispensing, or administering medications or as part of their professional practice. However, the applicable licensing boards, in their discretion, may impose such a requirement or obligation by regulations.

**(3) A licensed physician approved by the department who has authority to prescribe, dispense, or administer controlled substances may designate up to two employees who may access the database on the physician's behalf.**

(4) A licensed certified registered nurse practitioner or a licensed certified nurse midwife approved by the department who is authorized to prescribe, administer, or dispense pursuant to a Qualified Alabama Controlled Substances Registration Certificate; provided, however, that such access shall be limited to information concerning a current or prospective patient of the registered nurse practitioner or certified nurse midwife.

(5) A licensed assistant to physician approved by the department who is authorized to prescribe, administer, or dispense pursuant to a Qualified Alabama Controlled Substances Registration Certificate; provided, however, that such access shall be limited to information concerning a current patient of the assistant to the physician or an individual seeking treatment from the assistant to physician.

(6) A licensed pharmacist approved by the department, provided, however, that such access is limited to information related to the patient or prescribing practitioner designated on a controlled substance prescription that a pharmacist has been asked to fill. Pharmacists shall have no requirement or obligation to access or check the information in the controlled substances database prior to dispensing or administering medications or as part of their professional practices.

(7) State and local law enforcement authorities as authorized under Section 20-2-91, and federal law enforcement authorities authorized to access prescription information upon application to the department accompanied by a declaration that probable cause exists for the use of the requested information.

(8) Employees of the department and consultants engaged by the department for operational and review purposes.

(9) The prescription drug monitoring program of any of the other states or territories of the United States, if recognized by the Alliance for Prescription Drug Monitoring Programs under procedures developed by the United States Department of Justice or the Integrated Justice Information Systems Institute or successor entity subject to or consistent with limitations for access prescribed by this chapter for the Alabama Prescription Drug Monitoring Program.

(10) Authorized representatives of the Alabama Medicaid Agency; provided, however, that access shall be limited to inquiries concerning possible misuse or abuse of controlled substances by Medicaid recipients.

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Delaware

16 § 4798 (eff. until March 1, 2014)

16 § 4798 (eff. March 1, 2014)

Delaware Code Annotated (2013)

Title 16. Health and Safety

Part IV. Food and Drugs

Chapter 47. Uniform Controlled Substances Act

Subchapter VII. Miscellaneous

§ 4798. The Delaware Prescription Monitoring Program

<Text of section effective until March 1, 2014>

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**(2) The Office of Controlled Substances may provide data in the prescription monitoring program in the form of a report to the following persons:**

**a. A prescriber, or other person authorized by the prescriber, or a dispenser, or other person authorized by the dispenser, who requests information and certifies that the requested information is for the purpose of providing medical or pharmaceutical treatment to a bona fide patient;**

b. An individual who requests the individual's own prescription monitoring information in accordance with procedures established pursuant to regulations;

c. A designated representative of any Board or Commission pursuant to § 8735(a) of Title 29 responsible for the licensure, regulation, or discipline of prescribers, dispensers or other persons authorized to prescribe, administer, or dispense controlled substances and who is involved in a bona fide specific investigation involving a designated person;

d. A local, state, or federal law enforcement or prosecutorial official engaged in the administration, investigation, or enforcement of the laws governing controlled substances and who is involved in a bona fide specific drug-related investigation in which a report of suspected criminal activity involving controlled substances by an identified suspect has been made, and provided that such information be relevant and material to such investigation, limited in scope to the extent reasonably practicable in light of the purpose for which the information is sought, and include identifying information only if non-identifying information could not be used;

e. The Delaware Department of Health and Social Services regarding Medicaid program recipients;

f. A properly convened grand jury pursuant to a subpoena properly issued for the records;

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g. Personnel of the Division of Professional Regulation for purposes of administration and enforcement of this section;

h. Qualified personnel for the purpose of bona fide research or education; however, data elements that would reasonably identify a specific recipient, prescriber or dispenser must be deleted or redacted from such information prior to disclosure; and further provided that, release of the information may be made only pursuant to a written agreement between qualified personnel and the Office of Controlled Substances in order to ensure compliance with this subsection.

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West's Delaware Code Annotated (2013)  
Title 16. Health and Safety  
Part IV. Food and Drugs  
Chapter 47. Uniform Controlled Substances Act  
Subchapter VII. Miscellaneous

§ 4798. The Delaware Prescription Monitoring Program

<Text of section effective March 1, 2014>

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(k) Prescription information submitted to the PMP is protected health information, not subject to public or open records law, and not subject to disclosure, except as otherwise provided in this section.

(l) The Office of Controlled Substances shall maintain procedures to ensure that the privacy and confidentiality of patients and patient information collected, recorded, transmitted, and maintained is not disclosed, except as provided for in this section.

(1) If there is reasonable cause to believe a violation of law or breach of professional standards may have occurred, the Office of Controlled Substances shall notify the appropriate law-enforcement or professional licensure, certification, or regulatory agency or entity and shall provide prescription information required for an investigation.

**(2) The Office of Controlled Substances may provide data in the prescription monitoring program in the form of a report to the following persons:**

**a. A prescriber, or other person authorized by the prescriber, or a dispenser, or other person authorized by the dispenser, who requests information and certifies that the**

**requested information is for the purpose of providing medical or pharmaceutical treatment to a bona fide patient;**

- b. An individual who requests the individual's own prescription monitoring information in accordance with procedures established pursuant to regulations;
- c. A designated representative of any Board or Commission pursuant to § 8735(a) of Title 29 responsible for the licensure, regulation, or discipline of prescribers, dispensers or other persons authorized to prescribe, administer, or dispense controlled substances and who is involved in a bona fide specific investigation involving a designated person;
- d. A local, state, or federal law-enforcement or prosecutorial official engaged in the administration, investigation, or enforcement of the laws governing controlled substances and who is involved in a bona fide specific drug-related investigation in which a report of suspected criminal activity involving controlled substances by an identified suspect has been made, and provided that such information be relevant and material to such investigation, limited in scope to the extent reasonably practicable in light of the purpose for which the information is sought, and include identifying information only if nonidentifying information could not be used;
- e. The Delaware Department of Health and Social Services regarding Medicaid program recipients;
- f. A properly convened grand jury pursuant to a subpoena properly issued for the records;
- g. Personnel of the Division of Professional Regulation for purposes of administration and enforcement of this section;
- h. A licensed chemical dependency professional or licensed professional counselor of mental health who requests information and certifies that the requested information is for a patient enrolled in a substance abuse treatment program receiving treatment from, or under the direction of, the chemical dependency professional or professional counselor of mental health;
- i. The Chief Medical Examiner or licensed physician designee who requests information and certifies the request is for the purpose of investigating the death of an individual;
- j. Qualified personnel for the purpose of bona fide research or education; however, data elements that would reasonably identify a specific recipient, prescriber or dispenser must be deleted or redacted from such information prior to disclosure; and further provided that, release of the information may be made only pursuant to a written agreement between qualified personnel and the Office of Controlled Substances in order to ensure compliance with this subsection.

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## Idaho

Per the state PMP representative, Idaho will allow a prescriber to have one designated agent who must be employed at the prescriber's place of business and who must have their own login and password.

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## Indiana

### § 35-48-7-11.1

Annotated Indiana Code (2013)

Title 35. Criminal Law and Procedure

Article 48. Controlled Substances

Chapter 7. Central Repository for Controlled Substances Data

#### § 35-48-7-11.1 INSPECT program; confidentiality

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**(d) Except as provided in subsections (e) and (f), the board may release confidential information described in subsection (a) to the following persons:**

(1) A member of the board or another governing body that licenses practitioners and is engaged in an investigation, an adjudication, or a prosecution of a violation under any state or federal law that involves a controlled substance.

(2) An investigator for the consumer protection division of the office of the attorney general, a prosecuting attorney, the attorney general, a deputy attorney general, or an investigator from the office of the attorney general, who is engaged in:

(A) an investigation;

(B) an adjudication; or

(C) a prosecution;

of a violation under any state or federal law that involves a controlled substance.

(3) A law enforcement officer who is an employee of:

(A) a local, state, or federal law enforcement agency; or

(B) an entity that regulates controlled substances or enforces controlled substances rules or laws in another state;

that is certified to receive information from the INSPECT program.

**(4) A practitioner or practitioner's agent certified to receive information from the INSPECT program.**

(5) A controlled substance monitoring program in another state with which Indiana has established an interoperability agreement.

(6) The state toxicologist.

(7) A certified representative of the Medicaid retrospective and prospective drug utilization review program.

(8) A substance abuse assistance program for a licensed health care provider who:

(A) has prescriptive authority under IC 25; and

(B) is participating in the assistance program.

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Iowa  
§ 124.553  
ADC 657-37.2(124)  
ADC 657-37.4(124)

Iowa Code Annotated (2013)  
Title IV. Public Health  
Subtitle 1. Alcoholic Beverages and Controlled Substances  
Chapter 124. Controlled Substances  
Division VI. Drug Prescribing and Dispensing--Information Program

§ 124.553. Information access

**1. The board may provide information from the program to the following:**

a. (1) A pharmacist or prescribing practitioner who requests the information and certifies in a form specified by the board that it is for the purpose of providing medical or pharmaceutical care to a patient of the pharmacist or prescribing practitioner. **A pharmacist or a prescribing practitioner may delegate program information access to another authorized individual or agent only if that individual or agent registers for program information access, pursuant to board rules, as an agent of the pharmacist or prescribing practitioner.** Board rules shall identify the qualifications for a pharmacist's or prescribing practitioner's agent and shall limit the number of agents to whom each pharmacist or prescribing practitioner may delegate program information access.

(2) Notwithstanding subparagraph (1), a prescribing practitioner may delegate program information access to another licensed health care professional in emergency situations where the patient would be placed in greater jeopardy if the prescribing practitioner was required to access the information personally.

b. An individual who requests the individual's own program information in accordance with the procedure established in rules of the board and advisory council adopted under section 124.554.

c. Pursuant to an order, subpoena, or other means of legal compulsion for access to or release of program information that is issued based upon a determination of probable cause in the course of a specific investigation of a specific individual.

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Iowa Administrative Code (2013)  
Agency 657 Pharmacy Board  
Chapter 37 Iowa Prescription Monitoring Program

657-37.2(124) Definitions.

As used in this chapter:

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**“Practitioner’s agent” means a health care professional who is employed by or under the direct supervision of a health care practitioner and who is authorized by the practitioner to access PMP information as provided in subrule 37.4(1).**

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Iowa Administrative Code (2013)  
Agency 657 Pharmacy Board  
Chapter 37 Iowa Prescription Monitoring Program

657-37.4(124) Access to database information.

All information contained in the PMP database, including prescription information submitted for inclusion in the PMP database and records of requests for PMP information, shall be privileged and strictly confidential and not subject to public or open records laws. The board, council, and PMP administrator shall maintain procedures to ensure the privacy and confidentiality of patients, prescribers, dispensers, practitioners, practitioners’ agents, and patient information collected, recorded, transmitted, and maintained in the PMP database and to ensure that program information is not disclosed to persons except as provided in this rule.

**37.4(1) Prescribers and pharmacists. A health care practitioner authorized to prescribe or dispense controlled substances may obtain PMP information regarding the practitioner’s patient, or a patient seeking treatment from the practitioner, for the purpose of providing patient health care. A practitioner may authorize no more than three health care professionals to act as the practitioner’s agents for the purpose of requesting PMP information regarding a practitioner’s patients.**

**a. Prior to being granted access to PMP information, a practitioner or a practitioner’s agent shall submit an individual request for registration and program access. A practitioner or a practitioner’s agent with Internet access may register via a secure Web site established by the board for that purpose. A practitioner without Internet access shall submit a written registration request on a form provided by the PMP administrator. A practitioner without Internet access shall not authorize a practitioner’s agent to register for or to access PMP information on behalf of the practitioner. The PMP administrator shall**

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**take reasonable steps to verify the identity of a practitioner or practitioner's agent and to verify a practitioner's credentials prior to providing a practitioner with a secure login and initial password. Each practitioner or practitioner's agent registered to access PMP information shall securely maintain and use the login and password assigned to the individual practitioner or practitioner's agent. Except in an emergency when the patient would be placed in greater jeopardy by restricting PMP information access to the practitioner or practitioner's agent, a registered practitioner shall not share the practitioner's secure login and password information and shall not delegate PMP information access to another health care practitioner or to an unregistered agent. A registered practitioner's agent shall not delegate PMP information access to another individual.**

**b. A practitioner or practitioner's agent with Internet access may submit a request for PMP information via a secure Web site established by the board for that purpose. The requested information shall be provided to the requesting practitioner or practitioner's agent in a format established by the board and shall be delivered via the secure Web site.**

c. A practitioner without Internet access may submit to the PMP administrator a written request for PMP information via mail or facsimile transmission. The written request shall be in a format established by the board and shall be signed by the requesting practitioner. Prior to processing a written request for PMP information, the PMP administrator shall take reasonable steps to verify the request, which may include but not be limited to a telephone call to the practitioner at a telephone number known to be the number for the practitioner's practice.

d. A practitioner who requests and receives PMP information consistent with the requirements and intent of these rules may provide that information to another practitioner who is involved in the care of the patient who is the subject of the information. Information from the PMP database remains privileged and strictly confidential. Such disclosures among practitioners shall be consistent with these rules and federal and state laws regarding the confidentiality of patient information. The information shall be used for medical or pharmaceutical care purposes.

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Kansas  
ADC 68-21-5

Kansas Administrative Regulations (2013)  
Agency 68. Board of Pharmacy  
Article 21. Prescription Monitoring Program

68-21-5 Access to information.

All requests for, uses of, and disclosures of prescription monitoring information by authorized persons shall meet the requirements of K.S.A. 65-1685, and amendments thereto, and this article.

...

(c) By prescribers.

**(1) Any prescriber or health care practitioner authorized by a prescriber may obtain any program information relating to a patient under the prescriber's care, in accordance with this regulation and K.S.A. 65-1685 and amendments thereto. The information shall be provided in a format established by the board, which may include delivery by electronic means, facsimile, or telephone.**

**(2) Each prescriber or health care practitioner authorized by a prescriber who seeks access to program information shall submit a written request to the board by mail, hand delivery, or electronic means in a manner established by the board, using authentication. If the authentication is lost or missing or the security of the authentication is compromised, the prescriber shall cause the board to be notified by telephone and in writing as soon as reasonably possible. Information regarding more than one patient may be submitted in a single request.**

Each request shall be submitted in a format established by the board and shall include the following elements for each patient:

- (A) The patient's name and birth date;
- (B) if known to the prescriber, the patient's address and telephone number;
- (C) the time period for which information is being requested;
- (D) the prescriber's name;
- (E) the name and address of the prescriber's medical practice;

(F) the prescriber identification number; and

(G) the prescriber's signature.

(3) The authentication and identity of the dispenser shall be verified before allowing access to any program information.

