



TYPES OF AUTHORIZED RECIPIENTS – DE-IDENTIFIED DATA

Research current through June 2014.

This project was supported by Grant No. G1399ONDCP03A, awarded by the Office of National Drug Control Policy. Points of view or opinions in this document are those of the author and do not necessarily represent the official position or policies of the Office of National Drug Control Policy or the United States of Government.

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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 receipt of de-identified data from the database. In most cases, this information is used for research and statistical purposes, but other states limit access to de-identified data to certain state agencies or employees for statistical purposes or trend analysis. The following states either specifically include access to de-identified data or NAMSDDL was informed by the administrator of the state prescription monitoring program that de-identified data is available in that state.

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Arizona
§ 36-2604

Arizona Revised Statutes Annotated (2014)
Title 36. Public Health and Safety
Chapter 28. Controlled Substances Prescription Monitoring Program
Article 1. General Provisions

§ 36-2604. Use and release of confidential information

A. Except as otherwise provided in this section, prescription information submitted to the board pursuant to this article is confidential and is not subject to public inspection. The board shall establish procedures to ensure the privacy and confidentiality of patients and that patient information that is collected, recorded and transmitted pursuant to this article is not disclosed except as prescribed in this section.

B. The board or its designee shall review the prescription information collected pursuant to this article. If the board or its designee has reason to believe an act of unprofessional or illegal conduct has occurred, the board or its designee shall notify the appropriate professional licensing board or law enforcement or criminal justice agency and provide the prescription information required for an investigation.

C. The board may release data collected by the program to the following:

1. A person who is authorized to prescribe or dispense a controlled substance, or a delegate who is authorized by the prescriber or dispenser, to assist that person to provide medical or pharmaceutical care to a patient or to evaluate a patient.

2. An individual who requests the individual's own prescription monitoring information pursuant to § 12-2293.

3. A professional licensing board established pursuant to title 32, chapter 7, 11, 13, 14, 15, 16, 17, 18, 21, 25 or 29. Except as required pursuant to subsection B of this section, the board shall provide this information only if the requesting board states in writing that the information is necessary for an open investigation or complaint.

4. A local, state or federal law enforcement or criminal justice agency. Except as required pursuant to subsection B of this section, the board shall provide this information only if the requesting agency states in writing that the information is necessary for an open investigation or complaint.

5. The Arizona health care cost containment system administration regarding persons who are receiving services pursuant to chapter 29 of this title. [FN2] Except as required pursuant to

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subsection B of this section, the board shall provide this information only if the administration states in writing that the information is necessary for an open investigation or complaint.

6. A person who is serving a lawful order of a court of competent jurisdiction.

7. A person who is authorized to prescribe or dispense a controlled substance and who performs an evaluation on an individual pursuant to § 23-1026.

D. The board may provide data to public or private entities for statistical, research or educational purposes after removing information that could be used to identify individual patients or persons who received prescriptions from dispensers.

E. For the purposes of this section, “delegate” means a licensed health care professional who is employed in the office of or in a hospital with the prescriber or dispenser or an unlicensed medical records technician, medical assistant or office manager who is employed in the office of or in a hospital with the prescriber and who has received training regarding both the Health Insurance Portability and Accountability Act privacy standards, 45 Code of Federal Regulations Part 164, Subpart E, and security standards, 45 Code of Federal Regulations, Part 164, Subpart C.

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Arkansas
§ 20-7-607

West's Arkansas Code Annotated (2014)
Title 20. Public Health and Welfare
Subtitle 2. Health and Safety (Chapters 6 to 44)
Chapter 7. State Board of Health--Department of Health
Subchapter 6. Prescription Drug Monitoring Program Act

§ 20-7-607. Providing prescription monitoring information

(a)(1) The Department of Health may review the Prescription Drug Monitoring Program information, including without limitation a review to identify information that appears to indicate whether a person may be obtaining prescriptions in a manner that may represent misuse or abuse of controlled substances.

(2) If information of misuse or abuse is identified, the department shall notify the practitioners and dispensers who prescribed or dispensed the prescriptions.

(b) The department shall provide information in the Prescription Drug Monitoring Program upon request and at no cost only to the following persons:

(1) A person authorized to prescribe or dispense controlled substances for the purpose of providing medical or pharmaceutical care for his or her patients or for reviewing information regarding prescriptions that are recorded as having been issued or dispensed by the requester;

(2) A patient who requests his or her own prescription monitoring information;

(3) A parent or legal guardian of a minor child who requests the minor child's Prescription Drug Monitoring Program information;

(4)(A) A designated representative of a professional licensing board of the professions of the healing arts representing health care disciplines whose licensees are prescribers pursuant to an investigation of a specific individual, entity, or business licensed or permitted by that board.

(B) Except as permitted by subsection (a)(2) of this section, the department shall provide information under subsection (b)(4)(A) of this section only if the requesting board states in writing that the information is necessary for an investigation;

(5) The State Medical Examiner as authorized by law to investigate causes of deaths for cases under investigation pursuant to his or her official duties and responsibilities;

(6) Local, state, and federal law enforcement or prosecutorial officials engaged in the administration, investigation, or enforcement of the laws governing controlled substances required to be submitted under this subchapter pursuant to the agency's official duties and responsibilities; and

(7) Personnel of the department for purposes of administration and enforcement of this subchapter.

(c) Information collected under this subchapter shall be maintained for three (3) years.

(d) The department may provide information to public or private entities for statistical, research, or educational purposes after encrypting or removing the patient's name, street name and number, patient identification number, month and day of birth, and prescriber information that could be used to identify individual patients or persons who received prescriptions from dispensers, or both.

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California
Health and Safety Code § 11165

West's Annotated California Codes (2014)
Health and Safety Code
Division 10. Uniform Controlled Substances Act
Chapter 4. Prescriptions
Article 1. Requirements of Prescriptions

§ 11165. Controlled Substance Utilization Review and Evaluation System (CURES); electronic monitoring of Schedule II, Schedule III, and Schedule IV controlled substances; funding; confidentiality; reporting requirements for dispensers; stakeholder assistance in establishing rules and regulations and identifying CURES upgrades; education on access and use of CURES PDMP

(a) To assist health care practitioners in their efforts to ensure appropriate prescribing, ordering, administering, furnishing, and dispensing of controlled substances, law enforcement and regulatory agencies in their efforts to control the diversion and resultant abuse of Schedule II, Schedule III, and Schedule IV controlled substances, and for statistical analysis, education, and research, the Department of Justice shall, contingent upon the availability of adequate funds in the CURES Fund, maintain the Controlled Substance Utilization Review and Evaluation System (CURES) for the electronic monitoring of, and Internet access to information regarding, the prescribing and dispensing of Schedule II, Schedule III, and Schedule IV controlled substances by all practitioners authorized to prescribe, order, administer, furnish, or dispense these controlled substances.

(b) The Department of Justice may seek and use grant funds to pay the costs incurred by the operation and maintenance of CURES. The department shall annually report to the Legislature and make available to the public the amount and source of funds it receives for support of CURES.

(c)(1) The operation of CURES shall comply with all applicable federal and state privacy and security laws and regulations.

