

# NAMSDL



**National Alliance for Model State Drug Laws**

## **STATES THAT REQUIRE CERTAIN AUTHORIZED RECIPIENTS TO UNDERGO TRAINING AND/OR COMPLETION OF EDUCATIONAL COURSES BEFORE ACCESSING PMP DATA**

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## Introduction

This compilation of statutes and regulations only includes those states that *require* some type of training or educational course by certain authorized users before they can access the prescription monitoring program. Most states offer *optional* training courses for users, and those states are not included in this document.

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Arizona  
ADC R4-23-501

Arizona Administrative Code (2013)  
Title 4. Professions and Occupations  
Chapter 23. Board of Pharmacy  
Article 5. Controlled Substances Prescription Monitoring Program

R4-23-501. Controlled Substances Prescription Monitoring Program Registration

A. Under A.R.S. § 36-2606, a medical practitioner who is issued a license under A.R.S. Title 32, Chapter 7, 11, 13, 14, 15, 16, 17, 21, 25, or 29 and possesses a current DEA registration under the Federal Controlled Substances Act shall have a current CSPMP registration issued by the Board.

B. Application. To obtain a CSPMP registration, a person shall submit a completed application on a form furnished by the Board that includes:

1. Applicant's name, address, mailing address, if different, e-mail address, telephone number, facsimile number, license number issued under A.R.S. Title 32, Chapter 7, 11, 13, 14, 15, 16, 17, 21, 25, or 29, and DEA registration number;
2. Whether the applicant's license and DEA registration listed in subsection (B)(1) are current and in good standing, and if not, the status of the license and registration; and
3. Date signed and applicant's verified signature.

C. Registration. Within seven business days of receipt of a completed application specified in subsection (B), the Board office shall determine whether an application is complete. If the application is complete, the Board office shall issue a registration number and mail a current renewal receipt to the applicant. If the application is incomplete, the Board office shall issue a notice of incompleteness. An applicant with an incomplete application shall comply with the requirements of R4-23-502(F)(2) and (3).

D. Registration renewal. As specified in A.R.S. § 36-2606(C), the Board shall automatically suspend the registration of any registrant that fails to renew the registration on or before May 1 of the year in which the renewal is due. The Board shall vacate a suspension if the registrant submits a renewal application. A suspended registrant is prohibited from accessing information in the prescription monitoring program database.

E. CSPMP database access.

**1. A medical practitioner that chooses to use the CSPMP database shall request access from the CSPMP Director on forms provided on the Board's web site. Upon receipt of the request, the CSPMP Director or designee shall issue access credentials provided the medical practitioner is in compliance with the registration requirements of this Section and has completed the Board's CSPMP Online Training Program.**

**2. A pharmacist that chooses to use the CSPMP database shall request access from the CSPMP Director on forms provided on the Board's web site. Upon receipt of the request, the CSPMP Director or designee shall issue access credentials provided the pharmacist has a current active pharmacist license and has completed the Board's CSPMP Online Training Program.**

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# Kentucky

§ 218A.202

§ 218A.240

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

(1) The Cabinet for Health and Family Services shall establish an electronic system for monitoring Schedules II, III, IV, and V controlled substances that are dispensed within the Commonwealth by a practitioner or pharmacist or dispensed to an address within the Commonwealth by a pharmacy that has obtained a license, permit, or other authorization to operate from the Kentucky Board of Pharmacy. The cabinet may contract for the design, upgrade, or operation of this system if the contract preserves all of the rights, privileges, and protections guaranteed to Kentucky citizens under this chapter and the contract requires that all other aspects of the system be operated in conformity with the requirements of this or any other applicable state or federal law.

(2) A practitioner or a pharmacist authorized to prescribe or dispense controlled substances to humans shall register with the cabinet to use the system provided for in this section and shall maintain such registration continuously during the practitioner's or pharmacist's term of licensure and shall not have to pay a fee or tax specifically dedicated to the operation of the system.

(3) Every dispenser within the Commonwealth who is licensed, permitted, or otherwise authorized to prescribe or dispense a controlled substance to a person in Kentucky shall report to the Cabinet for Health and Family Services the data required by this section except that reporting shall not be required for:

(a) A drug administered directly to a patient in a hospital, a resident of a health care facility licensed under KRS Chapter 216B, a resident of a child-caring facility as defined by KRS 199.011, or an individual in a jail, correctional facility, or juvenile detention facility;

(b) A drug, other than any Schedule II controlled substance or a Schedule III controlled substance containing hydrocodone, dispensed by a practitioner at a facility licensed by the cabinet, provided that the quantity dispensed is limited to an amount adequate to treat the patient for a maximum of forty-eight (48) hours; or

(c) A drug administered or dispensed to a research subject enrolled in a research protocol approved by an institutional review board that has an active federalwide assurance number from

the United States Department of Health and Human Services, Office for Human Research Protections where the research involves single, double, or triple blind drug administration or is additionally covered by a certificate of confidentiality from the National Institutes of Health.