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Kentucky  
§ 218A.202

Baldwin's Kentucky Revised Statutes Annotated (2013)  
Title XVIII. Public Health  
Chapter 218A. Controlled Substances

§ 218A.202 Electronic system for monitoring controlled substances; required registration and reporting; penalty for illegal use of system; pilot or continuing project; continuing education programs; reports of failure to comply with section; administrative regulations

...

**(6) The Cabinet for Health and Family Services shall only disclose data to persons and entities authorized to receive that data under this section. Disclosure to any other person or entity, including disclosure in the context of a civil action where the disclosure is sought either for the purpose of discovery or for evidence, is prohibited unless specifically authorized by this section. The Cabinet for Health and Family Services shall be authorized to provide data to:**

(a) A designated representative of a board responsible for the licensure, regulation, or discipline of practitioners, pharmacists, or other person who is authorized to prescribe, administer, or dispense controlled substances and who is involved in a bona fide specific investigation involving a designated person;

(b) Employees of the Office of the Inspector General of the Cabinet for Health and Family Services who have successfully completed training for the electronic system and who have been approved to use the system, Kentucky Commonwealth's attorneys and assistant Commonwealth's attorneys, county attorneys and assistant county attorneys, a peace officer certified pursuant to KRS 15.380 to 15.404, a certified or full-time peace officer of another state, or a federal peace officer whose duty is to enforce the laws of this Commonwealth, of another state, or of the United States relating to drugs and who is engaged in a bona fide specific investigation involving a designated person;

(c) A state-operated Medicaid program in conformity with subsection (7) of this section;

(d) A properly convened grand jury pursuant to a subpoena properly issued for the records;

**(e) A practitioner or pharmacist, or employee of the practitioner's or pharmacist's practice acting under the specific direction of the practitioner or pharmacist, who requests information and certifies that the requested information is for the purpose of:**

**1. Providing medical or pharmaceutical treatment to a bona fide current or prospective patient; or**

**2. Reviewing and assessing the individual prescribing or dispensing patterns of the practitioner or pharmacist or to determine the accuracy and completeness of information contained in the monitoring system;**

**(f) The chief medical officer of a hospital or long-term-care facility, an employee of the hospital or long-term-care facility as designated by the chief medical officer and who is working under his or her specific direction, or a physician designee if the hospital or facility has no chief medical officer, if the officer, employee, or designee certifies that the requested information is for the purpose of providing medical or pharmaceutical treatment to a bona fide current or prospective patient or resident in the hospital or facility;**

(g) In addition to the purposes authorized under paragraph (a) of this subsection, the Kentucky Board of Medical Licensure, for any physician who is:

1. Associated in a partnership or other business entity with a physician who is already under investigation by the Board of Medical Licensure for improper prescribing or dispensing practices;

2. In a designated geographic area for which a trend report indicates a substantial likelihood that inappropriate prescribing or dispensing may be occurring; or

3. In a designated geographic area for which a report on another physician in that area indicates a substantial likelihood that inappropriate prescribing or dispensing may be occurring in that area;

(h) In addition to the purposes authorized under paragraph (a) of this subsection, the Kentucky Board of Nursing, for any advanced practice registered nurse who is:

1. Associated in a partnership or other business entity with a physician who is already under investigation by the Kentucky Board of Medical Licensure for improper prescribing or dispensing practices;

2. Associated in a partnership or other business entity with an advanced practice registered nurse who is already under investigation by the Board of Nursing for improper prescribing practices;

3. In a designated geographic area for which a trend report indicates a substantial likelihood that inappropriate prescribing or dispensing may be occurring; or

4. In a designated geographic area for which a report on a physician or another advanced practice registered nurse in that area indicates a substantial likelihood that inappropriate prescribing or dispensing may be occurring in that area;

(i) A judge or a probation or parole officer administering a diversion or probation program of a criminal defendant arising out of a violation of this chapter or of a criminal defendant who is documented by the court as a substance abuser who is eligible to participate in a court-ordered drug diversion or probation program; or

(j) A medical examiner engaged in a death investigation pursuant to KRS 72.026.

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Louisiana  
40 § 1007

West's Louisiana Statutes Annotated (2013)  
Louisiana Revised Statutes  
Title 40. Public Health and Safety  
Chapter 4. Food and Drugs  
Part X-A. Prescription Monitoring Program

§ 1007. Access to prescription monitoring information

A. Except as provided in Subsections C, D, E, F, G, H, and I of this Section, prescription monitoring information submitted to the board shall be protected health information, not subject to public or open records law, including but not limited to R.S. 44:1 et seq., and not subject to disclosure. Prescription monitoring information shall not be available for civil subpoena nor shall such information be disclosed, discoverable, or compelled to be produced in any civil proceeding nor shall such records be deemed admissible as evidence in any civil proceeding for any reason. Notwithstanding this provision, law enforcement and professional licensing, certification, or regulatory agencies may utilize prescription monitoring information in the course of any investigation and subsequent criminal and administrative proceedings, but only in accordance with federal and state law and the requirements of this Part.

B. The board shall maintain procedures to ensure that the privacy and confidentiality of patients and patient information collected, recorded, transmitted, and maintained is not disclosed to persons or entities except as in Subsections C, D, E, F, G, H, and I of this Section.

C. The board shall review the prescription monitoring information. If there is reasonable suspicion to believe a breach of professional or occupational standards may have occurred, the board shall notify the appropriate professional licensing agency with jurisdiction over prescribers or dispensers and shall provide prescription monitoring information required for an investigation.

D. The board shall provide prescription monitoring information to public or private entities, whether located in or outside of the state, for public research, policy, or educational purposes, but only after removing information that identifies or could be reasonably used to identify prescribers, dispensers, and individual patients or persons who received prescriptions from prescribers.

**E. The following persons, after successful completion of the educational courses identified in R.S. 40:1008, may access prescription monitoring information at no cost and in the same or similar manner, and for the same or similar purposes, as those persons are authorized to access similar protected health information under federal and state law and regulation:**

**(1) Persons authorized to prescribe or dispense controlled substances or drugs of concern, or their delegates as defined by rule, for the purpose of providing medical or pharmaceutical care for their patients, or for verifying their prescribing records.**

(2) Designated representatives from the professional licensing, certification, or regulatory agencies of this state or another state charged with administrative oversight of those professionals engaged in the prescribing or dispensing of controlled substances or other drugs of concern.

(3) Designated representatives from the Louisiana Medicaid program regarding Medicaid program recipients.

(4) Designated representatives of the board and any vendor or contractor establishing or maintaining the prescription monitoring program.

F. The board may provide a report containing prescription monitoring information upon application of local, state, out-of-state, and federal law enforcement or prosecutorial officials engaged in the administration, investigation, or enforcement of the laws governing controlled substances or other drugs of concern in compliance with and as limited by the relevant requirements of any of the following:

(1) A court order or court-ordered warrant, or a subpoena or summons issued by a judicial officer.

(2) A grand jury subpoena.

(3) An administrative request, including an administrative subpoena or summons, a civil or an authorized investigative demand, or similar process authorized under law, provided by law enforcement to the board, and further, provided all of the following:

(a) The information sought is relevant and material to a legitimate law enforcement inquiry.

(b) The request is specific and limited in scope to the extent reasonably practicable in light of the purpose for which the information is sought.

(c) De-identified information, or limited information that does not identify or could not reasonably lead to the identification of an individual patient, could not reasonably be used.

G. The board may provide prescription monitoring information in response to queries from prescription monitoring programs located in other states, through its participation in a secure interstate data exchange system. However, the board shall not provide prescription monitoring information to prescription monitoring programs located in other states unless the laws of the state receiving the information provide at a minimum both of the following:

(1) That the prescription monitoring information is protected health information, not subject to the Public Records Law, and not subject to disclosure.

(2) That the prescription monitoring information shall not be subject to civil subpoena, nor shall such information be disclosed, discoverable, or compelled to be produced in any civil proceeding, nor shall such records be deemed admissible as evidence in any civil proceeding for any reason.

H. The board may provide prescription monitoring information to authorized users of the prescription monitoring program via a state health information exchange or other third party conduit that has been approved by the board.

I. The board may provide prescription monitoring information to an individual who requests his personal prescription monitoring information in accordance with procedures established by board regulation.

J. The board and the advisory council shall be immune from civil liability arising from inaccuracy of any of the information submitted to the board pursuant to this Part.

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## Maine

ADC 14-118, Ch. 11, § 7

Code of Maine Rules (2013)

14. Department of Behavioral and Developmental Services

118. Office of Substance Abuse

Chapter 11. Rules Governing The Controlled Substances Prescription Monitoring Program

### Sec. 7. Access to Prescription Monitoring Information

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#### 2. By dispensers

**A. A dispenser, or a licensed pharmacy technician authorized by a supervising pharmacist, may obtain any prescription monitoring information insofar as the information relates to a customer of the dispenser seeking to have a prescription filled. The information shall be provided in a format established by the Office, which may include, but is not limited to, delivery by electronic means, facsimile transmission, or telephonic communication.**

B. A dispenser who seeks access to the information described above must register as a data requester in a manner specified by the Monitor or the Office. The Office or Monitor shall issue credentials to authorized dispensers. Dispensers may use these credentials to access the online database and submit requests. If the credentials issued by the Office are lost, missing, or the security of the credentials is compromised, the dispenser shall cause the Office or Monitor to be notified by telephone and in writing as soon as reasonably possible. Information regarding more than one customer may be submitted in a single request. Requests shall be in a format established by the Office or the Monitor and shall contain at least, but not limited to, the following elements for each customer:

- 1) The name and date of birth of the customer; and
- 2) The time period for which information is being requested.

C. The Office or the Monitor shall take reasonable steps to verify each registration, such as, but not limited to, making a telephone call to the dispenser or to an agent of the dispenser at a telephone number known to belong to the dispenser's place of business.

#### 3. By prescribers

**A. A prescriber, or any staff member duly authorized by a prescriber and the Office, may obtain any prescription monitoring information insofar as the information relates to a**

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**patient under the prescriber's care. The information shall be provided in a format established by the Office, which may include, but is not limited to delivery by electronic means, facsimile transmission, or telephonic communication.**

B. A prescriber, or any staff member duly authorized by a prescriber and the Office, who seeks access to the information described above must register as a data requester in a manner specified by the Monitor or the Office. The Office or Monitor shall issue credentials to authorized prescribers or their designees. Data requesters may use these credentials to access the online database and submit requests. If the credentials issued by the Office are lost, missing, or the security of the credentials is compromised, the data requester shall cause the Office or Monitor to be notified by telephone and in writing as soon as reasonably possible. Requests shall be in a format established by the Office or the Monitor and shall contain at least, but not limited to, the following elements for each patient:

- 1) The name and date of birth of the patient; and
- 2) The time period for which information is being requested.

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## Maryland

Health-General § 21-2A-06 (eff. until Oct. 1, 2013)

Health-General § 21-2A-06 (eff. Oct. 1, 2013)

ADC 10.47.07.02

ADC 10.47.07.04

West's Annotated Code of Maryland (2013)

Health--General

Title 21. Food, Drugs, and Cosmetics

Subtitle 2A. Prescription Drug Monitoring Program

§ 21-2A-06. Confidentiality of prescription monitoring data

<Text of section effective until October 1, 2013>

Data not subject to discovery or subpoena

(a) Prescription monitoring data:

(1) Are confidential and privileged, and not subject to discovery, subpoena, or other means of legal compulsion in civil litigation;

(2) Are not public records; and

(3) Except as provided in subsections (b) and (d) of this section or as otherwise provided by law, may not be disclosed to any person.

Allowable disclosure of prescription monitoring data

**(b) The Program shall disclose prescription monitoring data, in accordance with regulations adopted by the Secretary, to:**

**(1) A prescriber, or a licensed health care practitioner authorized by the prescriber, in connection with the medical care of a patient;**

**(2) A dispenser, or a licensed health care practitioner authorized by the dispenser, in connection with the dispensing of a monitored prescription drug;**

(3) A federal law enforcement agency or a State or local law enforcement agency, on issuance of a subpoena, for the purpose of furthering an existing bona fide individual investigation;

- (4) A licensing entity, on issuance of an administrative subpoena voted on by a quorum of the board of the licensing entity, for the purposes of furthering an existing bona fide individual investigation;
- (5) A rehabilitation program under a health occupations board, on issuance of an administrative subpoena;
- (6) A patient with respect to prescription monitoring data about the patient;
- (7) Subject to subsection (g) of this section, the authorized administrator of another state's prescription drug monitoring program;
- (8) The following units of the Department, on approval of the Secretary, for the purpose of furthering an existing bona fide individual investigation:
  - (i) The Office of the Chief Medical Examiner;
  - (ii) The Maryland Medical Assistance Program;
  - (iii) The Office of the Inspector General; and
  - (iv) The Office of Health Care Quality; or
- (9) The technical advisory committee established under § 21-2A-07 of this subtitle for the purposes set forth in subsection (c) of this section.

#### Review of requests for information

- (c) Before the Program discloses information under subsection (b)(3), (4), (5), (7), or (8) of this section, the technical advisory committee to the Program shall:
  - (1) Review the requests for information;
  - (2) Provide clinical guidance and interpretation of the information requested to the Secretary to assist in the Secretary's decision on how to respond to a judicial subpoena, administrative subpoena, or other request; and
  - (3) Provide clinical guidance and interpretation of the information requested to the authorized recipient of the information.

#### Persons who receive prescription monitoring data prohibited from disclosure

- (d) Except as provided by regulations adopted by the Secretary, a person who receives prescription monitoring data from the Program may not disclose the data.

## Disclosure of data for research, analysis, public reporting, and education

(e)(1) In addition to the disclosures required under subsection (b) of this section, the Program may disclose prescription monitoring data for research, analysis, public reporting, and education:

(i) After redaction of all information that could identify a patient, prescriber, dispenser, or any other individual; and

(ii) In accordance with regulations adopted by the Secretary.

(2) The Secretary may require submission of an abstract explaining the scope and purpose of the research, analysis, public reporting, or education before disclosing prescription monitoring data under this subsection.

## Injunctive relief

(f) The Office of the Attorney General may seek appropriate injunctive or other relief to maintain the confidentiality of prescription monitoring data as required under this section.

## Prescription monitoring data shared with other states

(g) The Program may provide prescription monitoring data to another state's prescription drug monitoring program only if the other state's prescription drug monitoring program agrees to use the prescription monitoring data in a manner consistent with the provisions of this subtitle.

## Request and receipt of prescription monitoring data from other states

(h) The Program may:

(1) Request and receive prescription monitoring data from another state's prescription drug monitoring program and use the prescription monitoring data in a manner consistent with the provisions of this subtitle; and

(2) Develop the capability to transmit prescription monitoring data to and receive prescription monitoring data from other prescription drug monitoring programs employing the standards of interoperability.

## Written agreements with other states

(i) The Program may enter into written agreements with other states' prescription drug monitoring programs for the purpose of establishing the terms and conditions for sharing prescription monitoring data under this section.

## Clinical practice standards

(j) Prescription monitoring data may not be used as the basis for imposing clinical practice standards.