(2) CURES shall operate under existing provisions of law to safeguard the privacy and confidentiality of patients. Data obtained from CURES shall only be provided to appropriate state, local, and federal public agencies for disciplinary, civil, or criminal purposes and to other agencies or entities, as determined by the Department of Justice, for the purpose of educating practitioners and others in lieu of disciplinary, civil, or criminal actions. **Data may be provided to public or private entities, as approved by the Department of Justice, for educational, peer review, statistical, or research purposes, provided that patient information, including any information that may identify the patient, is not compromised. Further, data disclosed to any individual or agency as described in this subdivision shall not be disclosed, sold, or**

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transferred to any third party. The Department of Justice shall establish policies, procedures, and regulations regarding the use, access, evaluation, management, implementation, operation, storage, disclosure, and security of the information within CURES, consistent with this subdivision.

(d) For each prescription for a Schedule II, Schedule III, or Schedule IV controlled substance, as defined in the controlled substances schedules in federal law and regulations, specifically Sections 1308.12, 1308.13, and 1308.14, respectively, of Title 21 of the Code of Federal Regulations, the dispensing pharmacy, clinic, or other dispenser shall report the following information to the Department of Justice as soon as reasonably possible, but not more than seven days after the date a controlled substance is dispensed, in a format specified by the Department of Justice:

(1) Full name, address, and, if available, telephone number of the ultimate user or research subject, or contact information as determined by the Secretary of the United States Department of Health and Human Services, and the gender, and date of birth of the ultimate user.

(2) The prescriber's category of licensure, license number, national provider identifier (NPI) number, if applicable, the federal controlled substance registration number, and the state medical license number of any prescriber using the federal controlled substance registration number of a government-exempt facility.

(3) Pharmacy prescription number, license number, NPI number, and federal controlled substance registration number.

(4) National Drug Code (NDC) number of the controlled substance dispensed.

(5) Quantity of the controlled substance dispensed.

(6) International Statistical Classification of Diseases, 9th revision (ICD-9) or 10th revision (ICD-10) Code, if available.

(7) Number of refills ordered.

(8) Whether the drug was dispensed as a refill of a prescription or as a first-time request.

(9) Date of origin of the prescription.

(10) Date of dispensing of the prescription.

(e) The Department of Justice may invite stakeholders to assist, advise, and make recommendations on the establishment of rules and regulations necessary to ensure the proper administration and enforcement of the CURES database. All prescriber and dispenser invitees shall be licensed by one of the boards or committees identified in subdivision (d) of Section 208

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of the Business and Professions Code, in active practice in California, and a regular user of CURES.

(f) The Department of Justice shall, prior to upgrading CURES, consult with prescribers licensed by one of the boards or committees identified in subdivision (d) of Section 208 of the Business and Professions Code, one or more of the boards or committees identified in subdivision (d) of Section 208 of the Business and Professions Code, and any other stakeholder identified by the department, for the purpose of identifying desirable capabilities and upgrades to the CURES Prescription Drug Monitoring Program (PDMP).

(g) The Department of Justice may establish a process to educate authorized subscribers of the CURES PDMP on how to access and use the CURES PDMP.

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Colorado
§ 12-42.5-404

West's Colorado Revised Statutes Annotated (2014)
Title 12. Professions and Occupations
Health Care
Article 42.5. Pharmacists, Pharmacy Businesses, and Pharmaceuticals
Part 4. Electronic Monitoring of Prescription Drugs

§ 12-42.5-404. Program operation--access--rules

- (1) The board shall operate and maintain the program.
- (2) The board shall adopt all rules necessary to implement the program.
- (3) The program is available for query only to the following persons or groups of persons:
 - (a) Board staff responsible for administering the program;
 - (b) Any practitioner with the statutory authority to prescribe controlled substances, or an individual designated by the practitioner to act on his or her behalf in accordance with Section 12-42.5-403(1.5)(b) to the extent the query relates to a current patient of the practitioner to whom the practitioner is prescribing or considering prescribing any controlled substance;
 - (c) A practitioner, or an individual designated by the practitioner to act on his or her behalf in accordance with Section 12-42.5-403(1.5)(b) engaged in a legitimate program to monitor a patient's drug abuse;
 - (c.5) The medical director, or his or her designee, at a facility that treats addiction with controlled substances, if an individual in treatment at the facility gives permission to the facility to access his or her program records;
 - (d) A pharmacist, an individual designated by the pharmacist in accordance with Section 12-42.5-403(1.5)(b) to act on his or her behalf, or a pharmacist licensed in another state, to the extent the information requested relates specifically to a current patient to whom the pharmacist is dispensing or considering dispensing a controlled substance or to whom the pharmacist is providing clinical patient care services;
 - (e) Law enforcement officials so long as the information released is specific to an individual patient or practitioner and is part of a bona fide investigation, and the request for information is accompanied by an official court order or subpoena;

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(f) The individual who is the recipient of a controlled substance prescription so long as the information released is specific to the individual;

(g) State regulatory boards within the division and the director of the division so long as the information released is specific to an individual practitioner and is part of a bona fide investigation, and the request for information is accompanied by an official court order or subpoena;

(h) A resident physician with an active physician training license issued by the Colorado medical board pursuant to section 12-36-122 and under the supervision of a licensed physician;

(i) The Department of Public Health and Environment for purposes of population-level analysis, but any use of the program data by the department is subject to the federal “Health Insurance Portability and Accountability Act of 1996”, Pub.L. 104-191, as amended, and implementing federal regulations, including the requirement to remove any identifying data unless exempted from the requirement.

(4) The board shall not charge a practitioner or pharmacy who transmits data in compliance with the operation and maintenance of the program a fee for the transmission of the data.

(5) The board, the Department of Public Health and Environment, or the Department of Health Care Policy and Financing, pursuant to a written agreement that ensures compliance with this part 4, may provide data to qualified personnel of a public or private entity for the purpose of bona fide research or education so long as the data does not identify a recipient of a practitioner who prescribed, or a prescription drug outlet that dispensed, a prescription drug.

(6) The board shall provide a means of sharing information about individuals whose information is recorded in the program with out-of-state health care practitioners and law enforcement officials that meet the requirements of paragraph (b), (c), or (e) of subsection (3) of this section.

(7) The board shall develop criteria for indicators of misuse, abuse, and diversion of controlled substances and, based on those criteria, provide unsolicited reports of dispensed controlled substances to prescribing practitioners and dispensing pharmacies for purposes of education and intervention to prevent and reduce occurrences of controlled substance misuse, abuse, and diversion. In developing the criteria, the board shall consult with the Colorado Dental Board, Colorado Medical Board, State Board of Nursing, State Board of Optometry, Colorado Podiatry Board, and State Board of Veterinary Medicine.