(4) Data for each controlled substance that is dispensed shall include but not be limited to the following:

- (a) Patient identifier;
- (b) National drug code of the drug dispensed;
- (c) Date of dispensing;
- (d) Quantity dispensed;
- (e) Prescriber; and
- (f) Dispenser.

(5) The data shall be provided in the electronic format specified by the Cabinet for Health and Family Services unless a waiver has been granted by the cabinet to an individual dispenser. The cabinet shall establish acceptable error tolerance rates for data. Dispensers shall ensure that reports fall within these tolerances. Incomplete or inaccurate data shall be corrected upon notification by the cabinet if the dispenser exceeds these error tolerance rates.

(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.

(7) The Department for Medicaid Services shall use any data or reports from the system for the purpose of identifying Medicaid providers or recipients whose prescribing, dispensing, or usage of controlled substances may be:

(a) Appropriately managed by a single outpatient pharmacy or primary care physician; or

(b) Indicative of improper, inappropriate, or illegal prescribing or dispensing practices by a practitioner or drug seeking by a Medicaid recipient.

(8) A person who receives data or any report of the system from the cabinet shall not provide it to any other person or entity except as provided in this section, in another statute, or by order of a court of competent jurisdiction and only to a person or entity authorized to receive the data or the report under this section, except that:

(a) A person specified in subsection (6)(b) of this section who is authorized to receive data or a report may share that information with any other persons specified in subsection (6)(b) of this section authorized to receive data or a report if the persons specified in subsection (6)(b) of this section are working on a bona fide specific investigation involving a designated person. Both the person providing and the person receiving the data or report under this paragraph shall document in writing each person to whom the data or report has been given or received and the day, month, and year that the data or report has been given or received. This document shall be maintained in a file by each agency engaged in the investigation;

(b) A representative of the Department for Medicaid Services may share data or reports regarding overutilization by Medicaid recipients with a board designated in subsection (6)(a) of this section, or with a law enforcement officer designated in subsection (6)(b) of this section;

(c) The Department for Medicaid Services may submit the data as evidence in an administrative hearing held in accordance with KRS Chapter 13B;

(d) If a state licensing board as defined in Section 4 of this Act initiates formal disciplinary proceedings against a licensee, and data obtained by the board is relevant to the charges, the board may provide the data to the licensee and his or her counsel, as part of the notice process required by KRS 13B.050, and admit the data as evidence in an administrative hearing conducted pursuant to KRS Chapter 13B, with the board and licensee taking all necessary steps to prevent further disclosure of the data; and

(e) A practitioner, pharmacist, or employee who obtains data under subsection (6)(e) of this section may share the report with the patient or person authorized to act on the patient's behalf and place the report in the patient's medical record, with that individual report then being deemed a medical record subject to disclosure on the same terms and conditions as an ordinary medical record in lieu of the disclosure restrictions otherwise imposed by this section.

(9) The Cabinet for Health and Family Services, all peace officers specified in subsection (6)(b) of this section, all officers of the court, and all regulatory agencies and officers, in using the data for investigative or prosecution purposes, shall consider the nature of the prescriber's and dispenser's practice and the condition for which the patient is being treated.

(10) The data and any report obtained therefrom shall not be a public record, except that the Department for Medicaid Services may submit the data as evidence in an administrative hearing held in accordance with KRS Chapter 13B.

(11) Intentional failure by a dispenser to transmit data to the cabinet as required by subsection (3), (4), or (5) of this section shall be a Class B misdemeanor for the first offense and a Class A misdemeanor for each subsequent offense.

(12) Intentional disclosure of transmitted data to a person not authorized by subsection (6) to subsection (8) of this section or authorized by KRS 315. 121, or obtaining information under this section not relating to a bona fide specific investigation, shall be a Class B misdemeanor for the first offense and a Class A misdemeanor for each subsequent offense.

(13) (a) The Commonwealth Office of Technology, in consultation with the Cabinet for Health and Family Services, may submit an application to the United States Department of Justice for a drug diversion grant to fund a pilot or continuing project to study, create, or maintain a real-time electronic monitoring system for Schedules II, III, IV, and V controlled substances.

(b) The pilot project shall:

1. Be conducted in two (2) rural counties that have an interactive real-time electronic information system in place for monitoring patient utilization of health and social services through a federally funded community access program; and

2. Study the use of an interactive system that includes a relational data base with query capability.

(c) Funding to create or maintain a real-time electronic monitoring system for Schedules II, III, IV, and V controlled substances may be sought for a statewide system or for a system covering any geographic portion or portions of the state.

(14) Provisions in this section that relate to data collection, disclosure, access, and penalties shall apply to the pilot project authorized under subsection (13) of this section.

(15) The Cabinet for Health and Family Services may, by promulgating an administrative regulation, limit the length of time that data remain in the electronic system. Any data removed from the system shall be archived and subject to retrieval within a reasonable time after a request from a person authorized to review data under this section.