West's Annotated Code of Maryland (2013)  
Health--General  
Title 21. Food, Drugs, and Cosmetics  
Subtitle 2A. Prescription Drug Monitoring Program

§ 21-2A-06. Confidentiality of prescription monitoring data

<Text of section effective October 1, 2013>

Data not subject to discovery or subpoena

(a) Prescription monitoring data:

(1) Are confidential and privileged, and not subject to discovery, subpoena, or other means of legal compulsion in civil litigation;

(2) Are not public records; and

(3) Except as provided in subsections (b) and (d) of this section or as otherwise provided by law, may not be disclosed to any person.

Allowable disclosure of prescription monitoring data

**(b) The Program shall disclose prescription monitoring data, in accordance with regulations adopted by the Secretary, to:**

**(1) A prescriber, or a licensed health care practitioner authorized by the prescriber, in connection with the medical care of a patient;**

**(2) A dispenser, or a licensed health care practitioner authorized by the dispenser, in connection with the dispensing of a monitored prescription drug;**

(3) A federal law enforcement agency or a State or local law enforcement agency, on issuance of a subpoena, for the purpose of furthering an existing bona fide individual investigation;

(4) A licensing entity, on issuance of an administrative subpoena voted on by a quorum of the board of the licensing entity, for the purposes of furthering an existing bona fide individual investigation;

(5) A rehabilitation program under a health occupations board, on issuance of an administrative subpoena;

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- (6) A patient with respect to prescription monitoring data about the patient;
- (7) Subject to subsection (g) of this section, the authorized administrator of another state's prescription drug monitoring program;
- (8) The following units of the Department, on approval of the Secretary, for the purpose of furthering an existing bona fide individual investigation:
  - (i) The Office of the Chief Medical Examiner;
  - (ii) The Maryland Medical Assistance Program;
  - (iii) The Office of the Inspector General;
  - (iv) The Office of Health Care Quality; and
  - (v) The Division of Drug Control; or
- (9) The technical advisory committee established under § 21-2A-07 of this subtitle for the purposes set forth in subsection (c) of this section.

#### Review of requests for information

- (c) Before the Program discloses information under subsection (b)(3), (4), (5), (7), or (8) of this section, the technical advisory committee to the Program shall:
  - (1) Review the requests for information;
  - (2) Provide clinical guidance and interpretation of the information requested to the Secretary to assist in the Secretary's decision on how to respond to a judicial subpoena, administrative subpoena, or other request; and
  - (3) Provide clinical guidance and interpretation of the information requested to the authorized recipient of the information.

#### Persons who receive prescription monitoring data prohibited from disclosure

- (d) Except as provided by regulations adopted by the Secretary, a person who receives prescription monitoring data from the Program may not disclose the data.

#### Disclosure of data for research, analysis, public reporting, and education

- (e)(1) In addition to the disclosures required under subsection (b) of this section, the Program may disclose prescription monitoring data for research, analysis, public reporting, and education:

(i) After redaction of all information that could identify a patient, prescriber, dispenser, or any other individual; and

(ii) In accordance with regulations adopted by the Secretary.

(2) The Secretary may require submission of an abstract explaining the scope and purpose of the research, analysis, public reporting, or education before disclosing prescription monitoring data under this subsection.

#### Injunctive relief

(f) The Office of the Attorney General may seek appropriate injunctive or other relief to maintain the confidentiality of prescription monitoring data as required under this section.

#### Prescription monitoring data shared with other states

(g) The Program may provide prescription monitoring data to another state's prescription drug monitoring program only if the other state's prescription drug monitoring program agrees to use the prescription monitoring data in a manner consistent with the provisions of this subtitle.

#### Request and receipt of prescription monitoring data from other states

(h) The Program may:

(1) Request and receive prescription monitoring data from another state's prescription drug monitoring program and use the prescription monitoring data in a manner consistent with the provisions of this subtitle; and

(2) Develop the capability to transmit prescription monitoring data to and receive prescription monitoring data from other prescription drug monitoring programs employing the standards of interoperability.

#### Written agreements with other states

(i) The Program may enter into written agreements with other states' prescription drug monitoring programs for the purpose of establishing the terms and conditions for sharing prescription monitoring data under this section.

#### Clinical practice standards

(j) Prescription monitoring data may not be used as the basis for imposing clinical practice standards.

Code of Maryland Regulations (2013)  
Title 10 Department of Health and Mental Hygiene  
Subtitle 47 Alcohol and Drug Abuse Administration  
Chapter 07 Prescription Drug Monitoring Program

.02 Definitions.

A. In this chapter, the following terms have the meanings indicated.

B. Terms Defined.

**(1) “Authorized licensed health care practitioner” means a licensed health care practitioner who is authorized by a prescriber or dispenser to access prescription monitoring data in connection with the medical care of a patient to whom the prescriber prescribes or the dispenser dispenses a monitored prescription drug.**

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Code of Maryland Regulations (2013)  
Title 10 Department of Health and Mental Hygiene  
Subtitle 47 Alcohol and Drug Abuse Administration  
Chapter 07 Prescription Drug Monitoring Program

.04 Disclosure of Prescription Monitoring Data.

**A. Registration of a Prescriber, a Dispenser, or an Authorized Licensed Health Care Practitioner to Request Prescription Monitoring Data.**

**(1) A prescriber, a dispenser, or an authorized licensed health care practitioner shall register with the Department or its agent, in a manner specified by the Department, in order to request disclosure of or otherwise access prescription monitoring data.**

**(2) The Department or its agent shall:**

**(a) Establish procedures to authenticate a prescriber, a dispenser, or an authorized licensed health care practitioner in accordance with Health-General Article, §21-2A-06(b)(1)-(2), Annotated Code of Maryland; and**

**(b) Issue credentials to a prescriber, a dispenser, or an authorized licensed health care practitioner that can be used to request disclosure of or otherwise access prescription monitoring data electronically.**

**(3) If the credentials issued to a registrant are lost, stolen, or otherwise compromised, the registrant shall notify the Department or its agent, by a method approved by the Department, as soon as reasonably possible.**

**(4) A prescriber or dispenser who authorizes the registration of a licensed health care practitioner to request disclosure of or otherwise access prescription monitoring data shall:**

**(a) Make every reasonable effort, including regularly reviewing and auditing any available logs of system access and use, to ensure the authorized licensed health care practitioner is requesting disclosure of, redisclosing, or otherwise accessing prescription monitoring data in clear compliance with Health-General Article, Title 21, Subtitle 2A, Annotated Code of Maryland, and all other State and federal laws and regulations governing the security and confidentiality of protected health information and personal medical records;**

**(b) Immediately notify the Department or its agent, by a method approved by the Department, as well as the licensing entity responsible for licensing, certifying, or registering the authorized licensed health care practitioner, if the prescriber or dispenser believes that the confidentiality of prescription monitoring data or the security of the Program has been compromised by an authorized licensed health care practitioner; and**

**(c) Immediately notify the Department or its agent, by a method approved by the Department, of any requested change in the registration status of an authorized licensed health care practitioner, including if that authorized licensed health care practitioner is no longer employed by or practicing under the authority of the prescriber or dispenser.**

#### **B. Disclosure of Prescription Monitoring Data to a Prescriber, a Dispenser, or an Authorized Licensed Health Care Practitioner.**

**(1) Upon request from a prescriber or a licensed health care practitioner authorized by a prescriber, the Program shall disclose patient-specific prescription monitoring data provided that the request is made solely for the purpose of the medical care or treatment of the patient about whom prescription monitoring data is being requested.**

**(2) Upon request from a prescriber, the Program may provide a report containing prescription monitoring data on all monitored prescription drugs dispensed pursuant to the prescriber's prescriptions, provided that the request is submitted on a form or in a manner approved by the Department.**

**(3) Upon request from a dispenser or a licensed health care practitioner authorized by a dispenser, the Program shall disclose patient-specific prescription monitoring data provided that the request is made pursuant to a dispenser's responsibility to perform due diligence and exercise professional judgment when presented with a prescription to dispense a monitored prescription drug for use by the patient about whom prescription monitoring data is being requested.**



**(4) The Department or its agent shall make available the electronic means by which a prescriber, a dispenser, or an authorized licensed health care practitioner may request disclosure of or otherwise access patient-specific prescription monitoring data.**

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Massachusetts  
94C § 24A

Massachusetts General Laws Annotated (2013)  
Part I. Administration of the Government (Ch. 1-182)  
Title XV. Regulation of Trade (Ch. 93-110H)  
Chapter 94C. Controlled Substances Act

§ 24A. Electronic monitoring of the prescribing and dispensing of controlled substances and certain additional drugs

(a)(1) The department shall establish and maintain an electronic system to monitor the prescribing and dispensing of all schedule II to V, inclusive, controlled substances and certain additional drugs by all professionals licensed to prescribe or dispense such substances. For the purposes of this section, “additional drugs” shall mean substances determined by the department to carry a bona fide potential for abuse.

(2) The department shall enter into reciprocal agreements with other states that have compatible prescription drug monitoring programs to share prescription drug monitoring information among the states.

(b) The requirements of this section shall not apply to the dispensing of controlled substances to inpatients in a hospital.

(c) For the purposes of monitoring the prescribing and dispensing of all schedule II to V, inclusive, controlled substances and additional drugs, as authorized in subsection (a), the department shall promulgate regulations including, but not limited to, (1) a requirement that each pharmacy that delivers a schedule II to V, inclusive, controlled substance or a substance classified as an additional drug by the department to the ultimate user shall submit to the department, by electronic means, information regarding each prescription dispensed for a drug included under subsection (a); and (2) a requirement that each pharmacy collects and reports, for each prescription dispensed for a drug under subsection (a), a customer identification number and other information associated with the customer identification number, as specified by the department. Each pharmacy shall submit the information in accordance with transmission methods and frequency requirements promulgated by the department; provided, however, that the information shall be submitted at least once every 7 days. The department may issue a waiver to a pharmacy that is unable to submit prescription information by electronic means. The waiver shall permit the pharmacy to submit prescription information by other means promulgated by the department; provided, however, that all information required in this section is submitted in this alternative format.

**The department, in consultation with all relevant licensing authorities, shall promulgate regulations** that require participants to utilize the prescription monitoring program prior to

seeing a new patient, including circumstances where participants would not be required to utilize the prescription monitoring program prior to seeing a new patient; a requirement that pharmacists be trained in the use of the prescription monitoring program as part of the continuing education requirements mandated for licensure by the board of registration in pharmacy, under section 24A of chapter 112 and **a requirement that allows authorized support staff to use the prescription monitoring program on behalf of a registered participant.**

(d) Prescription information submitted to the department under this section shall be confidential and exempt from disclosure under clause Twenty-sixth of section 7 of chapter 4 and chapter 66. The department shall maintain procedures to ensure that the privacy and confidentiality of patients and patient information collected, recorded, transmitted and maintained is not disclosed to persons except as provided for in this chapter.

(e) The department shall review the prescription and dispensing monitoring information. If there is reasonable cause to believe a violation of law or breach of professional standards may have occurred, the department shall notify the appropriate law enforcement or professional licensing, certification or regulatory agency or entity and provide prescription information required for an investigation.

(f) The department shall, upon request, provide data from the prescription monitoring program to the following:--

(1) persons authorized to prescribe or dispense controlled substances, for the purpose of providing medical or pharmaceutical care for their patients;

(2) individuals who request their own prescription monitoring information in accordance with procedures established under chapter 66A;

(3) persons authorized to act on behalf of state boards and regulatory agencies that supervise or regulate a profession that may prescribe controlled substances; provided, however, that the data request is in connection with a bona fide specific controlled substance or additional drug-related investigation;

(4) local, state and federal law enforcement or prosecutorial officials working with the executive office of public safety engaged in the administration, investigation or enforcement of the laws governing prescription drugs; provided, however, that the data request is in connection with a bona fide specific controlled substance or additional drug-related investigation;

(5) personnel of the executive office of health and human services regarding Medicaid program recipients; provided, however that the data request is in connection with a bona fide specific controlled substance or additional drug-related investigation; or

(6) personnel of the United States attorney, office of the attorney general or a district attorney; provided, however, that the data request is in connection with a bona fide specific controlled substance or additional drug related investigation.

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(g) The department may, at its initiative, provide data from the prescription monitoring program to practitioners in accordance with section 24.

(h) The department may provide de-identified, aggregate information to a public or private entity for statistical research or educational purposes.

(i) The department may contract with another agency or with a private vendor, as necessary, to ensure the effective operation of the prescription monitoring program. A contractor shall be bound to comply with the provisions regarding confidentiality of prescription information in this section.

(j) The department shall promulgate rules and regulations setting forth the procedures and methods for implementing this section.

(k) The department shall submit an annual report on the effectiveness of the prescription monitoring program with the clerks of the house and senate, the chairs of the joint committee on public health, the chairs of the joint committee on health care financing and the chairs of the joint committee on public safety and homeland security.

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Minnesota  
§ 152.126  
§ 245A.192

Minnesota Statutes Annotated (2013)  
Health (Ch. 144-159)  
Chapter 152. Drugs; Controlled Substances  
Prescriptions

§ 152.126. Controlled substances prescription electronic reporting system

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Subd. 5. Use of data by board. (a) The board shall develop and maintain a database of the data reported under subdivision 4. The board shall maintain data that could identify an individual prescriber or dispenser in encrypted form. The database may be used by permissible users identified under subdivision 6 for the identification of:

(1) individuals receiving prescriptions for controlled substances from prescribers who subsequently obtain controlled substances from dispensers in quantities or with a frequency inconsistent with generally recognized standards of use for those controlled substances, including standards accepted by national and international pain management associations; and

(2) individuals presenting forged or otherwise false or altered prescriptions for controlled substances to dispensers.

(b) No permissible user identified under subdivision 6 may access the database for the sole purpose of identifying prescribers of controlled substances for unusual or excessive prescribing patterns without a valid search warrant or court order.

(c) No personnel of a state or federal occupational licensing board or agency may access the database for the purpose of obtaining information to be used to initiate or substantiate a disciplinary action against a prescriber.

(d) Data reported under subdivision 4 shall be retained by the board in the database for a 12-month period, and shall be removed from the database no later than 12 months from the last day of the month during which the data was received.

Subd. 6. Access to reporting system data. (a) Except as indicated in this subdivision, the data submitted to the board under subdivision 4 is private data on individuals as defined in section 13.02, subdivision 12, and not subject to public disclosure.