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Connecticut
ADC 21a-254-6

Regulations of Connecticut State Agencies (2014)
Title 21A. Consumer Protection
Department of Consumer Protection
Electronic Prescription Drug Monitoring Program

Sec. 21a-254-6. Management of information

The department may provide prescription information obtained from pharmacies to:

- (a) Other regulatory, investigative or law enforcement agencies for disciplinary, civil, or criminal purposes;
- (b) Practitioners, for the purpose of education in lieu of disciplinary, civil or criminal action;
- (c) Practitioners and pharmacists, for the purposes of patient care, drug therapy management and monitoring of controlled substances obtained by the patient; and
- (d) Public or private entities, for statistical, research, or educational purposes, provided that the privacy of patients and confidentiality of patient information is not compromised.**

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Delaware
16 § 4798

West's Delaware Code Annotated (2014)
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 upon the availability of appropriations, or of other adequate funding to implement and maintain the Prescription Monitoring Program and upon 3-1-2014. See Historical and Statutory Notes below.>

...

(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;

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- 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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District of Columbia
§ 48-853.05

West's District of Columbia Code Annotated 2001 Edition (2014)
Division VIII. General Laws.
Title 48. Foods and Drugs.
Subtitle II. Prescription Drugs.
Chapter 8G. Prescription Drug Monitoring Program.

§ 48-853.05. Confidentiality of data; disclosure of information; discretionary authority of the Director.

(a) All data, records, and reports relating to the prescribing and dispensing of covered substances to patients and any abstracts from such data, records, and reports that are in the possession of the Program pursuant to this chapter and any materials relating to the operation or safety of the Program shall be confidential and shall be exempt from disclosure based on requests made pursuant to subchapter II of Chapter 2 of Title 5. Information obtained pursuant to the Program may only be disclosed as provided in this chapter.

(b) Upon receiving a request for information in accordance with the Department's regulations and in compliance with applicable District and federal laws and regulations, the Director shall disclose information relevant to:

(1) A specific investigation of a specific patient or of a specific dispenser or prescriber to an agent designated by the Chief of the Metropolitan Police Department to conduct drug diversion investigations;

(2) An investigation or inspection of or allegation of misconduct by a specific person licensed, certified, or registered by or an applicant for licensure, certification, or registration by a health occupations board or the Department;

(3) A disciplinary proceeding before a health occupations board or in any subsequent hearing, trial, or appeal of an action or board order to designated employees of the Department;

(4) The proceedings of any grand jury or additional grand jury that has been properly impaneled in accordance with § 11-1916; and

(5) A specific investigation of a specific dispenser or specific prescriber to an agent of the United States Drug Enforcement Administration with authority to conduct drug diversion investigations.

(c)(1) In accordance with the Department's regulations and applicable federal law and regulations, the Director may, at the Director's discretion, disclose:

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(A) Information in the possession of the Program concerning a patient who is over the age of 18 years to that patient, or to the parent or legal guardian of a child aged 18 years or under, unless otherwise prohibited by District or federal law;

(B) Information on a specific patient to a prescriber for the purpose of establishing the treatment history of the specific patient when the patient is either under care and treatment by the prescriber or the prescriber is initiating treatment of the patient;

(C) Information on a specific patient to a dispenser for the purpose of establishing a prescription history to assist the dispenser in determining the validity of a prescription when the patient is seeking a covered substance from the dispenser or the facility in which the dispenser practices;

(D) Information relevant to an investigation or regulatory proceeding of a specific dispenser or prescriber to other regulatory authorities concerned with granting, limiting, or denying licenses, certificates, or registrations to practice a health profession when the regulatory authority licenses the dispenser or prescriber, or the dispenser or prescriber is seeking licensure by a regulatory authority;

(E) Information relevant to an investigation relating to a specific dispenser or prescriber who is a participating provider in the District Medicaid program, DC Health Care Alliance, or any other public health care program; information relating to an investigation relating to a specific patient who is currently eligible for and receiving, or who has been eligible for and has received medical assistance services; information relevant to the Medicaid Fraud Control Unit of the Office of the Inspector General, or to designated employees of the Department of Health Care Finance, as appropriate;

(F) Information relevant to the determination of the cause of death of a specific patient to the designated employees of the Office of the Chief Medical Examiner; and

(G) Information for the purpose of bona fide research or education to qualified personnel; provided, that:

(i) Data elements that would reasonably identify a specific patient, prescriber, or dispenser shall be deleted or redacted from the information before disclosure; and

(ii) Release of the information shall only be made pursuant to a written agreement between qualified personnel and the Director to ensure compliance with this chapter.

(2) For the purposes of a disclosure under paragraph (1)(B) or (C) of this subsection:

(A) The request shall be made and the information shall be provided in the manner specified by the Director through rulemaking; and

(B) Notice shall be given to patients that the information described in paragraph (1)(B) or (C) of this subsection, as applicable, may be requested by a prescriber or dispenser participating with the Program.

(d) Confidential information that has been received, maintained, or developed by a health occupations board or disclosed by the health occupations board pursuant to this chapter shall not be available for discovery or court subpoena or introduced into evidence in any medical malpractice suit or other action for damages arising out of the provision of or failure to provide services; provided, that this section shall not be construed to inhibit any investigation or prosecution conducted pursuant to this chapter.

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Georgia
§ 16-13-60

West's Code of Georgia Annotated (2014)
Title 16. Crimes and Offenses
Chapter 13. Controlled Substances
Article 2. Regulation of Controlled Substances
Part 2. Controlled Substances Prescription Monitoring

§ 16-13-60. Confidentiality of information submitted

(a) Except as otherwise provided in subsections (c) and (d) of this Code section, prescription information submitted pursuant to Code Section 16-13-59 shall be confidential and shall not be subject to open records requirements, as contained in Article 4 of Chapter 18 of Title 50.

(b) The agency, in conjunction with the board, shall establish and maintain strict procedures to ensure that the privacy and confidentiality of patients, prescribers, and patient and prescriber information collected, recorded, transmitted, and maintained pursuant to this part are protected. Such information shall not be disclosed to any person or entity except as specifically provided in this part and only in a manner which in no way conflicts with the requirements of the federal Health Insurance Portability and Accountability Act (HIPAA) of 1996, P.L. 104-191.

(c) The agency shall be authorized to provide requested prescription information collected pursuant to this part only as follows:

(1) To persons authorized to prescribe or dispense controlled substances for the sole purpose of providing medical or pharmaceutical care to a specific patient;

(2) Upon the request of a patient, prescriber, or dispenser about whom the prescription information requested concerns or upon the request on his or her behalf of his or her attorney;

(3) To local, state, or federal law enforcement or prosecutorial officials pursuant to the issuance of a search warrant pursuant to Article 2 of Chapter 5 of Title 17; and

(4) To the agency or the Georgia Composite Medical Board upon the issuance of an administrative subpoena issued by a Georgia state administrative law judge.

(d) The board may provide data to government entities for statistical, research, educational, or grant application purposes after removing information that could be used to identify prescribers or individual patients or persons who received prescriptions from dispensers.

(e) Any person or entity who receives electronic data base prescription information or related reports relating to this part from the agency shall not provide such information or reports to any other person or entity except by order of a court of competent jurisdiction pursuant to this part.

(f) Any permissible user identified in this part who directly accesses electronic data base prescription information shall implement and maintain a comprehensive information security program that contains administrative, technical, and physical safeguards that are substantially equivalent to the security measures of the agency. 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 shall assess the sufficiency of any safeguards in place to control the risks.