(16) (a) The Cabinet for Health and Family Services shall work with each board responsible for the licensure, regulation, or discipline of practitioners, pharmacists, or other persons who are authorized to prescribe, administer, or dispense controlled substances for the development of a continuing education program about the purposes and uses of the electronic system for monitoring established in this section.

(b) The cabinet shall work with the Kentucky Bar Association for the development of a continuing education program for attorneys about the purposes and uses of the electronic system for monitoring established in this section.

(c) The cabinet shall work with the Justice and Public Safety Cabinet for the development of a continuing education program for law enforcement officers about the purposes and uses of the electronic system for monitoring established in this section.

(17) If the cabinet becomes aware of a prescriber's or dispenser's failure to comply with this section, the cabinet shall notify the licensing board or agency responsible for licensing the prescriber or dispenser. The licensing board shall treat the notification as a complaint against the licensee.

(18) The cabinet shall promulgate administrative regulations to implement the provisions of this section. Included in these administrative regulations shall be:

(a) An error resolution process allowing a patient to whom a report had been disclosed under subsection (8) of this section to request the correction of inaccurate information contained in the system relating to that patient; and

(b) Beginning July 1, 2013, a requirement that data be reported to the system under subsection (3) of this section within one (1) day of dispensing.

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

§ 218A.240 Controlled substances; duties and authority of state and local officers, Cabinet for Health and Family Services, and Kentucky Board of Pharmacy; civil proceedings; identification of trends; identification of prescribers, dispensers, and patients for licensing board; review of hospital's or health care facility's prescribing and dispensing practices

(1) All police officers and deputy sheriffs directly employed full-time by state, county, city, urban-county, or consolidated local governments, the Department of Kentucky State Police, the Cabinet for Health and Family Services, their officers and agents, and of all city, county, and Commonwealth's attorneys, and the Attorney General, within their respective jurisdictions, shall enforce all provisions of this chapter and cooperate with all agencies charged with the enforcement of the laws of the United States, of this state, and of all other states relating to controlled substances.

(2) For the purpose of enforcing the provisions of this chapter, the designated agents of the Cabinet for Health and Family Services shall have the full power and authority of peace officers in this state, including the power of arrest and the authority to bear arms, and shall have the power and authority to administer oaths; to enter upon premises at all times for the purpose of making inspections; to seize evidence; to interrogate all persons; to require the production of prescriptions, of books, papers, documents, or other evidence; to employ special investigators; and to expend funds for the purpose of obtaining evidence and to use data obtained under KRS 218A.202(7) in any administrative proceeding before the cabinet.

(3) The Kentucky Board of Pharmacy, its agents and inspectors, shall have the same powers of inspection and enforcement as the Cabinet for Health and Family Services.

(4) Designated agents of the Cabinet for Health and Family Services and the Kentucky Board of Pharmacy are empowered to remove from the files of a pharmacy or the custodian of records for that pharmacy any controlled substance prescription or other controlled substance record upon tendering a receipt. The receipt shall be sufficiently detailed to accurately identify the record. A receipt for the record shall be a defense to a charge of failure to maintain the record.

(5) Notwithstanding the existence or pursuit of any other remedy, civil or criminal, any law enforcement authority may maintain, in its own name, an action to restrain or enjoin any violation of this chapter or to forfeit any property subject to forfeiture under KRS 218A.410, irrespective of whether the owner of the property has been charged with or convicted of any offense under this chapter.

(a) Any civil action against any person brought pursuant to this section may be instituted in the Circuit Court in any county in which the person resides, in which any property owned by the person and subject to forfeiture is found, or in which the person has violated any provision of this chapter.

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(b) A final judgment rendered in favor of the Commonwealth in any criminal proceeding brought under this chapter shall estop the defendant from denying the essential allegations of the criminal offense in any subsequent civil proceeding brought pursuant to this section.

(c) The prevailing party in any civil proceeding brought pursuant to this section shall recover his or her costs, including a reasonable attorney's fee.

(d) Distribution of funds under this section shall be made in the same manner as in KRS 218A.420, except that if the Commonwealth's attorney has not initiated the forfeiture action under this section, his or her percentage of the funds shall go to the agency initiating the forfeiture action.

(6) The Cabinet for Health and Family Services shall make or cause to be made examinations of samples secured under the provisions of this chapter to determine whether any provision has been violated.

(7) (a) The Cabinet for Health and Family Services shall proactively use the data compiled in the electronic system created in KRS 218A.202 for investigations, research, statistical analysis, and educational purposes and shall proactively identify trends in controlled substance usage and other potential problem areas. **Only cabinet personnel who have undergone training for the electronic system and who have been approved to use the system shall be authorized access to the data and reports under this subsection.** The cabinet shall notify a state licensing board listed in KRS 218A.205 if a report or analysis conducted under this subsection indicates that further investigation about improper, inappropriate or illegal prescribing or dispensing may be necessary by the board. The board shall consider each report and may, after giving due consideration to areas of practice, specialties, board certifications, and appropriate standards of care, request and receive a follow-up report or analysis containing relevant information as to the prescriber or dispenser and his or her patients.