**(b) Except as specified in subdivision 5, the following persons shall be considered permissible users and may access the data submitted under subdivision 4 in the same or similar manner, and for the same or similar purposes, as those persons who are authorized to access similar private data on individuals under federal and state law:**

**(1) a prescriber or an agent or employee of the prescriber to whom the prescriber has delegated the task of accessing the data, to the extent the information relates specifically to a current patient, to whom the prescriber is prescribing or considering prescribing any controlled substance and with the provision that the prescriber remains responsible for the use or misuse of data accessed by a delegated agent or employee;**

**(2) a dispenser or an agent or employee of the dispenser to whom the dispenser has delegated the task of accessing the data, to the extent the information relates specifically to a current patient to whom that dispenser is dispensing or considering dispensing any controlled substance and with the provision that the dispenser remains responsible for the use or misuse of data accessed by a delegated agent or employee;**

(3) an individual who is the recipient of a controlled substance prescription for which data was submitted under subdivision 4, or a guardian of the individual, parent or guardian of a minor, or health care agent of the individual acting under a health care directive under chapter 145C;

(4) personnel of the board specifically assigned to conduct a bona fide investigation of a specific licensee;

(5) personnel of the board engaged in the collection of controlled substance prescription information as part of the assigned duties and responsibilities under this section;

(6) authorized personnel of a vendor under contract with the board who are engaged in the design, implementation, operation, and maintenance of the electronic reporting system as part of the assigned duties and responsibilities of their employment, provided that access to data is limited to the minimum amount necessary to carry out such duties and responsibilities;

(7) federal, state, and local law enforcement authorities acting pursuant to a valid search warrant;

(8) personnel of the medical assistance program assigned to use the data collected under this section to identify recipients whose usage of controlled substances may warrant restriction to a single primary care physician, a single outpatient pharmacy, or a single hospital; and

(9) personnel of the Department of Human Services assigned to access the data pursuant to paragraph (h).

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

(c) Any permissible user identified in paragraph (b), who directly accesses the data electronically, shall implement and maintain a comprehensive information security program that contains administrative, technical, and physical safeguards that are appropriate to the user's size and complexity, and the sensitivity of the personal information obtained. The permissible user shall identify reasonably foreseeable internal and external risks to the security, confidentiality, and integrity of personal information that could result in the unauthorized disclosure, misuse, or other compromise of the information and assess the sufficiency of any safeguards in place to control the risks.

(d) The board shall not release data submitted under this section unless it is provided with evidence, satisfactory to the board, that the person requesting the information is entitled to receive the data.

(e) The board shall not release the name of a prescriber without the written consent of the prescriber or a valid search warrant or court order. The board shall provide a mechanism for a prescriber to submit to the board a signed consent authorizing the release of the prescriber's name when data containing the prescriber's name is requested.

(f) The board shall maintain a log of all persons who access the data and shall ensure that any permissible user complies with paragraph (c) prior to attaining direct access to the data.

(g) Section 13.05, subdivision 6, shall apply to any contract the board enters into pursuant to subdivision 2. A vendor shall not use data collected under this section for any purpose not specified in this section.

(h) With available appropriations, the commission of human services shall establish and implement a system through which the Department of Human Services shall routinely access the data for the purpose of determining whether any client enrolled in an opioid treatment program licensed according to chapter 245A has been prescribed or dispensed a controlled substance in addition to that administered or dispensed by the opioid treatment program. When the commissioner determines there have been multiple prescribers or multiple prescriptions, the commissioner shall:

(1) inform the medical director of the opioid treatment program only that the commissioner determined the existence of multiple prescribers or multiple prescriptions of controlled substances; and

(2) direct the medical director of the opioid treatment program to access the data directly, review the effect of the multiple prescribers or multiple prescriptions, and document the review.

If determined necessary, the commission of human services shall seek a federal waiver of, or exception to, any applicable provision of Code of Federal Regulations, title 42, part 2.34, item (c), prior to implementing this paragraph.

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Minnesota Statutes Annotated (2013)  
Public Welfare and Related Activities (Ch. 245-267)  
Chapter 245A. Human Services Licensing

§ 245A.192. Providers licensed to provide treatment of opioid addiction.

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Subd. 11. Prescription monitoring program. (a) Upon admission to a methadone clinic outpatient treatment program, clients shall be notified that the Department of Human Services and the medical director will monitor the prescription monitoring program to review the prescribed controlled drugs the clients have received. **The medical director or the medical director's delegate must review data from the Minnesota Board of Pharmacy, prescription monitoring program (PMP) established under section 152.126 prior to the client being ordered any controlled substance as defined under section 152.126, subdivision 1, paragraph (b), including medications used for the treatment of opioid addiction.** The subsequent reviews of the PMP data must occur quarterly and be documented in the client's individual file. When the PMP data shows a recent history of multiple prescribers or multiple prescriptions for controlled substances, then subsequent reviews of the PMP data must occur monthly and be documented in the client's individual file. If, at any time, the medical director believes the use of the controlled substances places the client at risk of harm, the program must seek consent to discuss the client's opioid treatment with other prescribers and must seek consent for the other prescriber to disclose to the opioid treatment programs' medical director the client's condition that formed the basis of the other prescriptions. Additionally, any findings from the PMP data that are relevant to the medical director's course of treatment for the client must be documented in the client's individual file. A review of the PMP is not required for every medication dose adjustment.

(b) The commissioner shall collaborate with the Minnesota Board of Pharmacy to develop and implement an electronic system through which the commissioner shall routinely access the data from the Minnesota Board of Pharmacy prescription monitoring program established under section 152.126 for the purpose of determining whether any client enrolled in an opioid addiction treatment program licensed according to this section has also been prescribed or dispensed a controlled substance in addition to that administered or dispensed by the opioid addiction treatment program. When the commissioner determines there have been multiple prescribers or multiple prescriptions of controlled substances, the commissioner shall:

(1) inform the medical director of the opioid treatment program only that the commissioner determined the existence of multiple prescribers or multiple prescriptions of controlled substances; and

(2) direct the medical director of the opioid treatment program to access the data directly, review the effect of the multiple prescribers or multiple prescriptions, and document the review.



(c) If determined necessary, the commissioner shall seek a federal waiver of, or exception to, any applicable provision of Code of Federal Regulations, title 42, part 2.34, item (c), prior to implementing this paragraph.

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Montana  
ADC 24.174.1701

Administrative Rules of Montana (2012)  
Title 24. Labor and Industry  
Chapter 174. Board of Pharmacy  
Sub-chapter 17. Prescription Drug Registry

#### 24.174.1701 DEFINITIONS

(1) “Authorized user” means a prescriber, pharmacist, Board of Pharmacy staff, Montana Medicare or Medicaid programs, Tribal Health, Indian Health Service, and Veterans Affairs.

**(2) “Authorized agent” means a designated person authorized access by an authorized user. An authorized agent for a pharmacist must be a pharmacy intern or certified pharmacy technician.**

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New Mexico  
ADC 16.19.29

Code of New Mexico Rules (2013)  
Title 16. Occupational and Professional Licensing  
Chapter 19. Pharmacists  
Part 29. Controlled Substance Prescription Monitoring Program

#### 16.19.29. CONTROLLED SUBSTANCE PRESCRIPTION MONITORING PROGRAM

16.19.29.1 ISSUING AGENCY: Regulation and Licensing Department - Board of Pharmacy.

16.19.29.2 SCOPE: All persons or entities that dispense controlled substances pursuant to prescriptions from practitioners.

16.19.29.3 STATUTORY AUTHORITY: Section 30-31-16 of the Controlled Substance Act. 30-31-1 through 30-31-42 NMSA 1978, authorizes the board of pharmacy to promulgate regulations and charge reasonable fees regarding controlled substances. 30-31-16 authorizes the board to collect information regarding controlled substances.

16.19.29.4 DURATION: Permanent.

16.19.29.5 EFFECTIVE DATE: 07-15-04, unless a later date is cited at the end of a section.

16.19.29.6 OBJECTIVE: The objective of Part 29 of Chapter 19 is to promote the public health and welfare by detecting and preventing substance abuse and encouraging appropriate treatment of pain and other conditions for which controlled substances are prescribed. The purpose of the system is to improve access to controlled substances for legitimate medical needs by allowing a practitioner or a pharmacist to obtain a patient's pharmaceutical history related to controlled substances. The program's objectives will include education of the public and health care professionals regarding the nature and extent of the problem of drug abuse, appropriate prescribing and use of controlled substances, and the medical treatment options for abusers of controlled substances and pain management.

...

**16.19.29.9 ACCESS TO PRESCRIPTION INFORMATION: Practitioners registered with the program may designate one delegate per practice site to register with the program for the purpose of requesting and receiving reports for the practitioner.**

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New York  
Public Health Law § 3343-a

Mckinney's Consolidated Laws of New York Annotated (2013)  
Public Health Law  
Chapter 45. Of the Consolidated Laws  
Article 33. Controlled Substances  
Title IV. Dispensing to Ultimate Users

<Text of section effective August 27, 2013, except that pharmacists and designees may access the registry prior to that time to the extent practicable>

§ 3343-a. Prescription Monitoring Program Registry

1. Establishment of system. (A) The commissioner shall, in accordance with the provisions of this section, establish and maintain an electronic system for collecting, monitoring and reporting information concerning the prescribing and dispensing of controlled substances, to be known as the prescription monitoring program registry. The registry shall include information reported by pharmacies on a real time basis, as set forth in subdivision four of section thirty-three hundred thirty-three of this article.

...

**(D) The department shall establish and implement such protocols as are reasonably necessary to ensure that information contained in the registry is maintained in a secure and confidential manner and is accessible only by practitioners, pharmacists or their designees for the purposes established in subdivisions two and three of this section, or as otherwise set forth in sections thirty-three hundred seventy-one and thirty-three hundred seventy-one-a of this article.** Such protocols shall include a mechanism for the department to monitor and record access to the registry, which shall identify the authorized individual accessing and each controlled substance history accessed.

...

**(B) For purposes of this section, a practitioner may authorize a designee to consult the prescription monitoring program registry on his or her behalf, provided that:**

**(I) The designee so authorized is employed by the same professional practice or is under contract with such practice;**

**(II) The practitioner takes reasonable steps to ensure that such designee is sufficiently competent in the use of the registry;**

**(III) The practitioner remains responsible for ensuring that access to the registry by the designee is limited to authorized purposes and occurs in a manner that protects the confidentiality of the information obtained from the registry, and remains responsible for any breach of confidentiality; and**

**(IV) The ultimate decision as to whether or not to prescribe or dispense a controlled substance remains with the practitioner and is reasonably informed by the relevant controlled substance history information obtained from the registry.**

**The commissioner shall establish in regulation reasonable parameters with regard to a practitioner's ability to authorize designees pursuant to this section, which shall include processes necessary to allow the department to:**

**(a) Grant access to the registry in a reasonably prompt manner to as many designees as are authorized by practitioners, up to the number deemed appropriate by the commissioner for particular professional practices or types of practices, taking into account the need to maintain security of the registry and the patient-specific information maintained therein, and the objective of minimizing burdens to practitioners to the extent practicable;**

**(b) Require that practitioners notify the department upon terminating the authorization of any designee; and**

**(c) Establish a mechanism to prevent such terminated designees from accessing the registry in a reasonably prompt manner following such notification.**

3. Authority to consult prescription monitoring program registry; pharmacists. (A) A pharmacist may consult the prescription monitoring program registry in order to review the controlled substance history of an individual for whom one or more prescriptions for controlled substances is presented to such pharmacist.

**(B) For purposes of this section, a pharmacist may designate another pharmacist, a pharmacy intern, as defined by section sixty-eight hundred six of the Education Law, or other individual as may be permitted by the commissioner in regulation, to consult the prescription monitoring program registry on the pharmacist's behalf, provided that such designee is employed by the same pharmacy or is under contract with such pharmacy. The commissioner shall establish in regulation reasonable parameters with regard to a pharmacist's ability to authorize designees pursuant to this section, which shall include processes necessary to allow the department to:**

**(a) Grant access to the registry in a reasonably prompt manner to as many designees as are authorized by pharmacists, up to the number deemed appropriate by the commissioner for particular pharmacies, taking into account the need to maintain security of the registry and the patient-specific information maintained therein, and the objective of minimizing burdens to pharmacists to the extent practicable;**

**(b) Require that pharmacists notify the department upon terminating the authorization of any designee; and**

**(c) Establish a mechanism to prevent such terminated designees from accessing the registry in a reasonably prompt manner following such notification.**

4. Immunity. No practitioner or pharmacist, and no person acting on behalf of such practitioner or pharmacist as permitted under this section, acting with reasonable care and in good faith shall be subject to civil liability arising from any false, incomplete or inaccurate information submitted to or reported by the registry or for any resulting failure of the system to accurately or timely report such information; provided, however, that nothing in this subdivision shall be deemed to alter the obligation to submit or report prescription information to the department as otherwise set forth in this article or in regulations promulgated pursuant thereto.

5. Guidance to practitioners and pharmacists. The commissioner shall, in consultation with the commissioner of education, provide guidance to practitioners, pharmacists, and pharmacies regarding the purposes and uses of the registry established by this section and the means by which practitioners and pharmacists can access the registry. Such guidance shall reference educational information available pursuant to the prescription pain medication awareness program established pursuant to section thirty-three hundred nine-a of this article.

6. Individual access to controlled substance histories. The commissioner shall establish procedures by which an individual may:

(A) Request and obtain his or her own controlled substances history consisting of patient-specific information or, in appropriate circumstances, that of a patient who lacks capacity to make health care decisions and for whom the individual has legal authority to make such decisions and would have legal access to the patient's health care records; or

(B) Seek review of any part of his or her controlled substances history or, in appropriate circumstances, that of a patient who lacks capacity to make health care decisions and for whom the individual has legal authority to make such decisions and would have legal access to the patient's health care records, that such individual disputes.

Such procedures shall require the department to promptly revise any information accessible through the registry that the department determines to be inaccurate. Such procedures shall be described on the department's website and included with the controlled substances history provided to an individual pursuant to a request made under this subdivision or under subparagraph (IV) of paragraph (A) of subdivision two of section thirty-three hundred seventy-one of this article.

7. Department analysis of data. The department shall periodically analyze data contained in the prescription monitoring program registry to identify information that indicates that a violation of law or breach of professional standards may have occurred and, as warranted, provide any relevant information to appropriate entities as permitted under section thirty-three hundred

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seventy-one of this article. The department shall keep a record of the information provided, including, but not limited to, the specific information provided and the agency to which such information was provided, including the name and title of the person to whom such information was provided and an attestation from such person that he or she has authority to receive such information.

8. Funding the prescription monitoring program registry. (A) The commissioner shall make reasonable efforts to apply for monies available from the federal government and other institutions, to the extent deemed appropriate by the commissioner, and use any monies so obtained to supplement any other monies made available for the purposes of this title.

(B) Operation of the registry established by this section shall not be funded, in whole or in part, by fees imposed specifically for such purposes upon practitioners, pharmacists, designees or patients subject to this section.

9. Rules and regulations. The commissioner shall promulgate such rules and regulations as are necessary to effectuate the provisions of this section, in consultation with the work group established pursuant to subdivision three of section thirty-three hundred nine-a of this article.