(g) No provision in this part shall be construed to modify, limit, diminish, or impliedly repeal any authority existing on June 30, 2011, of a licensing or regulatory board or any other entity so authorized to obtain prescription information from sources other than the data base maintained pursuant to this part; provided, however, that the agency shall be authorized to release information from the data base only in accordance with the provisions of this part.

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Idaho
§ 37-2730A

West's Idaho Code Annotated (2014)
Title 37. Food, Drugs, and Oil
Chapter 27. Uniform Controlled Substances
Article III

§ 37-2730A. Prescription tracking program

(1) The board shall maintain a program to track the prescriptions for controlled substances that are filed with the board under section 37-2726, Idaho Code, for the purpose of assisting in identifying illegal activity related to the dispensing of controlled substances and for the purpose of assisting the board in providing information to patients, practitioners and pharmacists to assist in avoiding inappropriate use of controlled substances. The tracking program and any data created thereby shall be administered by the board.

(2) The board shall use the information obtained through the tracking program in identifying activity it reasonably suspects may be in violation of this chapter or medical assistance law. The board shall report this information to the individuals and persons set forth in section 37-2726(2), Idaho Code. The board may release unsolicited information to pharmacists and practitioners when the release of information may be of assistance in preventing or avoiding inappropriate use of controlled substances. The board may provide the appropriate law enforcement agency, medicaid or medicare agency or licensing board with the relevant information in the board's possession, including information obtained from the tracking program, for further investigation, or other appropriate law enforcement or administrative enforcement use.

(3) Information, which does not identify individual patients, practitioners or dispensing pharmacists or pharmacies, may be released by the board for educational, research or public information purposes.

(4) Nothing herein shall prevent a pharmacist or practitioner from furnishing another pharmacist or practitioner information obtained pursuant to and in compliance with this chapter.

(5) Unless there is shown malice or criminal intent or gross negligence or reckless, willful and wanton conduct as defined in section 6-904C, Idaho Code, the state of Idaho, the board, any other state agency, or any person, or entity in proper possession of information as herein provided shall not be subject to any liability or action for money damages or other legal or equitable relief by reason of any of the following:

(a) The furnishing of information under the conditions herein provided;

(b) The receiving and use of, or reliance on, such information;

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- (c) The fact that any such information was not furnished; or
- (d) The fact that such information was factually incorrect or was released by the board to the wrong person or entity.
- (6) The board 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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Illinois
720 § 570/318

West's Smith-Hurd Illinois Compiled Statutes Annotated (2014)
Chapter 720. Criminal Offenses
Offenses Against the Public
Act 570. Illinois Controlled Substances Act
Article III. Registration and Control of Manufacture, Distribution and Dispensing
570/318. Confidentiality of information

§ 318. Confidentiality of information.

...

(i) The Department may compile statistical reports from the information described in subsection (a). The reports must not include information that identifies, by name, license or address, any practitioner, dispenser, ultimate user, or other person administering a controlled substance.

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Indiana

§ 35-48-7-11.1

West's Annotated Indiana Code (2014)

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

Sec. 11.1. (a) Information received by the INSPECT program under section 8.1 of this chapter is confidential.

...

(j) The board may compile statistical reports from the information described in subsection (a). The reports must not include information that identifies any practitioner, ultimate user, or other person administering a controlled substance. Statistical reports compiled under this subsection are public records.

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Kansas
§ 65-1685

West's Kansas Statutes Annotated (2014)
Chapter 65. Public Health
Article 16. Regulation of Pharmacists

§ 65-1685. Same; database information privileged and confidential; persons authorized to receive data; advisory committee review of information

(a) The prescription monitoring program database, all information contained therein and any records maintained by the board, or by any entity contracting with the board, submitted to, maintained or stored as a part of the database, shall be privileged and confidential, shall not be subject to subpoena or discovery in civil proceedings and may only be used for investigatory or evidentiary purposes related to violations of state or federal law and regulatory activities of entities charged with administrative oversight of those persons engaged in the prescribing or dispensing of scheduled substances and drugs of concern, shall not be a public record and shall not be subject to the Kansas open records act, K.S.A. 45-215 et seq., and amendments thereto, except as provided in subsections (c) and (d).

(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 except as provided in subsections (c) and (d).

(c) The board is hereby authorized to provide data in the prescription monitoring program to the following persons:

(1) Persons authorized to prescribe or dispense scheduled substances and drugs of concern, for the purpose of providing medical or pharmaceutical care for their patients;

(2) an individual who requests the individual's own prescription monitoring information in accordance with procedures established by the board;

(3) designated representatives from the professional licensing, certification or regulatory agencies charged with administrative oversight of those persons engaged in the prescribing or dispensing of scheduled substances and drugs of concern;

(4) local, state and federal law enforcement or prosecutorial officials engaged in the administration, investigation or enforcement of the laws governing scheduled substances and drugs of concern subject to the requirements in K.S.A. 22-2502, and amendments thereto;

(5) designated representatives from the department of health and environment regarding authorized medicaid program recipients;

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(6) persons authorized by a grand jury subpoena, inquisition subpoena or court order in a criminal action;

(7) personnel of the prescription monitoring program advisory committee for the purpose of operation of the program;

(8) personnel of the board for purposes of administration and enforcement of this act or the uniform controlled substances act, K.S.A. 65-4101 et seq., and amendments thereto;

(9) persons authorized to prescribe or dispense scheduled substances and drugs of concern, when an individual is obtaining prescriptions in a manner that appears to be misuse, abuse or diversion of scheduled substances or drugs of concern; and

(10) medical examiners, coroners or other persons authorized under law to investigate or determine causes of death.

(d) The prescription monitoring program advisory committee established pursuant to K.S.A. 65-1689, and amendments thereto, is authorized to review and analyze the data for purposes of identifying patterns and activity of concern.

(1) If a review of information appears to indicate a person may be obtaining prescriptions in a manner that may represent misuse or abuse of controlled substances and drugs of concern, the advisory committee is authorized to notify the prescribers and dispensers who prescribed or dispensed the prescriptions. If the review identifies patterns or other evidence sufficient to create a reasonable suspicion of criminal activity, the advisory committee is authorized to notify the appropriate law enforcement agency.

(2) If a review of information appears to indicate that a violation of state or federal law relating to prescribing controlled substances and drugs of concern may have occurred, or that a prescriber or dispenser has knowingly prescribed, dispensed or obtained controlled substances and drugs of concern in a manner that is inconsistent with recognized standards of care for the profession, the advisory committee shall determine whether a report to the professional licensing, certification or regulatory agencies charged with administrative oversight of those persons engaged in prescribing or dispensing of controlled substances and drugs of concern or to the appropriate law enforcement agency is warranted.

(A) For purposes of such determination the advisory committee may, in consultation with the appropriate regulatory agencies and professional organizations, establish criteria regarding appropriate standards and utilize volunteer peer review committees of professionals with expertise in the particular practice to create such standards and review individual cases.