(b) The cabinet shall develop criteria, in collaboration with the Board of Medical Licensure, the Board of Nursing, the Office of Drug Control Policy, and the Board of Pharmacy, to be used to generate public trend reports from the data obtained by the system. Meetings at which the criteria are developed shall be meetings, as defined in KRS 61.805, that comply with the open meetings laws, KRS 61.805 to 61.850. The cabinet shall, on a quarterly basis, publish trend reports from the data obtained by the system. Except as provided in subsection (8) of this section, these trend reports shall not identify an individual prescriber, dispenser, or patient. Peace officers authorized to receive data under KRS 218A.202 may request trend reports not specifically published pursuant to this paragraph except that the report shall not identify an individual prescriber, dispenser, or patient.

(8) If the cabinet deems it to be necessary and appropriate, upon the request of a state licensing board listed in KRS 218A.205, the cabinet shall provide the requesting board with the identity of prescribers, dispensers, and patients used to compile a specific trend report.

(9) Any hospital or other health care facility may petition the cabinet to review data from the electronic system specified in KRS 218A.202 as it relates to employees of that facility to determine if inappropriate prescribing or dispensing practices are occurring. The cabinet may initiate any investigation in such cases as he or she determines is appropriate, and may request the assistance from the hospitals or health care facilities in the investigation.

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# Louisiana

40 § 1007

40 § 1008

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.

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

§ 1008. Education and treatment

**A. The board shall, in consultation with and upon the recommendation of the advisory council, implement the following education courses:**

**(1) An orientation course during the implementation phase of the prescription monitoring program.**

**(2) A course for persons who are authorized to access the prescription monitoring information, but who did not participate in the orientation course.**

**(3) A course for persons who are authorized to access the prescription monitoring information, but who have violated the laws or breached occupational standards involving the prescribing, dispensing, or use of any controlled substances or drugs monitored by the prescription monitoring program.**

**(4) A continuing education course for health care providers or professionals on prescribing practices, pharmacology, and the identification, treatment, and referral of a patient**

**addicted to or abusing controlled substances or drugs monitored by the prescription monitoring program.**

B. The board shall, in consultation with and upon recommendation of the advisory council, implement an educational program to inform the public about the use, diversion and abuse of, addiction to, and treatment for the addiction to controlled substances or drugs monitored by the prescription monitoring program.

C. The board shall, upon reasonable suspicion, refer potential or alleged impaired prescribers and dispensers to the appropriate professional licensing or certification agency to ensure intervention, treatment, and ongoing monitoring and follow-up.

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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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## Montana

Per the state PDMP representative, Montana requires all authorized users to complete a training program before being granted access.

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Nevada  
§ 453.1545

West's Nevada Revised Statutes Annotated (2013)  
Title 40. Public Health and Safety  
Chapter 453. Controlled Substances  
Uniform Controlled Substances Act  
General Provisions

§ 453.1545. Board and Division required to develop computerized program to track prescriptions for controlled substances and course of training for persons who access program; Board required to provide certain practitioners Internet access to database of program; reporting of illegal activity; agreements with state agency to receive or exchange information obtained by program; confidentiality of information obtained from program; immunity from liability for practitioner who transmits certain required information and reports; gifts, grants and donations

1. The Board and the Division shall cooperatively develop a computerized program to track each prescription for a controlled substance listed in schedule II, III or IV that is filled by a pharmacy that is registered with the Board or that is dispensed by a practitioner who is registered with the Board. The program must:

(a) Be designed to provide information regarding:

(1) The inappropriate use by a patient of controlled substances listed in schedules II, III and IV to pharmacies, practitioners and appropriate state agencies to prevent the improper or illegal use of those controlled substances; and

(2) Statistical data relating to the use of those controlled substances that is not specific to a particular patient.

(b) Be administered by the Board, the Division, the Health Division of the Department and various practitioners, representatives of professional associations for practitioners, representatives of occupational licensing boards and prosecuting attorneys selected by the Board and the Division.

(c) Not infringe on the legal use of a controlled substance for the management of severe or intractable pain.

(d) Include the contact information of each person who elects to access the database of the program pursuant to subsection 2, including, without limitation:

(1) The name of the person;

- (2) The physical address of the person;
- (3) The telephone number of the person; and
- (4) If the person maintains an electronic mail address, the electronic mail address of the person.

**2. The Board shall provide Internet access to the database of the program established pursuant to subsection 1 to each practitioner who is authorized to write prescriptions for and each person who is authorized to dispense controlled substances listed in schedule II, III or IV who:**

- (a) Elects to access the database of the program; and

**(b) Completes the course of instruction described in subsection 7.**

3. The Board and the Division must have access to the program established pursuant to subsection 1 to identify any suspected fraudulent or illegal activity related to the dispensing of controlled substances.