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North Carolina  
§ 90-113.74

West's North Carolina General Statutes Annotated (2013)  
Chapter 90. Medicine and Allied Occupations  
Article 5E. North Carolina Controlled Substances Reporting System Act

§ 90-113.74. Confidentiality

(a) Prescription information submitted to the Department is privileged and confidential, is not a public record pursuant to G.S. 132-1, is not subject to subpoena or discovery or any other use in civil proceedings, and except as otherwise provided below may only be used for investigative or evidentiary purposes related to violations of State or federal law and regulatory activities. Except as otherwise provided by this section, prescription information shall not be disclosed or disseminated to any person or entity by any person or entity authorized to review prescription information.

(b) The Department may use prescription information data in the controlled substances reporting system only for purposes of implementing this Article in accordance with its provisions.

(b1) The Department may review the prescription information data in the controlled substances reporting system and upon review may:

(1) Notify practitioners that a patient may have obtained prescriptions for controlled substances in a manner that may represent abuse, diversion of controlled substances, or an increased risk of harm to the patient.

(2) Report information regarding the prescribing practices of a practitioner to the agency responsible for licensing, registering, or certifying the practitioner pursuant to rules adopted by the agency as set forth below in subsection (b2) of this section.

(b2) In order to receive a report pursuant to subdivision (2) of subsection (b1) of this section, an agency responsible for licensing, registering, or certifying a practitioner with prescriptive or dispensing authority shall adopt rules setting the criteria by which the Department may report the information to the agency. The criteria for reporting established by rule shall not establish the standard of care for prescribing or dispensing, and it shall not be a basis for disciplinary action by an agency that the Department reported a practitioner to an agency based on the criteria.

**(c) The Department shall release data in the controlled substances reporting system to the following persons only:**

**(1) Persons authorized to prescribe or dispense controlled substances for the purpose of providing medical or pharmaceutical care for their patients. A person authorized to**



**receive data pursuant to this paragraph may delegate the authority to receive the data to other persons working under his or her direction and supervision, provided the Department approves the delegation.**

(2) An individual who requests the individual's own controlled substances reporting system information.

(3) Special agents of the North Carolina State Bureau of Investigation who are assigned to the Diversion & Environmental Crimes Unit and whose primary duties involve the investigation of diversion and illegal use of prescription medication. SBI agents assigned to the Diversion & Environmental Crimes Unit may then provide this information to other SBI agents who are engaged in a bona fide specific investigation related to enforcement of laws governing licit drugs. The SBI shall notify the Office of the Attorney General of North Carolina of each request for inspection of records maintained by the Department.

(4) Primary monitoring authorities for other states pursuant to a specific ongoing investigation involving a designated person, if information concerns the dispensing of a Schedule II through V controlled substance to an ultimate user who resides in the other state or the dispensing of a Schedule II through V controlled substance prescribed by a licensed health care practitioner whose principal place of business is located in the other state.

(5) To a sheriff or designated deputy sheriff or a police chief or a designated police investigator who is assigned to investigate the diversion and illegal use of prescription medication or pharmaceutical products identified in Article 5 of this Chapter of the General Statutes as Schedule II through V controlled substances and who is engaged in a bona fide specific investigation related to the enforcement of laws governing licit drugs pursuant to a lawful court order specifically issued for that purpose.

(6) The Division of Medical Assistance for purposes of administering the State Medical Assistance Plan.

(7) Licensing boards with jurisdiction over health care disciplines pursuant to an ongoing investigation by the licensing board of a specific individual licensed by the board.

(8) Any county medical examiner appointed by the Chief Medical Examiner pursuant to G.S. 130A-382 and the Chief Medical Examiner, for the purpose of investigating the death of an individual.

(d) The Department may provide data to public or private entities for statistical, research, or educational purposes only after removing information that could be used to identify individual patients who received prescription medications from dispensers.

(e) In the event that the Department finds patterns of prescribing medications that are unusual, the Department shall inform the Attorney General's Office of its findings. The Office of the Attorney General shall review the Department's findings to determine if the findings should be

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reported to the SBI and the appropriate sheriff for investigation of possible violations of State or federal law relating to controlled substances.

(f) The Department shall purge from the controlled substances reporting system database all information more than six years old.

(g) Nothing in this Article shall prohibit a person authorized to prescribe or dispense controlled substances pursuant to Article 1 of Chapter 90 of the General Statutes from disclosing or disseminating data regarding a particular patient obtained under subsection (c) of this section to another person (i) authorized to prescribe or dispense controlled substances pursuant to Article 1 of Chapter 90 of the General Statutes and (ii) authorized to receive the same data from the Department under subsection (c) of this section.

(h) Nothing in this Article shall prevent persons licensed or approved to practice medicine or perform medical acts, tasks, and functions pursuant to Article 1 of Chapter 90 of the General Statutes from retaining data received pursuant to subsection (c) of this section in a patient's confidential health care record.

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## North Dakota

Per the state PMP representative, North Dakota will allow the designation of agents to access the PMP database.

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Ohio  
§ 4729.80

Baldwin's Ohio Revised Code Annotated (2013)  
Title XLVII. Occupations--Professions  
Chapter 4729. Pharmacists; Dangerous Drugs  
Miscellaneous Provisions

§ 4729.80 Disclosure of database information; disclosure of requests for database information

**(A) If the state board of pharmacy establishes and maintains a drug database pursuant to section 4729.75 of the Revised Code, the board is authorized or required to provide information from the database in accordance with the following:**

(1) On receipt of a request from a designated representative of a government entity responsible for the licensure, regulation, or discipline of health care professionals with authority to prescribe, administer, or dispense drugs, the board may provide to the representative information from the database relating to the professional who is the subject of an active investigation being conducted by the government entity.

(2) On receipt of a request from a federal officer, or a state or local officer of this or any other state, whose duties include enforcing laws relating to drugs, the board shall provide to the officer information from the database relating to the person who is the subject of an active investigation of a drug abuse offense, as defined in section 2925.01 of the Revised Code, being conducted by the officer's employing government entity.

(3) Pursuant to a subpoena issued by a grand jury, the board shall provide to the grand jury information from the database relating to the person who is the subject of an investigation being conducted by the grand jury.

(4) Pursuant to a subpoena, search warrant, or court order in connection with the investigation or prosecution of a possible or alleged criminal offense, the board shall provide information from the database as necessary to comply with the subpoena, search warrant, or court order.

**(5) On receipt of a request from a prescriber or the prescriber's delegate approved by the board, the board may provide to the prescriber information from the database relating to a patient who is either of the following, if the prescriber certifies in a form specified by the board that it is for the purpose of providing medical treatment to the patient who is the subject of the request;**

**(a) A current patient of the prescriber;**

**(b) A potential patient of the prescriber based on a referral of the patient to the prescriber.**

**(6) On receipt of a request from a pharmacist or the pharmacist's delegate approved by the board, the board may provide to the pharmacist information from the database relating to a current patient of the pharmacist, if the pharmacist certifies in a form specified by the board that it is for the purpose of the pharmacist's practice of pharmacy involving the patient who is the subject of the request.**

(7) On receipt of a request from an individual seeking the individual's own database information in accordance with the procedure established in rules adopted under section 4729.84 of the Revised Code, the board may provide to the individual the individual's own database information.

(8) On receipt of a request from the medical director of a managed care organization that has entered into a data security agreement with the board required by section 5167.14 of the Revised Code, the board shall provide to the medical director information from the database relating to a medicaid recipient enrolled in the managed care organization, including information in the database related to prescriptions for the recipient that were not covered or reimbursed under a program administered by the department of medicaid.

(9) On receipt of a request from the medicaid director, the board shall provide to the director information from the database relating to a recipient of a program administered by the department of medicaid, including information in the database related to prescriptions for the recipient that were not covered or paid by a program administered by the department.

(10) On receipt of a request from the administrator of workers' compensation, the board may provide to the administrator information from the database relating to a claimant under Chapter 4121., 4123., 4127., or 4131. of the Revised Code.

(11) On receipt of a request from a requestor described in division (A)(1), (2), (5), or (6) of this section who is from or participating with another state's prescription monitoring program, the board may provide to the requestor information from the database, but only if there is a written agreement under which the information is to be used and disseminated according to the laws of this state.

(B) The state board of pharmacy shall maintain a record of each individual or entity that requests information from the database pursuant to this section. In accordance with rules adopted under section 4729.84 of the Revised Code, the board may use the records to document and report statistics and law enforcement outcomes.

The board may provide records of an individual's requests for database information to the following:

(1) A designated representative of a government entity that is responsible for the licensure, regulation, or discipline of health care professionals with authority to prescribe, administer, or dispense drugs who is involved in an active investigation being conducted by the government entity of the individual who submitted the requests for database information;

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(2) A federal officer, or a state or local officer of this or any other state, whose duties include enforcing laws relating to drugs and who is involved in an active investigation being conducted by the officer's employing government entity of the individual who submitted the requests for database information.

(C) Information contained in the database and any information obtained from it is not a public record. Information contained in the records of requests for information from the database is not a public record. Information that does not identify a person may be released in summary, statistical, or aggregate form.

(D) A pharmacist or prescriber shall not be held liable in damages to any person in any civil action for injury, death, or loss to person or property on the basis that the pharmacist or prescriber did or did not seek or obtain information from the database.

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## Oregon

§ 431.966 (eff. Jan. 1, 2014)

West's Oregon Revised Statutes Annotated (2013)

Title 36. Public Health and Safety

Chapter 431. State and Local Administration and Enforcement of Health Laws

Prescription Monitoring Program

(Program)

§ 431.966. Prescription monitoring information disclosure; limitations

<Text of section effective January 1, 2014>

(1)(a) Except as provided under subsection (2) of this section, prescription monitoring information submitted under ORS 431.964 to the prescription monitoring program established in ORS 431.962:

(A) Is protected health information under ORS 192.553 to 192.581.

(B) Is not subject to disclosure pursuant to ORS 192.410 to 192.505.

(b) Except as provided under subsection (2)(a)(E) of this section, prescription monitoring information submitted under ORS 431.964 to the prescription monitoring program may not be used to evaluate a practitioner's professional practice.

**(2)(a) To the extent that the law or regulation is applicable to the prescription monitoring program, if a disclosure of prescription monitoring information, other than the sex of a patient for whom a drug was prescribed, complies with the federal Health Insurance Portability and Accountability Act of 1996 (P.L. 104-191) and regulations adopted under it, including 45 C.F.R. parts 160 and 164, federal alcohol and drug treatment confidentiality laws and regulations adopted under those laws, including 42 C.F.R. part 2, and state health and mental health confidentiality laws, including ORS 179.505, 192.517 and 192.553 to 192.581, the Oregon Health Authority shall disclose the information:**

**(A) To a practitioner or pharmacist, or, if a practitioner or pharmacist authorizes the authority to disclose the information to a member of the practitioner's or pharmacist's staff, to a member of the practitioner's or pharmacist's staff. If a practitioner or pharmacist authorizes disclosing the information to a member of the practitioner's or pharmacist's staff under this subparagraph, the practitioner or pharmacist remains responsible for the use or misuse of the information by the staff member. To receive information under this subparagraph, or to authorize the receipt of information by a staff member under this subparagraph, a practitioner or pharmacist must certify that the requested information is for the purpose of evaluating the need for or providing medical or**

**pharmaceutical treatment for a patient to whom the practitioner or pharmacist anticipates providing, is providing or has provided care.**

(B) To a practitioner in a form that catalogs all prescription drugs prescribed by the practitioner according to the number assigned to the practitioner by the Drug Enforcement Administration of the United States Department of Justice.

(C) To designated representatives of the authority or any vendor or contractor with whom the authority has contracted to establish or maintain the electronic system of the prescription monitoring program.

(D) Pursuant to a valid court order based on probable cause and issued at the request of a federal, state or local law enforcement agency engaged in an authorized drug-related investigation involving a person to whom the requested information pertains.

(E) To a health professional regulatory board that certifies in writing that the requested information is necessary for an investigation related to licensure, renewal or disciplinary action involving the applicant, licensee or registrant to whom the requested information pertains.

(F) To a prescription monitoring program of another state if the confidentiality, security and privacy standards of the requesting state are determined by the authority to be equivalent to those of the authority.

(G) To the State Medical Examiner or designee of the State Medical Examiner, for the purpose of conducting a medicolegal investigation or autopsy.

(b) The authority may disclose information from the prescription monitoring program that does not identify a patient, practitioner or drug outlet:

(A) For educational, research or public health purposes;

(B) To a local public health authority, as defined in ORS 431.260; or

(C) To officials of the authority who are conducting special epidemiologic morbidity and mortality studies in accordance with ORS 432.060 and rules adopted under ORS 431.110.

(c) The authority shall disclose information relating to a patient maintained in the electronic system operated pursuant to the prescription monitoring program established under ORS 431.962 to that patient at no cost to the patient within 10 business days after the authority receives a request from the patient for the information.

(d)(A) A patient may request the authority to correct any information about the patient that is erroneous. The authority shall grant or deny a request to correct information within 10 business days after the authority receives the request.



(B) If the authority denies a patient's request to correct information under this paragraph, or fails to grant a patient's request to correct information under this paragraph within 10 business days after the authority receives the request, the patient may appeal the denial or failure to grant the request. Upon receipt of an appeal under this subparagraph, the authority shall conduct a contested case hearing as provided in ORS chapter 183. Notwithstanding ORS 183.450, in the contested case hearing, the authority has the burden of establishing that the information included in the prescription monitoring program is correct.

(e) The information in the prescription monitoring program may not be used for any commercial purpose.

(f) In accordance with ORS 192.553 to 192.581 and federal privacy regulations, any person authorized to prescribe or dispense a prescription drug and who is entitled to access a patient's prescription monitoring information may discuss or release the information to other health care providers involved with the patient's care, in order to provide safe and appropriate care coordination.

(3)(a) The authority shall maintain records of the information disclosed through the prescription monitoring program including, but not limited to:

(A) The identity of each person who requests or receives information from the program and the organization, if any, the person represents;

(B) The information released to each person or organization; and

(C) The date and time the information was requested and the date and time the information was provided.

(b) Records maintained as required by this subsection may be reviewed by the Prescription Monitoring Program Advisory Commission.

(4) Information in the prescription monitoring program that identifies an individual patient must be removed no later than three years from the date the information is entered into the program.

(5) The authority shall notify the Attorney General and each affected individual of an improper disclosure of information from the prescription monitoring program.

(6)(a) If the authority or a person or entity required to report or authorized to receive or release controlled substance prescription information under this section violates this section or ORS 431.964 or 431.968, a person injured by the violation may bring a civil action against the authority, person or entity and may recover damages in the amount of \$1,000 or actual damages, whichever is greater.

(b) Notwithstanding paragraph (a) of this subsection, the authority and a person or entity required to report or authorized to receive or release controlled substance prescription

information under this section are immune from civil liability for violations of this section or ORS 431.964 or 431.968 unless the authority, person or entity acts with malice, criminal intent, gross negligence, recklessness or willful intent.

(7) Nothing in ORS 431.962 to 431.978 and 431.992 requires a practitioner or pharmacist who prescribes or dispenses a prescription drug to obtain information about a patient from the prescription monitoring program. A practitioner or pharmacist who prescribes or dispenses a prescription drug may not be held liable for damages in any civil action on the basis that the practitioner or pharmacist did or did not request or obtain information from the prescription monitoring program.