(B) The peer review committee or committees appointed herein shall have authority to request and receive information in the prescription monitoring program database from the director of the prescription monitoring program.

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(C) If the determination is made that a referral to a regulatory or law enforcement agency is not warranted but educational or professional advising might be appropriate, the advisory committee may refer the prescribers or dispensers to other such resources.

(e) The board is hereby authorized to provide data in the prescription monitoring program to public or private entities for statistical, research or educational purposes after removing information that could be used to identify individual practitioners, dispensers, patients or persons who received prescriptions from dispensers.

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Kentucky
§ 218A.240

Baldwin's Kentucky Revised Statutes Annotated (2014)
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

...

(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.

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Louisiana
§ 40:1007

West's Louisiana Statutes Annotated (2014)
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.

...

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.

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Maine
22 § 7250

Maine Revised Statutes Annotated (2014)
Title 22. Health and Welfare
Subtitle 4. Human Services
Part 3. Drug Abuse
Chapter 1603. Controlled Substances Prescription Monitoring

§ 7250. Access to prescription monitoring information and confidentiality

...

6. Treatment pattern data. The department may provide to a prescriber who treats a member under the MaineCare program prescription monitoring information on the prescriber and other prescribers that is de-identified as to prescriber and patient and that indicates treatment patterns in comparison among peers. If the department has shared with a prescriber treatment pattern data under this subsection, the department shall allow the prescriber time to align the prescriber's prescribing patterns with the patterns of the peers of the prescriber. The department may take appropriate actions with regard to a prescriber who is unable to achieve treatment pattern alignment as provided in this subsection.

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Maryland

Health-General § 21-2A-06 (eff. until Oct. 1, 2014)

Health-General § 21-2A-06 (eff. Oct. 1, 2014)

West's Annotated Code of Maryland (2014)

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, 2014>

...

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.

...

West's Annotated Code of Maryland (2014)

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, 2014>

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

(f)(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.

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Massachusetts
94C § 24A

Massachusetts General Laws Annotated (2014)
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

...

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

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Mississippi
§ 73-21-127

West's Annotated Mississippi Code (2014)
Title 73. Professions and Vocations
Chapter 21. Pharmacists
Mississippi Pharmacy Practice Act

§ 73-21-127. Computer program to track prescriptions for controlled substances and report illegal activity

The Board of Pharmacy shall develop and implement a computerized program to track prescriptions for controlled substances and to report suspected abuse and misuse of controlled substances in compliance with the federal regulations promulgated under authority of the National All Schedules Prescription Electronic Reporting Act of 2005 and in compliance with the federal HIPAA law, under the following conditions:

...

(e)(i) Access to collected data shall be confidential and not subject to the provisions of the federal Freedom of Information Act or the Mississippi Open Records Act. Upon request, the State Board of Pharmacy shall provide collected information to: pharmacists or practitioners who are properly registered with the State Board of Pharmacy and are authorized to prescribe or dispense controlled substances for the purpose of providing medical and pharmaceutical care for their patients; local, state and federal law enforcement officials engaged in the administration, investigation or enforcement of the laws governing illicit drug use; regulatory and licensing boards in this state; Division of Medicaid regarding Medicaid and Medicare Program recipients; judicial authorities under grand jury subpoena; an individual who requests the individual's own prescription monitoring information; and prescription monitoring programs in other states through mutual agreement adhering to State Board of Pharmacy policies.

(ii) The Director of the Mississippi Bureau of Narcotics, or his designee, shall have access to the Prescription Monitoring Program (PMP) database for the purpose of investigating the potential illegal acquisition, distribution, dispensing, prescribing or administering of the controlled and noncontrolled substances monitored by the program, subject to all legal restrictions on further dissemination of the information obtained.

(iii) The State Board of Pharmacy may also provide generic, nonidentifying statistical data for research or educational purposes.

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Montana
ADC 24.174.1713

Administrative Rules of Montana (2013)
Title 24. Labor and Industry
Chapter 174. Board of Pharmacy
Sub-chapter 17. Prescription Drug Registry

24.174.1713 RELEASE OF PRESCRIPTION DRUG REGISTRY INFORMATION TO
OTHER ENTITIES

(1) The board shall provide prescription registry information to public or private entities for public research, policy, or educational purposes, but only after removing information that identifies or could reasonably be used to identify individuals or entities whose information is contained in the registry.

(2) The board may charge a fee to a person who requests information under this rule.

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Nevada
§ 453.1545

West's Nevada Revised Statutes Annotated (2014)
Title 40. Public Health and Safety (Chapters 439-461A)
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.

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New Jersey
§ 45:1-46

New Jersey Statutes Annotated (2014)
Title 45. Professions and Occupations
Subtitle 1. Professions and Occupations Regulated by State Boards of Registration and Examination
Chapter 1. General Provisions
Article 3. Record Background Checks for Health Care Professionals

§ 45:1-46. Access to prescription information

...

g. The director may provide nonidentifying prescription drug monitoring information to public or private entities for statistical, research or educational purposes.

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New Mexico
ADC 16.19.29

Code of New Mexico Rules (2014)
Title 16. Occupational and Professional Licensing
Chapter 19. Pharmacists
Part 29. Controlled Substance Prescription Monitoring Program

...

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.

...

E. The board shall be authorized to provide data in the prescription monitoring program to the following persons:

- (1) persons authorized to prescribe or dispense controlled substances, for the purpose of providing medical or pharmaceutical care for their patients;
- (2) an individual who request's their own prescription monitoring information in accordance with procedures established under 61-11-2.D NMSA, 1978 and Subsection G of 16.19.6.23 NMAC;
- (3) New Mexico medical board, New Mexico board of nursing, New Mexico board of veterinary medicine, New Mexico board of dental health care, board of examiners in optometry, osteopathic examiners board, acupuncture & oriental medicine board, and podiatry board for their licensees;
- (4) professional licensing authorities of other states if their licensees practice in the state or prescriptions provided by their licensees are dispensed in the state;
- (5) local, state and federal law enforcement or prosecutorial officials engaged in an ongoing investigation of an individual in the enforcement of the laws governing licit drugs;
- (6) human services department regarding medicaid program recipients;
- (7) metropolitan, district, state or federal court(s) under grand jury subpoena or criminal court order;
- (8) personnel of the board for purposes of administration and enforcement of this regulation, or 16.19.20 NMAC or;

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(9) the controlled substance monitoring program of another state or group of states with whom the state has established an interoperability agreement;

(10) a parent to have access to the prescription records about his or her minor child, as his or her minor child's personal representative when such access is not inconsistent with state or other laws;

(11) the board shall use de-identified data obtained from the prescription drug monitoring database to identify and report to state and local public health authorities the geographic areas of the state where anomalous prescribing dispensing or use of controlled substances is occurring.

(12) the board shall share prescription drug monitoring database data with the department of health for the purpose of tracking inappropriate prescribing and misuse of controlled substances, including drug overdose.

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North Carolina
§ 90-113.74

West's North Carolina General Statutes Annotated (2014)
Chapter 90. Medicine and Allied Occupations
Article 5E. North Carolina Controlled Substances Reporting System Act

§ 90-113.74. Confidentiality

...