4. The Board or the Division shall report any activity it reasonably suspects may be fraudulent or illegal to the appropriate law enforcement agency or occupational licensing board and provide the law enforcement agency or occupational licensing board with the relevant information obtained from the program for further investigation.

5. The Board and the Division may cooperatively enter into a written agreement with an agency of any other state to provide, receive or exchange information obtained by the program with a program established in that state which is substantially similar to the program established pursuant to subsection 1, including, without limitation, providing such state access to the database of the program or transmitting information to and receiving information from such state. Any information provided, received or exchanged as part of an agreement made pursuant to this section may only be used in accordance with the provisions of this chapter.

6. Information obtained from the program relating to a practitioner or a patient is confidential and, except as otherwise provided by this section and NRS 239.0115, must not be disclosed to any person. That information must be disclosed:

- (a) Upon the request of a person about whom the information requested concerns or upon the request on behalf of that person by his or her attorney; or
- (b) Upon the lawful order of a court of competent jurisdiction.

**7. The Board and the Division shall cooperatively develop a course of training for persons who elect to access the database of the program pursuant to subsection 2 and require each such person to complete the course of training before the person is provided with Internet access to the database pursuant to subsection 2.**

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8. A practitioner who is authorized to write prescriptions for each person who is authorized to dispense controlled substances listed in schedule II, III or IV who acts with reasonable care when transmitting to the Board or the Division a report or information required by this section or a regulation adopted pursuant thereto is immune from civil and criminal liability relating to such action.

9. The Board and the Division may apply for any available grants and accept any gifts, grants or donations to assist in developing and maintaining the program required by this section.

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## New Jersey

Pursuant to the state PMP representative, New Jersey requires authorized users to receive training prior to accessing the PMP database.

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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.

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### 16.19.29.12 REGISTRATION FOR ACCESS TO PRESCRIPTION INFORMATION:

A. Practitioners with individual drug enforcement administration (DEA) issued numbers will complete and submit a hard copy written, signed and notarized application. After verification of submitted information, a username and password will be issued to the practitioner. One subaccount per practitioner account is authorized for an agent of the practitioner. The agent designated by the practitioner will complete and submit a hard copy written, signed and

notarized application. After verification of submitted information, a username and password will be issued to the agent.

B. Pharmacies with DEA issued numbers will complete and submit a hard copy written, signed and notarized application. After verification of submitted information, a username and password will be issued. Pharmacies will designate one individual who will complete and submit a hard copy written, signed and notarized application. After verification of submitted information, a username and password will be issued to the individual. Pharmacies will not be permitted to obtain a subaccount.

C. All registrations will be renewed every three years by completing and submitting a new application.

**D. All registrants to the prescription monitoring program will complete a web based training program approved by the board.**

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# Ohio

ADC 4723-9-02

ADC 4723-9-13

Baldwin's Ohio Administrative Code Annotated (2013)

4723 Nursing Board

Chapter 4723-9. Prescriptive Authority

4723-9-02 Requirements for a course of study in advanced pharmacology

(A) To be acceptable to the board, a course of study shall meet the following requirements:

(1) Be a minimum of forty-five contact hours in length and include content which ensures sufficient preparation for the safe and effective prescribing of drugs and therapeutic devices;

(2) Include content which is specific to the participant's nursing specialty and which includes all of the following:

(a) A minimum of thirty-six hours of training, obtained from a single provider, in:

(i) Pharmacokinetic principles and clinical application; and

(ii) Principles of the use of drugs and therapeutic devices in the prevention of illness and maintenance of health;

(b) A combined six hours of instruction in:

(i) The fiscal and ethical implications of prescribing drugs and therapeutic devices; and

(ii) The state and federal laws that apply to the authority to prescribe;

**(c) Six hours of instruction that is specific to schedule II controlled substances, including instruction in all of the following:**

(i) Indications and contraindications for the use of schedule II controlled substances in drug therapies, including risk, evaluation and mitigation strategies for the use of opiates in the treatment of chronic pain for non-terminal conditions, and the need for periodic assessment and documentation of the patient's functional status;

(ii) The most recent guidelines and recommendations for pain management therapies and education, as established by state and national organizations such as the Ohio pain initiative, the American pain society and the United States food and drug administration (FDA);