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## South Dakota

Per the state PMP representative, South Dakota will allow a prescriber to “sponsor” a designated agent. At this time, they do not allow pharmacists to designate an agent.

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Tennessee

§ 53-10-302 (eff. until Oct. 1, 2013)

§ 53-10-302 (eff. Oct. 1, 2013)

§ 53-10-306 (eff. until July 1, 2016)

West's Tennessee Code Annotated (2013)

Title 53. Food, Drugs and Cosmetics

Chapter 10. Legend Drugs

Part 3. Tennessee Prescription Safety Act of 2012

§ 53-10-302. Definitions

<Text of section effective until October 1, 2013>

As used in this part:

...

**(10) “Healthcare practitioner extender” means any registered or licensed healthcare professional, and up to two (2) unlicensed persons designated by the prescriber or dispenser, who act as agents of that prescriber or dispenser. The prescriber or dispenser shall be responsible for all actions taken by their agents pursuant to this part;**

...

West's Tennessee Code Annotated (2013)

Title 53. Food, Drugs and Cosmetics

Chapter 10. Legend Drugs

Part 3. Tennessee Prescription Safety Act of 2012

§ 53-10-302. Definitions

<Text of section effective October 1, 2013>

As used in this part:

...

**(10) “Healthcare practitioner extender” means any registered or licensed healthcare professional, and up to two (2) unlicensed persons per prescriber or dispenser designated by the prescriber or dispenser to act as agents of such prescriber or dispenser. A prescriber shall have the ability to authorize a healthcare practitioner extender to check**

**the controlled substance database as stipulated in § 53-10-310(e) for other prescribers in the authorizing prescriber's practice. Notwithstanding the provisions of Section 28 of Chapter 880 of the Public Acts of 2012, any one-time costs required to be made to effectuate the provisions of this act specific to system modifications required by changes in this subdivision shall be shared on a pro-rata basis, excluding the pharmacy board, by the appropriate prescribing boards as enumerated in title 53, chapter 10, part 3. The prescriber or dispenser shall be responsible for actions taken by their agents pursuant to this part;**

...

West's Tennessee Code Annotated (2013)  
Title 53. Food, Drugs and Cosmetics  
Chapter 10. Legend Drugs  
Part 3. Tennessee Prescription Safety Act of 2012

§ 53-10-306. Confidentiality; disclosure; penalties

<Text of section effective until July 1, 2016>

**(a) Information sent to, contained in, and reported from the database in any format is confidential and not subject to title 10, chapter 7, regarding public records, and not subject to subpoena from any court and shall be made available only as provided for in § 53-10-308 and to the following persons in accordance with the limitations stated and rules promulgated pursuant to this part, or as otherwise provided for in § 53-10-311:**

- (1) Personnel of the committee specifically assigned to conduct analysis or research;
- (2) Authorized committee, board, or department of health personnel or any designee appointed by the committee engaged in analysis of controlled substances prescription information as a part of the assigned duties and responsibilities of their employment;
- (3) A prescriber conducting medication history reviews who is actively involved in the care of the patient; a prescriber or supervising physician of the prescriber conducting a review of all medications dispensed by prescription attributed to that prescriber; or a prescriber having authority to prescribe or dispense controlled substances, to the extent the information relates specifically to a current or bona fide prospective patient of the prescriber, to whom the prescriber has prescribed or dispensed, is prescribing or dispensing, or considering prescribing or dispensing any controlled substance. Each authorized individual referenced under this subdivision shall have a separate identifiable authentication for access;
- (4) A dispenser or pharmacist not authorized to dispense controlled substances conducting drug utilization or medication history reviews who is actively involved in the care of the patient; or a dispenser having authority to dispense controlled substances to the extent the information relates

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specifically to a current or a bona fide prospective patient to whom that dispenser has dispensed, is dispensing, or considering dispensing any controlled substance. Each authorized individual referenced under this subdivision shall have a separate identifiable authentication for access;

(5) A county medical examiner appointed pursuant to § 38-7-104 when acting in an official capacity as established in § 38-7-109; provided, any access to information from the database shall be subject to the confidentiality provisions of this part except where information obtained from the database is appropriately included in any official report of the county medical examiners, toxicological reports or autopsy reports issued by the county medical examiner under § 38-7-110(c);

(6) Personnel of the following entities actively engaged in analysis of controlled substances prescription information as a part of their assigned duties and responsibilities related directly to TennCare:

(A) The office of inspector general;

(B) The medicaid fraud control unit; and

(C) The bureau of TennCare's chief medical officer, associate chief medical directors, director of quality oversight, and associate director of pharmacy;

(7) A quality improvement committee as defined in § 68-11-272 of a hospital licensed under title 68 or title 33, as part of the committee's confidential and privileged activities under § 68-11-272(b)(4) with respect to the evaluation, supervision or discipline of a healthcare provider employed by the hospital or any of its affiliates or subsidiaries, who is known or suspected by the hospital's administrator to be prescribing controlled substances for the prescriber's personal use;

(8) Law enforcement personnel; provided, that such personnel are engaged in the official investigation and enforcement of state or federal laws involving controlled substances or violations under this part; and that any law enforcement personnel receiving information from the database pursuant to this section shall comply with the requirements of this subsection (a):

(A)(i) Any law enforcement agency or judicial district drug task force that wants one (1) or more of its officers or agents to have the authorization to request information from the database shall first pre-approve each such officer. Pre-approval shall be by the applicant's supervisor, who shall be either the chief of police, county sheriff or the judicial district drug task force director. The list of pre-approved applicants shall be sent to the district attorney general in the judicial district in which the agency or task force has jurisdiction;

(ii) By December 1 of each year, each district attorney general shall send to the board of pharmacy a list of applicants authorized to request information from the database from that general's judicial district for the next calendar year;

(B)(i) If the Tennessee bureau of investigation (TBI) wants one (1) or more of its agents to have the authorization to request information from the database each such agent shall first be pre-approved by the agent's immediate supervisor and division head. Approved applicants shall be sent to the board by the director;

(ii) By December 1 of each year, the TBI director shall send to the board of pharmacy a list of applicants authorized to request information from the database from the bureau for the next calendar year;

(C) An application submitted by law enforcement personnel shall include, but not be limited to the:

(i) Applicant's name; title; agency; agency address; agency contact number; agency supervisor; and badge number, identification number or commission number, and the business email address of each applicant officer or agent, the appropriate district attorney general and, if a TBI agent, the TBI director and their business email addresses; and

(ii) Signatures of the applicant, the applicants approving supervisor and the district attorney general of the judicial district in which the applicant has jurisdiction or the approving division head and the TBI director;

(D) It shall be a duty of the board, as part of its duties to maintain the database pursuant to § 53-10-305(c), to receive and verify the lists of authorized applications sent to it by the district attorneys general and the director of the TBI pursuant to this subsection (a); or

**(9) A healthcare practitioner extender, who is acting under the direction and supervision of a prescriber or dispenser, and only to the extent the information relates specifically to a current or bona fide prospective patient to whom the prescriber or dispenser has prescribed or dispensed, is prescribing or dispensing, or considering prescribing or dispensing any controlled substance. Each authorized individual referenced under this subdivision shall have a separate identifiable authentication for access.**

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Texas

Health & Safety § 481.076 (eff. Sept. 1, 2013)

Vernon's Texas Statutes and Codes Annotated (2013)

Health and Safety Code

Title 6. Food, Drugs, Alcohol, and Hazardous Substances

Subtitle C. Substance Abuse Regulation and Crimes

Chapter 481. Texas Controlled Substances Act

Subchapter C. Regulation of Manufacture, Distribution, and Dispensation of Controlled Substances, Chemical Precursors, and Chemical Laboratory Apparatus

§ 481.076. Official Prescription Information

<Text of section effective September 1, 2013>

**(a) The director may not permit any person to have access to information submitted to the director under Section 481.074(q) or 481.075 except:**

(1) an investigator for the Texas Medical Board, the Texas State Board of Podiatric Medical Examiners, the State Board of Dental Examiners, the State Board of Veterinary Medical Examiners, the Texas Board of Nursing, or the Texas State Board of Pharmacy;

(2) an authorized officer or member of the department engaged in the administration, investigation, or enforcement of this chapter or another law governing illicit drugs in this state or another state; or

(3) if the director finds that proper need has been shown to the director:

(A) a law enforcement or prosecutorial official engaged in the administration, investigation, or enforcement of this chapter or another law governing illicit drugs in this state or another state;

**(B) a pharmacist or a pharmacy technician, as defined by Section 551.003, Occupations Code, acting at the direction of a pharmacist or a practitioner who is a physician, dentist, veterinarian, podiatrist, or advanced practice nurse or is a physician assistant described by Section 481.002(39)(D) or a nurse licensed under Chapter 301, Occupations Code, acting at the direction of a practitioner and is inquiring about a recent Schedule II, III, IV, or V prescription history of a particular patient of the practitioner; or**

(C) a pharmacist or practitioner who is inquiring about the person's own dispensing or prescribing activity.



(a-1) A person authorized to receive information under Subsection (a)(3)(B) or (C) may access that information through a health information exchange, subject to proper security measures to ensure against disclosure to unauthorized persons.

(a-2) A person authorized to receive information under Subsection (a)(3)(B) may include that information in any form in the medical or pharmacy record of the patient who is the subject of the information. Any information included in the patient's medical or pharmacy record under this subsection is subject to any applicable state or federal confidentiality or privacy laws.

(b) This section does not prohibit the director from creating, using, or disclosing statistical data about information received by the director under this section if the director removes any information reasonably likely to reveal the identity of each patient, practitioner, or other person who is a subject of the information.

(c) The director by rule shall design and implement a system for submission of information to the director by electronic or other means and for retrieval of information submitted to the director under this section and Sections 481.074 and 481.075. The director shall use automated information security techniques and devices to preclude improper access to the information. The director shall submit the system design to the Texas State Board of Pharmacy and the Texas Medical Board for review and approval or comment a reasonable time before implementation of the system and shall comply with the comments of those agencies unless it is unreasonable to do so.

(d) Information submitted to the director under this section may be used only for:

(1) the administration, investigation, or enforcement of this chapter or another law governing illicit drugs in this state or another state;

(2) investigatory or evidentiary purposes in connection with the functions of an agency listed in Subsection (a)(1); or

(3) dissemination by the director to the public in the form of a statistical tabulation or report if all information reasonably likely to reveal the identity of each patient, practitioner, or other person who is a subject of the information has been removed.

(e) The director shall remove from the information retrieval system, destroy, and make irretrievable the record of the identity of a patient submitted under this section to the director not later than the end of the 36th calendar month after the month in which the identity is entered into the system. However, the director may retain a patient identity that is necessary for use in a specific ongoing investigation conducted in accordance with this section until the 30th day after the end of the month in which the necessity for retention of the identity ends.

(f) If the director permits access to information under Subsection (a)(2) relating to a person licensed or regulated by an agency listed in Subsection (a)(1), the director shall notify and cooperate with that agency regarding the disposition of the matter before taking action against

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the person, unless the director determines that notification is reasonably likely to interfere with an administrative or criminal investigation or prosecution.

(g) If the director permits access to information under Subsection (a)(3)(A) relating to a person licensed or regulated by an agency listed in Subsection (a)(1), the director shall notify that agency of the disclosure of the information not later than the 10th working day after the date the information is disclosed.

(h) If the director withholds notification to an agency under Subsection (f), the director shall notify the agency of the disclosure of the information and the reason for withholding notification when the director determines that notification is no longer likely to interfere with an administrative or criminal investigation or prosecution.

(i) Information submitted to the director under Section 481.074(q) or 481.075 is confidential and remains confidential regardless of whether the director permits access to the information under this section.

(j) Repealed by Acts 1999, 76th Leg., ch. 145, § 5(3), eff. Sept. 1, 1999.

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Utah  
§ 58-37f-301  
ADC R156-37f

West's Utah Code Annotated (2013)  
Title 58. Occupations and Professions  
Chapter 37F. Controlled Substance Database Act  
Part 3. Access

§ 58-37f-301. Access to database

(1) The division shall make rules, in accordance with Title 63G, Chapter 3, Utah Administrative Rulemaking Act, to:

(a) effectively enforce the limitations on access to the database as described in this part; and

(b) establish standards and procedures to ensure accurate identification of individuals requesting information or receiving information without request from the database.

**(2) The division shall make information in the database and information obtained from other state or federal prescription monitoring programs by means of the database available only to the following individuals, in accordance with the requirements of this chapter and division rules:**

(a) personnel of the division specifically assigned to conduct investigations related to controlled substance laws under the jurisdiction of the division;

(b) authorized division personnel engaged in analysis of controlled substance prescription information as a part of the assigned duties and responsibilities of their employment;

(c) in accordance with a written agreement entered into with the department, employees of the Department of Health:

(i) whom the director of the Department of Health assigns to conduct scientific studies regarding the use or abuse of controlled substances, if the identity of the individuals and pharmacies in the database are confidential and are not disclosed in any manner to any individual who is not directly involved in the scientific studies; or

(ii) when the information is requested by the Department of Health in relation to a person or provider whom the Department of Health suspects may be improperly obtaining or providing a controlled substance;

(d) in accordance with a written agreement entered into with the department, a designee of the director of the Department of Health, who is not an employee of the Department of Health, whom the director of the Department of Health assigns to conduct scientific studies regarding the use or abuse of controlled substances pursuant to an application process established in rule by the Department of Health, if:

(i) the designee provides explicit information to the Department of Health regarding the purpose of the scientific studies;

(ii) the scientific studies to be conducted by the designee:

(A) fit within the responsibilities of the Department of Health for health and welfare;

(B) are reviewed and approved by an Institutional Review Board that is approved for human subject research by the United States Department of Health and Human Services; and

(C) are not conducted for profit or commercial gain; and

(D) are conducted in a research facility, as defined by division rule, that is associated with a university or college in the state accredited by the Northwest Commission on Colleges and Universities;

(iii) the designee protects the information as a business associate of the Department of Health; and

(iv) the identity of the prescribers, patients, and pharmacies in the database are de-identified, confidential, not disclosed in any manner to the designee or to any individual who is not directly involved in the scientific studies;

(e) a licensed practitioner having authority to prescribe controlled substances, to the extent the information:

(i)(A) relates specifically to a current or prospective patient of the practitioner; and

(B) is provided to or sought by the practitioner for the purpose of:

(I) prescribing or considering prescribing any controlled substance to the current or prospective patient;

(II) diagnosing the current or prospective patient;

(III) providing medical treatment or medical advice to the current or prospective patient; or

(IV) determining whether the current or prospective patient:

(Aa) is attempting to fraudulently obtain a controlled substance from the practitioner; or

(Bb) has fraudulently obtained, or attempted to fraudulently obtain, a controlled substance from the practitioner;

(ii)(A) relates specifically to a former patient of the practitioner; and

(B) is provided to or sought by the practitioner for the purpose of determining whether the former patient has fraudulently obtained, or has attempted to fraudulently obtain, a controlled substance from the practitioner;