(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.

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North Dakota
§ 19-03.5-03

West's North Dakota Century Code Annotated (2014)
Title 19. Foods, Drugs, Oils, and Compounds
Chapter 19-03.5. Prescription Drug Monitoring Program

§ 19-03.5-03. Access to prescription information

...

3. Unless disclosure is prohibited by law, the board may provide data in the central repository to:

- a. A prescriber for the purpose of providing medical care to a patient, a dispenser for the purpose of filling a prescription or providing pharmaceutical care for a patient, a prescriber or dispenser inquiring about the prescriber's or dispenser's own prescribing activity, or a prescriber or dispenser in order to further the purposes of the program;
- b. An individual who requests the prescription information of the individual or the individual's minor child;
- c. State boards and regulatory agencies that are responsible for the licensing of individuals authorized to prescribe or dispense controlled substances if the board or regulatory agency is seeking information from the central repository that is relevant to an investigation of an individual who holds a license issued by that board or regulatory agency;
- d. Local, state, and federal law enforcement or prosecutorial officials engaged in the enforcement of laws relating to controlled substances who seek information for the purpose of an investigation or prosecution of the drug-related activity or probation compliance of an individual;
- e. The department of human services for purposes regarding the utilization of controlled substances by a medicaid recipient or establishment and enforcement of child support and medical support;
- f. Workforce safety and insurance for purposes regarding the utilization of controlled substances by a claimant;
- g. Judicial authorities under grand jury subpoena or court order or equivalent judicial process for investigation of criminal violations of controlled substances laws;

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h. Public or private entities for statistical, research, or educational purposes after the information is de-identified with respect to any prescriber, dispenser, or patient who received a prescription for a controlled substance;

i. A peer review committee which means any committee of a health care organization, composed of health care providers, employees, administrators, consultants, agents, or members of the health care organization's governing body, which conducts professional peer review as defined in chapter 23-34; or

j. A licensed addiction counselor for the purpose of providing services for a licensed treatment program in this state.

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Ohio

§ 4729.80 (eff. until Sept. 17, 2014)

§ 4729.80 (eff. Sept. 17, 2014)

Baldwin's Ohio Revised Code Annotated (2014)

Title XLVII. Occupations--Professions

Chapter 4729. Pharmacists; Dangerous Drugs

Miscellaneous Provisions

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

<Text of Section Effective Until September 17, 2014>

...

(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.**

...

Baldwin's Ohio Revised Code Annotated (2014)

Title XLVII. Occupations--Professions

Chapter 4729. Pharmacists; Dangerous Drugs

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Oklahoma

63 § 2-309D (eff. until Nov. 1, 2014)

63 § 2-309D (eff. Nov. 1, 2014)

Oklahoma Statutes Annotated (2014)

Title 63. Public Health and Safety

Chapter 2. Uniform Controlled Dangerous Substances Act

Article III. Regulation of Manufacture, Distribution, Dispensing, Prescribing, Administering and Using for Scientific Purposes of Controlled Dangerous Substances

Anti-Drug Diversion Act

§ 2-309D. Central repository information--Confidentiality--Access-- Disclosure--Penalties--Liability

<Text of Section Effective Until November 1, 2014>

...

C. This section shall not prevent the disclosure, at the discretion of the Director of the Oklahoma State Bureau of Narcotics and Dangerous Drugs Control, of statistical information gathered from the central repository to the general public which shall be limited to types and quantities of controlled substances dispensed and the county where dispensed.

...

Oklahoma Statutes Annotated (2014)

Title 63. Public Health and Safety

Chapter 2. Uniform Controlled Dangerous Substances Act

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Oregon
§ 431.966

West's Oregon Revised Statutes Annotated (2014)
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 subject to final change by the Oregon Office of the Legislative Counsel.>

...

(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.

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(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.

...

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Rhode Island
§ 21-28-3.32
ADC 31-2-1:3.0

West's General Laws of Rhode Island Annotated (2014)
Title 21. Food and Drugs
Chapter 28. Uniform Controlled Substances Act
Article III. Regulation of Manufacturing, Distributing, Prescribing, Administering, and
Dispensing Controlled Substances

§ 21-28-3.32. Electronic prescription database

(a) The information contained in any prescription drug monitoring database maintained by the department of health pursuant to section 3.18 of this chapter shall be disclosed only:

(1) To a practitioner who certifies that the requested information is for the purpose of evaluating the need for or providing medical treatment for a current patient to whom the practitioner is prescribing or considering prescribing a controlled substance;

(2) To a pharmacist who certifies that the requested information is for a current client to whom the pharmacist is dispensing or considering dispensing a controlled substance;

(3) To an authorized designee of the practitioner and/or pharmacist to consult the prescription drug monitoring database on the practitioner's and/or pharmacist's behalf, provided that:

(i) The designee so authorized is employed by the same professional practice or pharmacy;

(ii) The practitioner or pharmacist takes reasonable steps to ensure that such designee is sufficiently competent in the use of the database;

(iii) The practitioner or pharmacist remains responsible for ensuring that access to the database by the designee is limited to authorized purposes as provided for in subsections (a)(1) and (a)(2) of this section;

(iv) The practitioner or pharmacist remains responsible for ensuring access to the database by the designee occurs in a manner that protects the confidentiality of information obtained from the database, and remains responsible for any breach of confidentiality;

(v) The practitioner or pharmacist terminates the designee's access to the database at the termination of the designee's employment; and

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

(4) Pursuant to a valid search warrant based on probable cause to believe a violation of federal or state criminal law has occurred and that specified information contained in the database would assist in the investigation of the crime;

(5) To a patient who requests his or her own prescription information, or the parent or legal guardian of a minor child who requests the minor child's prescription information;

(6) To a health professional regulatory board that documents, 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;

(7) To any vendor or contractor with whom the department has contracted to establish or maintain the electronic system of the prescription drug monitoring database; or

(8) To public or private entities for statistical, research, or educational purposes, after removing the patient and prescriber information that could be used to identify individual patients. This shall not include entities receiving a waiver from the institutional review board.

...

West's Rhode Island Administrative Code (2014)

Title 31. Health Department

Division 2. Drug Control

Rule 1. Rules and Regulations Governing Electronic Data Transfer of Controlled Substances in Schedules II and III

31-2-1:3.0. Data Collection

3.1 The electronic system shall provide for the method of data collection; transmission from all dispensers to the Department; maintenance and use of data; and shall be as set forth in the latest edition of the ASAP Telecommunications Format for Controlled Substances of reference 1 herein.

3.2 Required data shall be transmitted by direct computer link, double sided/high density micro floppy disk, or microcassette. All computerized pharmacies shall submit the required data no later than 1 July 1997.

3.3 The Department shall:

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3.3.1 be authorized to provide data in the electronic prescription system to other regulatory, investigative or law enforcement agencies for disciplinary, civil, or criminal purposes, and for the purposes of educating practitioners in lieu of disciplinary, civil or criminal action.

3.3.2 be authorized to provide data to appropriate public or private entities for statistical, research, or educational purposes provided that the privacy and confidentiality of patients and patient information is not compromised.