- (iii) The most recent guidelines and recommendations for stimulant therapies utilized in the management of attention-deficit or hyperactivity disorder, as adopted by state and national organizations such as the American academy of pediatrics;
  - (iv) Fiscal and ethical implications of prescribing schedule II controlled substances;
  - (v) State and federal laws that apply to the authority to prescribe schedule II controlled substances, including state medical board of Ohio rules governing controlled substances and the treatment of chronic pain, and Ohio state board of pharmacy rules governing the manner of issuance of a prescription;
  - (vi) Prevention of abuse and diversion of schedule II controlled substances, including identification of the risk of abuse, addiction and diversion, recognition of abuse, addiction and diversion, types of assistance available for prevention of abuse, addiction and diversion, **the use of the Ohio automated rx reporting system ( OARRS)**, and other methods of establishing safeguards against abuse and diversion; and
- (d) Up to three hours of instruction specific to schedule II controlled substances as set forth in paragraphs (A)(2)(c)(iii) and (A)(2)(c)(iv) of this rule may be credited toward satisfying the six hours of instruction required by paragraphs (A)(2)(b)(i) and (A)(2)(b)(ii) of this rule.
- (3) Include a process for interaction of the participants with instructional personnel;
  - (4) Include a process for evaluating the participants' learning of the content required by this rule that includes:
    - (a) Successful completion of case studies or written assignments;
    - (b) Successful completion of a comprehensive written examination;
    - (c) A mechanism to assure the security of the evaluation process; and
  - (5) Be faculty-directed and obtained either from:
    - (a) An accredited educational institution acceptable to the board; or
    - (b) A continuing education program in pharmacology that meets the requirements of Chapter 4723-14 of the Administrative Code.

Baldwin's Ohio Administrative Code Annotated (2013)  
4723 Nursing Board  
Chapter 4723-9. Prescriptive Authority

4723-9-13 Instruction specific to schedule II controlled substances

(A) All clinical nurse specialists, certified nurse practitioners, and certified nurse-midwives, who hold a certificate to prescribe, including an externship certificate, prior to the effective date of this rule must obtain six contact hours of instruction specific to schedule II controlled substances on or before August 31, 2013 in order to be eligible to renew their certificate to prescribe, and present documentation satisfactory to the board of having completed the instruction.

**(B) To meet this requirement, the course of instruction must:**

**(1) Include the following content:**

(a) Indications and contraindications for the use of schedule II controlled substances in drug therapies, including risk, evaluation and mitigation strategies for the use of opiates in the treatment of chronic pain for non-terminal conditions, and the need for periodic assessment and documentation of the patient's functional status;

(b) The most recent guidelines and recommendations for pain management therapies and education, as established by state and national organizations such as the Ohio pain initiative, the American pain society and the United States food and drug administration (FDA);

(c) The most recent guidelines and recommendations for stimulant therapies utilized in the management of attention-deficit or hyperactivity disorder, as adopted by state and national organizations such as the American academy of pediatrics;

(d) Fiscal and ethical implications of prescribing schedule II controlled substances;

(e) State and federal laws that apply to the authority to prescribe schedule II controlled substances, including state medical board of Ohio rules governing controlled substances and the treatment of chronic pain, and Ohio state board of pharmacy rules governing the manner of issuance of a prescription; and

(f) Prevention of abuse and diversion of schedule II controlled substances, including identification of the risk of abuse, addiction and diversion, recognition of abuse, addiction and diversion, types of assistance available for prevention of abuse, addiction and diversion, **the use of the Ohio automated rx reporting system ( OARRS)**, and other methods of establishing safeguards against abuse and diversion;

(2) Meet the requirements of Chapter 4723-14 of the Administrative Code; and

(3) Be at minimum six hours in length.

(C) Applicants must submit documentation of successful completion to the board in the form of an original certificate, issued by the provider of the course of instruction that includes:

(1) Name of the attendee;

(2) Title of the program;

(3) Date of the program;

(4) Name and address of the provider and OBN approver number, if applicable; and

(5) Verification of completion of at least six hours each of sixty minutes in duration.

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## Pennsylvania

Pursuant to the PMP administrator, Pennsylvania requires authorized users to undergo training in the use of the PMP.

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## South Carolina

Pursuant to their website, the South Carolina Department of Health and Environmental Control requires all practitioners and pharmacists requesting direct access to the PMP database to complete a SCRIPTS online training course prior to submitting an access request form.

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# Utah

§ 58-37f-402

West's Utah Code Annotated (2013)  
Title 58. Occupations and Professions  
Chapter 37F. Controlled Substance Database Act  
Part 4. Registration and Training

§ 58-37f-402. Online tutorial and test relating to the database--Fees--Rulemaking authority--  
Continuing professional education credit

**(1) The division shall develop an online tutorial and an online test for registration to use the database that provides instruction regarding, and tests, the following:**

**(a) the purpose of the database;**

**(b) how to access and use the database;**

**(c) the law relating to:**

**(i) the use of the database; and**

**(ii) the information submitted to, and obtained from, the database; and**

**(d) basic knowledge that is important for all people who prescribe controlled substances to know in order to help ensure the health and safety of an individual to whom a controlled substance is prescribed.**

**(2) The division shall design the test described in this section as follows:**

**(a) an individual shall answer all of the questions correctly in order to pass the test;**

**(b) an individual shall be permitted to immediately retake the portion of the test that the individual answers incorrectly as many times as necessary for the individual to pass the test; and**