(iii) relates specifically to an individual who has access to the practitioner's Drug Enforcement Administration identification number, and the practitioner suspects that the individual may have used the practitioner's Drug Enforcement Administration identification number to fraudulently acquire or prescribe a controlled substance;

(iv) relates to the practitioner's own prescribing practices, except when specifically prohibited by the division by administrative rule;

(v) relates to the use of the controlled substance database by an employee of the practitioner, described in Subsection (2)(f); or

(vi) relates to any use of the practitioner's Drug Enforcement Administration identification number to obtain, attempt to obtain, prescribe, or attempt to prescribe, a controlled substance;

**(f) in accordance with Subsection (3)(a), an employee of a practitioner described in Subsection (2)(e), for a purpose described in Subsection (2)(e)(i) or (ii), if:**

**(i) the employee is designated by the practitioner as an individual authorized to access the information on behalf of the practitioner;**

**(ii) the practitioner provides written notice to the division of the identity of the employee; and**

**(iii) the division:**

**(A) grants the employee access to the database; and**

**(B) provides the employee with a password that is unique to that employee to access the database in order to permit the division to comply with the requirements of Subsection 58-37f-203(3)(b) with respect to the employee;**

**(g) an employee of the same business that employs a licensed practitioner under Subsection (2)(e) if:**

**(i) the employee is designated by the practitioner as an individual authorized to access the information on behalf of the practitioner;**

**(ii) the practitioner and the employing business provide written notice to the division of the identity of the designated employee; and**

**(iii) the division:**

**(A) grants the employee access to the database; and**

**(B) provides the employee with a password that is unique to that employee to access the database in order to permit the division to comply with the requirements of Subsection 58-37f-203(3)(b) with respect to the employee;**

(h) a licensed pharmacist having authority to dispense a controlled substance to the extent the information is provided or sought for the purpose of:

(i) dispensing or considering dispensing any controlled substance; or

(ii) determining whether a person:

(A) is attempting to fraudulently obtain a controlled substance from the pharmacist; or

(B) has fraudulently obtained, or attempted to fraudulently obtain, a controlled substance from the pharmacist;

(i) federal, state, and local law enforcement authorities, and state and local prosecutors, engaged as a specified duty of their employment in enforcing laws:

(i) regulating controlled substances;

(ii) investigating insurance fraud, Medicaid fraud, or Medicare fraud; or

(iii) providing information about a criminal defendant to defense counsel, upon request during the discovery process, for the purpose of establishing a defense in a criminal case;

(j) employees of the Office of Internal Audit and Program Integrity within the Department of Health who are engaged in their specified duty of ensuring Medicaid program integrity under Section 26-18-2.3;

(k) a mental health therapist, if:

(i) the information relates to a patient who is:

**(A) enrolled in a licensed substance abuse treatment program; and**

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(B) receiving treatment from, or under the direction of, the mental health therapist as part of the patient's participation in the licensed substance abuse treatment program described in Subsection (2)(k)(i)(A);

(ii) the information is sought for the purpose of determining whether the patient is using a controlled substance while the patient is enrolled in the licensed substance abuse treatment program described in Subsection (2)(k)(i)(A); and

(iii) the licensed substance abuse treatment program described in Subsection (2)(k)(i)(A) is associated with a practitioner who:

(A) is a physician, a physician assistant, an advance practice registered nurse, or a pharmacist; and

(B) is available to consult with the mental health therapist regarding the information obtained by the mental health therapist, under this Subsection (2)(k), from the database;

(l) an individual who is the recipient of a controlled substance prescription entered into the database, upon providing evidence satisfactory to the division that the individual requesting the information is in fact the individual about whom the data entry was made;

(m) the inspector general, or a designee of the inspector general, of the Office of Inspector General of Medicaid Services, for the purpose of fulfilling the duties described in Title 63A, Chapter 13, Part 2, Office and Powers; and

(n) the following licensed physicians for the purpose of reviewing and offering an opinion on an individual's request for workers' compensation benefits under Title 34A, Chapter 2, Workers' Compensation Act, or Title 34A, Chapter 3, Utah Occupational Disease Act:

(i) a member of the medical panel described in Section 34A-2-601; or

(ii) a physician offering a second opinion regarding treatment.

(3)(a) A practitioner described in Subsection (2)(e) may designate up to three employees to access information from the database under Subsection (2)(f), (2)(g), or (4)(c).

(b) The division shall make rules, in accordance with Title 63G, Chapter 3, Utah Administrative Rulemaking Act, to:

(i) establish background check procedures to determine whether an employee designated under Subsection (2)(f), (2)(g), or (4)(c) should be granted access to the database; and

(ii) establish the information to be provided by an emergency room employee under Subsection (4).

**(c) The division shall grant an employee designated under Subsection (2)(f), (2)(g), or (4)(c) access to the database, unless the division determines, based on a background check, that the employee poses a security risk to the information contained in the database.**

**(4)(a) An individual who is employed in the emergency room of a hospital may exercise access to the database under this Subsection (4) on behalf of a licensed practitioner if the individual is designated under Subsection (4)(c) and the licensed practitioner:**

**(i) is employed in the emergency room;**

**(ii) is treating an emergency room patient for an emergency medical condition; and**

**(iii) requests that an individual employed in the emergency room and designated under Subsection (4)(c) obtain information regarding the patient from the database as needed in the course of treatment.**

**(b) The emergency room employee obtaining information from the database shall, when gaining access to the database, provide to the database the name and any additional identifiers regarding the requesting practitioner as required by division administrative rule established under Subsection (3)(b).**

**(c) An individual employed in the emergency room under this Subsection (4) may obtain information from the database as provided in Subsection (4)(a) if:**

**(i) the employee is designated by the practitioner as an individual authorized to access the information on behalf of the practitioner;**

**(ii) the practitioner and the hospital operating the emergency room provide written notice to the division of the identity of the designated employee; and**

**(iii) the division:**

**(A) grants the employee access to the database; and**

**(B) provides the employee with a password that is unique to that employee to access the database in order to permit the division to comply with the requirements of Subsection 58-37f-203(3)(b) with respect to the employee.**

**(d) The division may impose a fee, in accordance with Section 63J-1-504, on a practitioner who designates an employee under Subsection (2)(f), (2)(g), or (4)(c) to pay for the costs incurred by the division to conduct the background check and make the determination described in Subsection (3)(b).**



(5)(a) An individual who is granted access to the database based on the fact that the individual is a licensed practitioner or a mental health therapist shall be denied access to the database when the individual is no longer licensed.

**(b) An individual who is granted access to the database based on the fact that the individual is a designated employee of a licensed practitioner shall be denied access to the database when the practitioner is no longer licensed.**

Utah Administrative Code (2013)

Commerce

R156. Occupational and Professional Licensing.

R156-37f. Controlled Substance Database Act Rule.

R156-37f-101. Title.

This rule shall be known as the “Controlled Substance Database Act Rule”.

...

R156-37f-301. Access to Database Information.

In accordance with Subsections 58-37f-301(1)(a) and (b):

(1) The Division Director shall designate in writing those individuals employed by the Division who shall have access to the information in the Database (Database staff).

(2)(a) A request for information from the Database may be made:

(i) directly to the Database by electronic submission, if the requester is registered to use the Database; or

(ii) by oral or written submission to the Database staff, if the requester is not registered to use the Database.

(b) An oral request may be submitted by telephone or in person.

(c) A written request may be submitted by facsimile, email, regular mail, or in person except as otherwise provided herein.

(d) The Division may in its discretion require a requestor to verify the requestor's identity.

(3) The following Database information may be disseminated to a verified requestor who is permitted to obtain the information:

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(a) dispensing/reporting pharmacy ID number/name;

(b) subject's birth date;

(c) date prescription was filled;

(d) prescription (Rx) number;

(e) metric quantity;

(f) days supply;

(g) NDC code/drug name;

(h) prescriber ID/name;

(i) date prescription was written;

(j) subject's last name;

(k) subject's first name; and

(l) subject's street address;

(4) Federal, state and local law enforcement authorities and state and local prosecutors requesting information from the Database under Subsection 58-37f-301(2)(d) must provide a valid case number of the investigation or prosecution.

(5) An individual whose records are contained within the Database may not receive an accounting of persons or entities that have requested or received Database information about the individual.

(6) An individual whose records are contained within the Database may obtain his or her own information and records by:

(a) personally appearing before the Database staff with government-issued picture identification confirming the requester's identity; or

(b) submitting a signed and notarized request that includes the requester's:

(i) full name;

(ii) complete home address;

(iii) date of birth; and

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(iv) driver license or state identification card number.

(7) A requester holding power of attorney for an individual whose records are contained within the Database may obtain the individual's information and records by:

(a) personally appearing before the Database staff with government-issued picture identification confirming the requester's identity; and

(b) providing:

(i) an original, properly executed power of attorney designation; and

(ii) a signed and notarized request, executed by the individual whose information is contained within the Database, and including the individual's:

(A) full name;

(B) complete home address;

(C) date of birth; and

(D) driver license or state identification card number verifying the individual's identity.

(8) A requestor who is the legal guardian of a minor or incapacitated individual whose records are contained within the Database may obtain the individual information and records by:

(a) personally appearing before the Database staff with government-issued picture identification confirming the requester's identity;

(b) submitting the minor or incapacitated individual's:

(i) full name;

(ii) complete home address;

(iii) date of birth; and

(iv) if applicable, state identification card number verifying the individual's identity; and

(c) submitting legal proof that the requestor is the guardian of the individual who is the subject of the request for information from the Database.

(9) A requestor who has a release-of-records from an individual whose records are contained within the Database may obtain the individual's information and records by:

- (a) submitting a request in writing;
- (b) submitting an original, signed and notarized release-of-records in a format acceptable to the Database staff, identifying the purpose of the release; and
- (c) submitting the individual's:
  - (i) full name;
  - (ii) complete home address;
  - (iii) telephone number;
  - (iv) date of birth; and
  - (v) driver license or state identification card number verifying the identity of the person who is the subject of the request.

**(10) An employee of a licensed practitioner who is authorized to prescribe controlled substances may obtain Database information to the extent permissible under Subsection 58-37f-301(2)(d)if, prior to making the request:**

**(a) the licensed practitioner has provided to the Division a written designation that includes the designating practitioner's DEA number and the designated employee's:**

- (i) full name;**
- (ii) complete home address;**
- (iii) e-mail address;**
- (iv) date of birth; and**
- (v) driver license number or state identification card number;**

**(b) the designated employee has registered for an account for access to the Database and provided a unique user identification and password;**

**(c) the designated employee has passed a Database background check of available criminal court and Database records; and**

**(d) the Database has issued the designated employee a user personal identification number (PIN) and activated the employee's Database account. (11) An employee of a business that employs a licensed practitioner who is authorized to prescribe controlled substances may**

**obtain Database information to the extent permissible under Subsection 58-37f-301(2)(d) if, prior to making the request:**

**(a) the licensed practitioner and employing business have provided to the Division a written designation that includes:**

**(i) the designating practitioner's DEA number;**

**(ii) the name of the employing business; and**

**(iii) the designated employee's:**

**(A) full name;**

**(B) complete home address;**

**(C) e-mail address;**

**(D) date of birth; and**

**(E) driver license number or state identification card number;**

**(b) the designated employee has registered for an account for access to the Database and provided a unique user identification and password;**

**(c) the designated employee has passed a Database background check of available criminal court and Database records; and**

**(d) the Database has issued the designated employee a user personal identification number (PIN) and activated the employee's Database account.**

**(12) An individual who is employed in the emergency room of a hospital that employs a licensed practitioner who is authorized to prescribe controlled substances may obtain Database information to the extent permissible under Subsection 58-37f-301(2)(d) if, prior to making the request:**

**(a) the practitioner and the hospital operating the emergency room have provided to the Division a written designation that includes:**

**(i) the designating practitioner's DEA number;**

**(ii) the name of the hospital;**

**(iii) the names of all emergency room practitioners employed at the hospital; and**

**(iv) the designated employee's:**

**(A) full name;**

**(B) complete home address;**

**(C) e-mail address;**

**(C) date of birth; and**

**(D) driver license number or state identification card number;**

**(b) the designated employee has registered for an account for access to the Database and provided a unique user identification and password;**

**(c) the designated employee has passed a Database background check of available criminal court and Database records; and**

**(d) the Database has issued the designated employee a user personal identification number (PIN) and activated the employee's Database account.**

(13) The Utah Department of Health may access Database information for purposes of scientific study regarding public health. To access information, the scientific investigator shall:

(a) demonstrate to the satisfaction of the Division that the research is part of an approved project of the Utah Department of Health;

(b) provide a description of the research to be conducted, including:

(i) a research protocol for the project; and

(ii) a description of the data needed from the Database to conduct that research;

(c) provide assurances and a plan that demonstrates all Database information will be maintained securely, with access being strictly restricted to the requesting scientific investigator;

(d) provide for electronic data to be stored on a secure database computer system with access being strictly restricted to the requesting scientific investigator; and

(e) pay all relevant expenses for data transfer and manipulation.

(14) Database information that may be disseminated under Section 58-37f-301 may be disseminated by the Database staff either:

(a) verbally;

© 2013 Research is current as of July 2013. In order to ensure that the information contained herein is as current as possible, research is conducted using both nationwide legal database software, individual state legislative websites, and direct communications with state PDMP representatives. Please contact Sarah Kelsey at (703) 836-6100, ext. 119 or skelsey@namsdl.org with any additional updates or information that may be relevant to this document. This document is intended for educational purposes only and does not constitute legal advice or opinion. Headquarters Office: THE NATIONAL ALLIANCE FOR MODEL STATE DRUG LAWS. 215 Lincoln Ave. Suite 201, Santa Fe, NM 87501.

(b) by facsimile;

(c) by email;

(d) by U.S. mail; or

(e) where adequate technology is in place to ensure that a record will not be compromised, intercepted, or misdirected, by electronic access.

#### R156-37f-801a. Reporting of Information by Pharmacies Participating in the Pilot Program for Real-time Reporting.

(1) In accordance with Subsection 58-37f-801(1)(a), the pilot area is designated as the entire state of Utah. Any pharmacy or pharmacy group that submits information to the Database is eligible and may participate in the Real-time Pilot Program.

(2) In accordance with Subsection 58-37f-801(8), each licensed pharmacy participating in the pilot program for real-time reporting shall, in conjunction with controlled substance point of sale, submit from the pharmacy's database to the Controlled Substance Database, the information required by Section 58-37f-203 as implemented by Section R156-37f-203, through real-time interface and reporting software developed by the Division's contract provider.

#### R156-37f-801b. Access to Information in the Database Submitted by Pharmacies Participating in the Pilot Program for Real-time Reporting.

In accordance with Subsection 58-37f-801(8), access to information in the Database submitted by pharmacies participating in the pilot program for real-time reporting shall be the same as set forth in Section 58-37f-301 as implemented by Section R156-37f-301.