3.3.3 in using the information for investigative or prosecutorial purposes, consider the nature of the prescriber's or dispenser's practice and the condition(s) for which the patient is being treated.

3.3.4 ensure the privacy and confidentiality of patients and shall ensure that patient information collected, recorded, transmitted, and stored in the prescription system is maintained in accordance with applicable state and federal laws, rules and regulations.

3.3.5 ensure that the EDT program does not infringe on the legal use of any schedule II or III controlled substance.

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South Carolina

§ 44-53-1650

Code of Laws of South Carolina 1976 Annotated (2014)

Title 44. Health

Chapter 53. Poisons, Drugs and Other Controlled Substances

Article 15. Prescription Monitoring Program

§ 44-53-1650. Confidentiality; persons to whom data may be released.

(A) Prescription information submitted to drug control is confidential and not subject to public disclosure under the Freedom of Information Act or any other provision of law, except as provided in subsections (C) and (D).

(B) Drug control 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 subsections (C) and (D).

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

(D) Drug control may provide data in the prescription monitoring program to the following persons:

(1) a practitioner or pharmacist or authorized delegate who requests information and certifies that the requested information is for the purpose of providing medical or pharmaceutical treatment to a bona fide patient;

(2) an individual who requests the individual's own prescription monitoring information in accordance with procedures established pursuant to state law;

(3) a designated representative of the South Carolina Department of Labor, Licensing and Regulation responsible for the licensure, regulation, or discipline of practitioners, pharmacists, 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;

(4) a local, state, or federal law enforcement or prosecutorial official engaged in the administration, investigation, or enforcement of the laws governing licit drugs and who is involved in a bona fide specific drug related investigation involving a designated person;

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(5) the South Carolina Department of Health and Human Services regarding Medicaid program recipients;

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

(7) personnel of drug control for purposes of administration and enforcement of this article;

(8) 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. Further, release of the information only may be made pursuant to a written agreement between qualified personnel and the department in order to ensure compliance with this Subsection.

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South Dakota
§ 34-20E-7

South Dakota Codified Laws (2014)
Title 34. Public Health and Safety
Chapter 34-20E. Prescription Drug Monitoring Program

§ 34-20E-7. Disclosure of data in central repository to certain persons and entities

Unless disclosure is prohibited by law, the board may provide data in the central repository to:

- (1) Any prescriber for the purpose of providing medical care to a patient, a dispenser for the purpose of filling a prescription or providing pharmaceutical care for a patient, a prescriber or dispenser inquiring about the prescriber's or dispenser's own prescribing activity, or a prescriber or dispenser in order to further the purposes of the program;
- (2) Any individual who requests the prescription information of the individual or the individual's minor child;
- (3) Any state board or regulatory agency that is responsible for the licensing of individuals authorized to prescribe or dispense controlled substances if the board or regulatory agency is seeking information from the central repository that is relevant to an investigation of an individual who holds a license issued by that board or regulatory agency;
- (4) Any local, state, and federal law enforcement or prosecutorial officials engaged in the enforcement of laws relating to controlled substances who seek information for the purpose of an investigation or prosecution of the drug-related activity or probation compliance of an individual;
- (5) The Department of Social Services for purposes regarding the utilization of controlled substances by a medicaid recipient;
- (6) Any insurer for purposes regarding the utilization of controlled substances by a claimant;
- (7) Any judicial authority under grand jury subpoena or court order or equivalent judicial process for investigation of criminal violations of controlled substances laws;
- (8) Any public or private entity for statistical, research, or educational purposes after the information is de-identified with respect to any prescriber, dispenser, or patient who received a prescription for a controlled substance; or**

(9) Any peer review committee, which means any committee of a health care organization, composed of health care providers, employees, administrators, consultants, agents, or members of the health care organization's governing body, which conducts professional peer review.

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Tennessee
§ 53-10-306

West's Tennessee Code Annotated (2014)
Title 53. Food, Drugs and Cosmetics
Chapter 10. Legend Drugs
Part 3. Tennessee Prescription Safety Act of 2012

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

...

(o) Authorized committee, board, or department personnel and 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 may publish, or otherwise make available to prescribers, dispensers and to the general public, aggregate unidentifiable personal data contained in or derived from the database for the purpose of educational outreach.

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Texas
Health & Safety Code § 481.076

Vernon's Texas Statutes and Codes Annotated (2014)
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

(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.

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(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 a 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.

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Utah
§ 58-37f-301
ADC R156-37f
ADC R384-203

West's Utah Code Annotated (2014)
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;

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(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;

...

Utah Administrative Code (2014)

Commerce

R156. Occupational and Professional Licensing.

R156-37f. Controlled Substance Database Act Rule.

...

(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:

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(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.

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Utah Administrative Code (2014)

Health

R384. Health, Disease Control and Prevention, Health Promotion.

R384-203. Prescription Drug Database Access.

R384-203-1. Authority and Purpose.

This rule establishes procedures and application processes pursuant to Title 58-37f-301(2)(d) for Utah Department of Health Executive Director to allow access to the Prescription Drug database by a designated and assigned person to conduct scientific studies regarding the use or abuse of controlled substances, who is not an employee of the Department of Health.

R384-203-2. Definitions.

The following definitions apply to this rule:

(1) “Department” means the Utah Department of Health.

(2) “Director” means the Utah Department of Health Executive Director.

(3) “Prescription Drug Database” means the Utah Controlled Substance Database.

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(4) “Research facility” means a research facility associated with a university or college in the state accredited by the Northwest Commission on Colleges and Universities.

(5) “Institutional Review Board” means a board that is approved for human subject research by the United States Department of Health and Human Services.

(6) “Designee” means a person designated and assigned by the Director to have access to the Prescription Drug database in order to conduct scientific studies regarding the use or abuse of controlled substances, who is not an employee of the Department.

(7) “Business associate” means a business associate as defined under the HIPAA privacy, security, and breach notification rules in 45 CFR 164.502(a), 164.504(e), and 164.532(d) and (e).

(8) “De-identified” means information as defined in 45 CFR 164.502(d) and 164.514(a), (b), and (c).

R384-203-3. Criteria for Application to Access Prescription Drug Database.

(1) The study must fit within the responsibilities of the Department for health and welfare.

(2) De-identified prescriber, patient and pharmacy data will meet the research needs.

(3) The research facility designee must provide:

(a) written assurances that the studies are not conducted for and will not be used for profit or commercial gain;

(b) written assurances that the designee shall protect the information as a business associate of the Department of Health; and

(c) documentation of an Institutional Review Board approval.

R384-203-4. Research Application Process.

(1) The research facility designee will prepare and submit for Department approval an application as designated by the Department detailing explicit information regarding the scientific studies to be conducted including the:

(a) purpose of the study;

(b) research protocol for the project;

(c) description of the data needed from the database to conduct that research;

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(d) plan that demonstrates all database information will be maintained securely, with access being strictly restricted to the designee and research study staff; and

(e) provisions for electronic data to be stored on a secure database computer system with access being strictly restricted to the designee and research study staff.

(2) Application will be reviewed by the Department's Institutional Review Board and recommendation made to the director for or against approval.