**(c) after an individual takes the test, the test software shall:**

**(i) immediately inform the individual of the number of questions that were answered incorrectly;**

**(ii) provide the correct answers;**

- (iii) replay the portion of the tutorial that relates to the incorrectly answered questions; and**
- (iv) ask the individual the incorrectly answered questions again.**

**(3) The division shall design the tutorial and test so that it is possible to take the tutorial and complete the test in 20 minutes or less, if the individual answers all of the questions correctly on the first attempt.**

**(4) The division shall ensure that the tutorial and test described in this section are fully functional and available for use online on or before November 1, 2010.**

**(5) The division shall impose a fee, in accordance with Section 63J-1-504, on an individual who takes the test described in this section, to pay the costs incurred by the division to:**

- (a) develop, implement, and administer the tutorial and test described in this section; and**
- (b) fulfill the other duties imposed on the division under this part.**

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

- (a) develop, implement, and administer the tutorial and test described in this section; and**
- (b) fulfill the other duties imposed on the division under this part.**

**(7) The Department of Health shall assist the division in developing the portion of the test described in Subsection (1)(d).**

**(8) Completing the online tutorial and passing the online test described in this section shall count as 1/2 hour of continuing professional education under Subsection 58-37-6.5(2).**

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Vermont  
18 § 4282  
ADC 12-5-21:4

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:

- (1) “Dispenser” shall mean any person who “dispenses” or engages in “dispensing” as those terms are defined in 26 V.S.A. § 2022(5).
- (2) “Health care provider” shall mean an individual licensed, certified, or authorized by law to provide professional health care service in this state to an individual during that individual's medical or dental care, treatment, or confinement.
- (3) “VPMS” shall mean the Vermont prescription monitoring system established under 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.
- (5) “Department” means the Department of Health.
- (6) “Drug diversion investigator” means an employee of the Department of Public Safety whose primary duties include investigations involving violations of laws regarding prescription drugs or the diversion of prescribed controlled substances, and who has completed a training program established by the Department of Health by rule that is designed to ensure that officers have the training necessary to use responsibly and properly any information that they receive from the VPMS.**
- (7) “Evidence-based” means based on criteria and guidelines that reflect high-quality, cost-effective care. The methodology used to determine such guidelines shall meet recognized standards for systematic evaluation of all available research and shall be free from conflicts of interest. Consideration of the best available scientific evidence does not preclude consideration

of experimental or investigational treatment or services under a clinical investigation approved by an institutional review board.

West's Vermont Administrative Code (2013)  
Title 12. Agency of Human Services  
Subtitle 5. Department of Health  
General  
Rule 21. Prescription Monitoring System  
12-5-21:4. TRAINING

Section 4.1 Designation of Training Programs.

**The Department, in consultation with the Advisory Committee and one or more individuals with medical expertise relating to prescribing controlled substances and treatment of drug addiction and dependence, will periodically designate one or more training programs for law enforcement officers relating to responsible and proper use of VPMS data. The Department will maintain a list of current trained law enforcement officers qualified to receive a report from a professional licensure board as authorized by 18 V.S.A. § 4284(b)(5).**

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## West Virginia

### § 60A-9-5

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.

(2) Subject to the provisions of subdivision (1) of this subsection, the board shall also review the West Virginia Controlled Substance Monitoring Program database and issue reports that identify abnormal or unusual practices of patients who exceed parameters as determined by the advisory committee established in this section. The board shall communicate with prescribers and dispensers to more effectively manage the medications of their patients in the manner recommended by the advisory committee. All other reports produced by the board shall be kept confidential. The board shall maintain the information required by this article for a period of not

less than five years. Notwithstanding any other provisions of this code to the contrary, data obtained under the provisions of this article may be used for compilation of educational, scholarly or statistical purposes, and may be shared with the West Virginia Department of Health and Human Resources for those purposes, as long as the identities of persons or entities and any personally identifiable information, including protected health information, contained therein shall be redacted, scrubbed or otherwise irreversibly destroyed in a manner that will preserve the confidential nature of the information. No individual or entity required to report under section four of this article may be subject to a claim for civil damages or other civil relief for the reporting of information to the Board of Pharmacy as required under and in accordance with the provisions of this article.

(3) The board shall establish an advisory committee to develop, implement and recommend parameters to be used in identifying abnormal or unusual usage patterns of patients in this state. This advisory committee shall:

(A) Consist of the following members: A physician licensed by the West Virginia Board of Medicine, a dentist licensed by the West Virginia Board of Dental Examiners, a physician licensed by the West Virginia Board of Osteopathy, a licensed physician certified by the American Board of Pain Medicine, a licensed physician board certified in medical oncology recommended by the West Virginia State Medical Association, a licensed physician board certified in palliative care recommended by the West Virginia Center on End of Life Care, a pharmacist licensed by the West Virginia Board of Pharmacy, a licensed physician member of the West Virginia Academy of Family Physicians, an expert in drug diversion and such other members as determined by the board.