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Vermont  
18 § 4282  
18 § 4284

West's Vermont Statutes Annotated (2013)  
Title Eighteen. Health  
Part 5. Foods and Drugs  
Chapter 84A. Vermont Prescription Monitoring System

§ 4282. Definitions

As used in this chapter:

...

**(4) “Delegate” means an individual employed by a health care provider or pharmacy or in the Office of the Chief Medical Examiner and authorized by a health care provider or dispenser or by the Chief Medical Examiner to request information from the VPMS relating to a bona fide current patient of the health care provider or dispenser or to a bona fide investigation or inquiry into an individual’s death.**

...

West's Vermont Statutes Annotated (2013)  
Title Eighteen. Health  
Part 5. Foods and Drugs  
Chapter 84A. Vermont Prescription Monitoring System

§ 4284. Protection and disclosure of information

(a) The data collected pursuant to this chapter and all related information and records shall be confidential, except as provided in this chapter, and shall not be subject to the Public Records Act. The Department shall maintain procedures to protect patient privacy, ensure the confidentiality of patient information collected, recorded, transmitted, and maintained, and ensure that information is not disclosed to any person except as provided in this section.

**(b)(1) The Department shall provide only the following persons with access to query the VPMS:**

**(A) A health care provider, dispenser, or delegate who is registered with the VPMS and certifies that the requested information is for the purpose of providing medical or pharmaceutical treatment to a bona fide current patient.**



(B) Personnel or contractors, as necessary for establishing and maintaining the VPMS.

(C) The Medical Director of the Department of Vermont Health Access, for the purposes of Medicaid quality assurance, utilization, and federal monitoring requirements with respect to Medicaid recipients for whom a Medicaid claim for a Schedule II, III, or IV controlled substance has been submitted.

**(D) A medical examiner or delegate from the Office of the Chief Medical Examiner, for the purpose of conducting an investigation or inquiry into the cause, manner, and circumstances of an individual's death.**

(E) A health care provider or medical examiner licensed to practice in another state, to the extent necessary to provide appropriate medical care to a Vermont resident or to investigate the death of a Vermont resident.

(2) The Department shall provide reports of data available to the Department through the VPMS only to the following persons:

(A) A patient or that person's health care provider, or both, when VPMS reveals that a patient may be receiving more than a therapeutic amount of one or more regulated substances.

(B) A designated representative of a board responsible for the licensure, regulation, or discipline of health care providers or dispensers pursuant to a bona fide specific investigation.

(C) A patient for whom a prescription is written, insofar as the information relates to that patient.

(D) The relevant occupational licensing or certification authority if the Commissioner reasonably suspects fraudulent or illegal activity by a health care provider. The licensing or certification authority may report the data that are the evidence for the suspected fraudulent or illegal activity to a drug diversion investigator.

(E)(i) The Commissioner of Public Safety, personally, or the Deputy Commissioner of Public Safety, personally, if the Commissioner of Health, personally, or a Deputy Commissioner of Health, personally, makes the disclosure and has consulted with at least one of the patient's health care providers, when the disclosure is necessary to avert a serious and imminent threat to a person or the public.

(ii) The Commissioner of Public Safety, personally, or the Deputy Commissioner of Public Safety, personally, when he or she requests data from the Commissioner of Health, and the Commissioner of Health believes, after consultation with at least one of the patient's health care providers, that disclosure is necessary to avert a serious and imminent threat to a person or the public.

(iii) The Commissioner or Deputy Commissioner of Public Safety may disclose such data received pursuant to this subdivision (E) as is necessary, in his or her discretion, to avert the serious and imminent threat.

<Text of subsection (F) effective October 1, 2013>

(F) A prescription monitoring system or similar entity in another state pursuant to a reciprocal agreement to share prescription monitoring information with the Vermont Department of Health as described in section 4288 of this title.

(c) A person who receives data or a report from VPMS or from the Department shall not share that data or report with any other person or entity not eligible to receive that data pursuant to subsection (b) of this section, except as necessary and consistent with the purpose of the disclosure and in the normal course of business. Nothing shall restrict the right of a patient to share his or her own data.

(d) The Commissioner shall offer health care providers and dispensers training in the proper use of information they may receive from VPMS. Training may be provided in collaboration with professional associations representing health care providers and dispensers.

(e) A drug diversion investigator who may receive information pursuant to this section shall not have access to VPMS except for information provided to the officer by the licensing or certification authority.

(f) The Department is authorized to use information from VPMS for research, trend analysis, and other public health promotion purposes provided that data are aggregated or otherwise de-identified. The Department shall post the results of trend analyses on its website for use by health care providers, dispensers, and the general public. When appropriate, the Department shall send alerts relating to identified trends to health care providers and dispensers by electronic mail.

(g) Following consultation with the Unified Pain Management System Advisory Council and an opportunity for input from stakeholders, the Department shall develop a policy that will enable it to use information from VPMS to determine if individual prescribers and dispensers are using VPMS appropriately.

(h) Following consultation with the Unified Pain Management System Advisory Council and an opportunity for input from stakeholders, the Department shall develop a policy that will enable it to evaluate the prescription of regulated drugs by prescribers.

(i) Knowing disclosure of transmitted data to a person not authorized by subsection (b) of this section, or obtaining information under this section not relating to a bona fide specific investigation, shall be punishable by imprisonment for not more than one year or a fine of not more than \$1,000.00, or both, in addition to any penalties under federal law.

(j) All information and correspondence relating to the disclosure of information by the Commissioner to a patient's health care provider pursuant to subdivision (b)(2)(A) of this section shall be confidential and privileged, exempt from public inspection and copying under the Public Records Act, immune from subpoena or other disclosure, and not subject to discovery or introduction into evidence.

(k) Each request for disclosure of data pursuant to subdivision (b)(2)(B) of this section shall document a bona fide specific investigation and shall specify the case number of the investigation.

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Virginia  
§ 54.1-2523.2

West's Annotated Code of Virginia (2013)

Title 54.1. Professions and Occupations

Subtitle III. Professions and Occupations Regulated by Boards Within the Department of Health Professions

Chapter 25.2. Prescription Monitoring Program

§ 54.1-2523.2. Authority to access database

**Any prescriber authorized to access the information in the possession of the Prescription Monitoring Program pursuant to this chapter may, pursuant to regulations promulgated by the Director to implement the provisions of this section, delegate such authority to health care professionals who are (i) licensed, registered, or certified by a health regulatory board under the Department of Health Professions and (ii) employed at the same facility and under the direct supervision of the prescriber.**

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Washington  
ADC 246-470-050

Washington Administrative Code (2013)  
Title 246. Health, Department of  
Chapter 246-470. Prescription Monitoring Program

246-470-050. Pharmacist, prescriber or other health care practitioner access to information from the program.

**A pharmacist, prescriber, or licensed health care practitioner authorized by a prescriber may obtain prescription monitoring information relating to their patients, for the purpose of providing medical or pharmaceutical care.**

**(1) Registration for access. A pharmacist, prescriber, or licensed health care practitioner authorized by a prescriber shall register with the department in order to receive an authentication to access the electronic system. The registration process shall be established by the department.**

**(2) Verification by the department. The department shall verify the authentication and identity of the pharmacist, prescriber, or licensed health care practitioner authorized by a prescriber before allowing access to any prescription monitoring information.**

**(3) Procedure for accessing prescription information. A pharmacist, prescriber, or licensed health care practitioner authorized by a prescriber may access information from the program electronically, using the authentication issued by the department.**

**(4) A pharmacist, prescriber, or licensed health care practitioner authorized by a prescriber may alternately submit a written request via mail or facsimile transmission in a manner and format established by the department.**

**(5) Reporting lost or stolen authentication. If the authentication issued by the department is lost, missing, or the security of the authentication is compromised, the pharmacist, prescriber, or licensed health care practitioner authorized by a prescriber shall notify the department by telephone and in writing as soon as reasonably possible.**

**(6) All requests for, uses of, and disclosures of prescription monitoring information by authorized persons must be consistent with the program's mandate as outlined in RCW 70.225.040 and this chapter.**

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West Virginia  
§ 60A-9-5  
ADC § 15-8-7

West's Annotated Code of West Virginia (2013)  
Chapter 60A. Uniform Controlled Substances Act  
Article 9. Controlled Substances Monitoring

§ 60A-9-5. Confidentiality; limited access to records; period of retention; no civil liability for required reporting

(a)(1) The information required by this article to be kept by the State Board of Pharmacy is confidential and not subject to the provisions of chapter twenty-nine-b of this code or obtainable as discovery in civil matters absent a court order and is open to inspection only by inspectors and agents of the State Board of Pharmacy, members of the West Virginia State Police expressly authorized by the Superintendent of the West Virginia State Police to have access to the information, authorized agents of local law-enforcement agencies as members of a federally affiliated drug task force, authorized agents of the federal Drug Enforcement Administration, duly authorized agents of the Bureau for Medical Services, duly authorized agents of the Office of the Chief Medical Examiner for use in post-mortem examinations, duly authorized agents of licensing boards of practitioners in this state and other states authorized to prescribe Schedules II, III and IV controlled substances, prescribing practitioners and pharmacists and persons with an enforceable court order or regulatory agency administrative subpoena: Provided, That all law-enforcement personnel who have access to the Controlled Substances Monitoring Program database shall be granted access in accordance with applicable state laws and Board of Pharmacy legislative rules, shall be certified as a West Virginia law-enforcement officer and shall have successfully completed United States Drug Enforcement Administration Diversion Training and National Association of Drug Diversion Investigation Training. All information released by the State Board of Pharmacy must be related to a specific patient or a specific individual or entity under investigation by any of the above parties except that practitioners who prescribe or dispense controlled substances may request specific data related to their Drug Enforcement Administration controlled substance registration number or for the purpose of providing treatment to a patient: Provided, however, That the West Virginia Controlled Substances Monitoring Program Database Review Committee established in subsection (b) of this section is authorized to query the database to comply with said subsection.

...

**(f) Persons or entities with access to the West Virginia Controlled Substances Monitoring Program database pursuant to this section may, pursuant to rules promulgated by the Board of Pharmacy, delegate appropriate personnel to have access to said database;**

...

West Virginia Code of State Rules (2013)  
Title 15. West Virginia Board of Pharmacy  
Legislative Rule (Ser. 8)  
Series 8. Controlled Substances Monitoring

§ 15-8-7. Confidentiality.

7.1. The board shall carry out a program to protect the confidentiality of the information received by the central repository.

7.2. The board may disclose confidential information received by the central repository to any person who is engaged in receiving, processing, or storing the information.

**7.3. The board may release confidential information received by the central repository to the following persons:**

(a) A duly authorized agent of a board in this state or another state that licenses practitioners authorized to prescribe Schedules II, III, and IV controlled substances who is engaged in an investigation, an adjudication, or a prosecution of a violation under any state or federal law that involves a controlled substance;

(b) Members of the West Virginia State Police expressly authorized by the superintendent of the West Virginia State Police to have access to the information;

(c) An authorized agent of a local law-enforcement agency who is acting as a member of a Federally affiliated drug task force;

(d) Authorized agents of the federal Drug Enforcement Administration;

(e) The Chief Medical Examiner for the State of West Virginia or his or her duly authorized agent for use in post-mortem examinations;

(f) A person with an enforceable court order or regulatory agency administrative subpoena;

(g) Inspectors and agents of the Board;

**(h) Prescribing practitioners or their duly authorized agents;**

**(i) Pharmacists or a registered pharmacy technician as the agent of the pharmacist; and**

(j) A person using the data for compilation of educational, scholarly, or statistical purposes so long as the individually identifiable data of the persons or entities stored in the central repository remains confidential.

7.4. All information released by the board shall be related to a specific patient or a specific individual or entity under investigation by any of the persons set forth in subsection 7.3 (a) through (i) of this section except that practitioners who prescribe controlled substances may request specific data related to their drug enforcement administration controlled substance registration number or for the purpose of providing treatment to a patient.

7.5. All access to the data collected by the central repository shall be limited to regular business hours of the board's office unless an individual authorized to receive the information proves that an immediate danger to the public exists and immediate access is necessary to prevent further harm, Provided That the board may permit access at any time to authorized users through the use of a secure connection and through the use of proper security features designed to protect the integrity and confidentiality of the information from unauthorized access or disclosure.

**7.6. Any person or entity having access to the central repository and who is permitted to designate a duly authorized agent to have access to the central repository pursuant to this rule shall make the designation on a form to be supplied by the board. It is the responsibility of the designating individual to insure that the designated agent maintains the confidentiality of the information in the central repository as required. Further, should the designating individual remove the authority of the designated agent to act as the duly authorized agent, or should the designated agent leave the employment of the designating individual or entity such that he or she is no longer eligible to act as the duly authorized agent, then the designating individual shall immediately notify the board, at which time the designee's access to the central repository shall be removed.**

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Wisconsin  
ADC Phar. 18.09

Wisconsin Administrative Code (2013)  
Pharmacy Examining Board  
Chapter Phar 18. Prescription Drug Monitoring Program

Phar 18.09 Direct access to PDMP information.

**(1) Dispensers, dispenser delegates, practitioners, and practitioner delegates may access PDMP information in the same or similar manner, and for the same or similar purposes, as those persons are authorized to access similar confidential patient health care records under ss. 146.82 and 450.19, Stats., this chapter and other state or federal laws and regulations relating to the privacy of patient health care records.**

**(2) To obtain access to PDMP information, dispensers, dispenser delegates, practitioners, and practitioner delegates shall create an account with the board on a form provided by the board.**

**Note: The application to create an account may be completed online at [www.dsps.wi.gov](http://www.dsps.wi.gov) or obtained at no charge from the Department of Safety and Professional Services, 1400 East Washington Avenue, P.O. Box 8935, Madison, WI 53708.**

(3) The board may deny, suspend, revoke or otherwise restrict or limit a dispenser's, dispenser delegate's, practitioner's, or practitioner delegate's direct access to PDMP information for any of the following reasons:

(a) The dispenser, dispenser delegate, practitioner, or practitioner delegate uses PDMP information in violation of s. 146.82 or 450.19, Stats., this chapter, or other state or federal laws or regulations relating to the privacy of patient health care records.

(b) The dispenser, dispenser delegate, practitioner, or practitioner delegate is no longer licensed in this state or another state and recognized by this state as a person authorized to prescribe or dispense monitored prescription drugs.

(c) The board, other licensing board, or regulatory agency takes adverse action against the dispenser, dispenser delegate, practitioner, or practitioner delegate.

(d) A licensing board or equivalent regulatory agency in another jurisdiction takes adverse action against the dispenser, dispenser delegate, practitioner, or practitioner delegate.

(e) The federal department of justice, drug enforcement administration takes adverse action against the dispenser, dispenser delegate, practitioner, or practitioner delegate.

(f) The dispenser, dispenser delegate, practitioner, or practitioner delegate is convicted of a crime substantially related to the prescribing or dispensing of a monitored prescription drug.

(g) The dispenser delegate or practitioner delegate is no longer delegated the task of inputting or accessing PDMP information.

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