(B) Recommend parameters to identify abnormal or unusual usage patterns of controlled substances for patients in order to prepare reports as requested in accordance with subsection (a), subdivision (2) of this section.

(C) Make recommendations for training, research and other areas that are determined by the committee to have the potential to reduce inappropriate use of prescription drugs in this state, including, but not limited to, studying issues related to diversion of controlled substances used for the management of opioid addiction.

(D) Monitor the ability of medical services providers, health care facilities, pharmacists and pharmacies to meet the twenty-four hour reporting requirement for the Controlled Substances Monitoring Program set forth in section three of this article, and report on the feasibility of requiring real-time reporting.

(E) Establish outreach programs with local law enforcement to provide education to local law enforcement on the requirements and use of the Controlled Substances Monitoring Program database established in this article.

(b) The Board of Pharmacy shall create a West Virginia Controlled Substances Monitoring Program Database Review Committee of individuals consisting of two prosecuting attorneys

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from West Virginia counties, two physicians with specialties which require extensive use of controlled substances and a pharmacist who is trained in the use and abuse of controlled substances. The review committee may determine that an additional physician who is an expert in the field under investigation be added to the team when the facts of a case indicate that the additional expertise is required. The review committee, working independently, may query the database based on parameters established by the advisory committee. The review committee may make determinations on a case-by-case basis on specific unusual prescribing or dispensing patterns indicated by outliers in the system or abnormal or unusual usage patterns of controlled substances by patients which the review committee has reasonable cause to believe necessitates further action by law enforcement or the licensing board having jurisdiction over the prescribers or dispensers under consideration. The review committee shall also review notices provided by the chief medical examiner pursuant to subsection (h), section ten, article twelve, chapter sixty-one of this code and determine on a case-by-case basis whether a practitioner who prescribed or dispensed a controlled substance resulting in or contributing to the drug overdose may have breached professional or occupational standards or committed a criminal act when prescribing the controlled substance at issue to the decedent. Only in those cases in which there is reasonable cause to believe a breach of professional or occupational standards or a criminal act may have occurred, the review committee shall notify the appropriate professional licensing agency having jurisdiction over the applicable prescriber or dispenser and appropriate law-enforcement agencies and provide pertinent information from the database for their consideration. The number of cases identified shall be determined by the review committee based on a number that can be adequately reviewed by the review committee. The information obtained and developed may not be shared except as provided in this article and is not subject to the provisions of chapter twenty-nine-b of this code or obtainable as discovering in civil matters absent a court order.

(c) The Board of Pharmacy is responsible for establishing and providing administrative support for the advisory committee and the West Virginia Controlled Substances Monitoring Program Database Review Committee. The advisory committee and the review committee shall elect a chair by majority vote. Members of the advisory committee and the review committee may not be compensated in their capacity as members but shall be reimbursed for reasonable expenses incurred in the performance of their duties.

(d) The board shall promulgate rules with advice and consent of the advisory committee, in accordance with the provisions of article three, chapter twenty-nine-a of this code on or before June 1, 2013. The legislative rules must include, but shall not be limited to, the following matters: (1) Identifying parameters used in identifying abnormal or unusual prescribing or dispensing patterns; (2) processing parameters and developing reports of abnormal or unusual prescribing or dispensing patterns for patients, practitioners and dispensers; (3) establishing the information to be contained in reports and the process by which the reports will be generated and disseminated; and (4) setting up processes and procedures to ensure that the privacy, confidentiality, and security of information collected, recorded, transmitted and maintained by the review committee is not disclosed except as provided in this section.

(e) All practitioners, as that term is defined in section one hundred-one, article two of this chapter who prescribe or dispense schedule II, III or IV controlled substances shall, on or before

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July 1, 2011, have online or other form of electronic access to the West Virginia Controlled Substances Monitoring Program database;

(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;

(g) Good faith reliance by a practitioner on information contained in the West Virginia Controlled Substances Monitoring Program database in prescribing or dispensing or refusing or declining to prescribe or dispense a schedule II, III or IV controlled substance shall constitute an absolute defense in any civil or criminal action brought due to prescribing or dispensing or refusing or declining to prescribe or dispense; and

(h) A prescribing or dispensing practitioner may notify law enforcement of a patient who, in the prescribing or dispensing practitioner's judgment, may be in violation of section four hundred ten, article four of this chapter, based on information obtained and reviewed from the controlled substances monitoring database. A prescribing or dispensing practitioner who makes a notification pursuant to this subsection is immune from any civil, administrative or criminal liability that otherwise might be incurred or imposed because of the notification if the notification is made in good faith.

(i) Nothing in the article may be construed to require a practitioner to access the West Virginia Controlled Substances Monitoring Program database except as provided in section five-a of this article.

(j) The Board of Pharmacy shall provide an annual report on the West Virginia Controlled Substance Monitoring Program to the Legislative Oversight Commission on Health and Human Resources Accountability with recommendations for needed legislation no later than January 1 of each year.

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