(3) Director will determine approval status of the application.

(4) Designee will sign the Department's data sharing agreement if application is approved by the Director.

R384-203-5. Data Provision and Fees.

(1) Department will obtain, de-identify and provide the data set requested in the application.

(2) Research facility and designee shall pay all relevant expenses for data transfer and manipulation.

R384-203-6. Audit Provisions.

Research facility and designee shall submit, upon request, to a Department audit of the recipients' compliance with the terms of the data sharing agreement.

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Vermont
18 §4284

West's Vermont Statutes Annotated (2014)
Title Eighteen. Health
Part 5. Foods and Drugs
Chapter 84A. Vermont Prescription Monitoring System

§ 4284. Protection and disclosure of information

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(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.

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Virginia
§ 54.1-2523

West's Annotated Code of Virginia (2014)

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. Confidentiality of data; disclosure of information; discretionary authority of Director

A. All data, records, and reports relating to the prescribing and dispensing of covered substances to recipients and any abstracts from such data, records, and reports that are in the possession of the Prescription Monitoring Program pursuant to this chapter and any material relating to the operation or security of the program shall be confidential and shall be exempt from the Virginia Freedom of Information Act (§ 2.2-3700 et seq.) pursuant to subdivision 15 of § 2.2-3705.5. Further, the Director shall only have discretion to disclose any such information as provided in subsections B and C.

B. Upon receiving a request for information in accordance with the Department's regulations and in compliance with applicable federal law and regulations, the Director shall disclose the following:

1. Information relevant to a specific investigation of a specific recipient or of a specific dispenser or prescriber to an agent who has completed the Virginia State Police Drug Diversion School designated by the superintendent of the Department of State Police or designated by the chief law-enforcement officer of any county, city, or town or campus police department to conduct drug diversion investigations pursuant to § 54.1-3405.
2. Information relevant to an investigation or inspection of or allegation of misconduct by a specific person licensed, certified, or registered by or an applicant for licensure, certification, or registration by a health regulatory board; information relevant to a disciplinary proceeding before a health regulatory board or in any subsequent trial or appeal of an action or board order to designated employees of the Department of Health Professions; or to designated persons operating the Health Practitioners' Monitoring Program pursuant to Chapter 25.1 (§ 54.1-2515 et seq.).
3. Information relevant to the proceedings of any investigatory grand jury or special grand jury that has been properly impaneled in accordance with the provisions of Chapter 13 (§ 19.2-191 et seq.) of Title 19.2.

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4. Information relevant to a specific investigation of a specific recipient, dispenser, or prescriber to an agent of a federal law-enforcement agency with authority to conduct drug diversion investigations.

C. In accordance with the Department's regulations and applicable federal law and regulations, the Director may, in his discretion, disclose:

1. Information in the possession of the program concerning a recipient who is over the age of 18 to that recipient. The information shall be mailed to the street or mailing address indicated on the recipient request form.

2. Information on a specific recipient to a prescriber, as defined in this chapter, for the purpose of establishing the treatment history of the specific recipient when such recipient is either under care and treatment by the prescriber or the prescriber is initiating treatment of such recipient. In a manner specified by the Director in regulation, notice shall be given to patients that information may be requested by the prescriber from the Prescription Monitoring Program.

3. Information on a specific recipient to a dispenser for the purpose of establishing a prescription history to assist the dispenser in determining the validity of a prescription in accordance with § 54.1-3303 when the recipient is seeking a covered substance from the dispenser or the facility in which the dispenser practices. In a manner specified by the Director in regulation, notice shall be given to patients that information may be requested by the dispenser from the Prescription Monitoring Program.

4. Information relevant to an investigation or regulatory proceeding of a specific dispenser or prescriber to other regulatory authorities concerned with granting, limiting or denying licenses, certificates or registrations to practice a health profession when such regulatory authority licenses such dispenser or prescriber or such dispenser or prescriber is seeking licensure by such other regulatory authority.

5. Information relevant to an investigation relating to a specific dispenser or prescriber who is a participating provider in the Virginia Medicaid program or information relevant to an investigation relating to a specific recipient who is currently eligible for and receiving or who has been eligible for and has received medical assistance services to the Medicaid Fraud Control Unit of the Office of the Attorney General or to designated employees of the Department of Medical Assistance Services, as appropriate.

6. Information relevant to determination of the cause of death of a specific recipient to the designated employees of the Office of the Chief Medical Examiner.

7. Information for the purpose of bona fide research or education to qualified personnel; however, data elements that would reasonably identify a specific recipient, prescriber, or dispenser shall be deleted or redacted from such information prior to disclosure. Further, release of the information shall only be made pursuant to a written agreement between

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such qualified personnel and the Director in order to ensure compliance with this subdivision.

8. Information relating to prescriptions for covered substances issued by a specific prescriber, which have been dispensed and reported to the Program, to that prescriber.

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Washington
§ 70.225.040

West's Revised Code of Washington Annotated (2014)
Title 70. Public Health and Safety
Chapter 70.225. Prescription Monitoring Program

§ 70.225.040. Confidentiality of prescription information--Procedures--Immunity when acting in good faith

...

(4) The department may provide data to public or private entities for statistical, research, or educational purposes after removing information that could be used to identify individual patients, dispensers, prescribers, and persons who received prescriptions from dispensers.

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West Virginia
§ 60A-9-5

West's Annotated Code of West Virginia (2014)
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**

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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.

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Wisconsin
ADC Phar. 18.11

Wisconsin Administrative Code (2014)
Pharmacy Examining Board
Chapter Phar 18. Prescription Drug Monitoring Program

Phar 18.11 Methods of obtaining PDMP information.

...

(9) The board shall disclose the minimum amount of PDMP information necessary to a researcher 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 if the person does all of the following:

(a) Creates an account with the board on a form provided by the board.

(b) Provides proof sufficient to the board that the person is entitled to the information under s. 146.82 (2) (a) 6. or 20., Stats.

(c) Makes a request for the PDMP information through its account with the board.

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Wyoming
§ 35-7-1060

West's Wyoming Statutes Annotated (2013)
Title 35. Public Health and Safety
Chapter 7. Food and Drugs
Article 10. Controlled Substances
Article X

§ 35-7-1060. Controlled substances prescription tracking program

...

(c) The tracking program shall not be used to infringe on the legal use of a controlled substance. Information obtained through the controlled substance prescription tracking program is confidential and may not be released and is not admissible in any judicial or administrative proceeding, except as follows:

(i) The board may release information to practitioners and pharmacists when the release of the information may be of assistance in preventing or avoiding inappropriate use of controlled substances;

(ii) The board shall report any information that it reasonably suspects may relate to fraudulent or illegal activity to the appropriate law enforcement agency and the relevant occupational licensing board;

(iii) The board may release information to the patient to whom the information pertains or his agent or, if the patient is a minor, to his parents or guardian;

(iv) The board may release information to a third party if the patient has signed a consent specifically for the release of his controlled substance prescription information to the specific third party;

(v) The board may release information that does not identify individual patients, practitioners, pharmacists or pharmacies, for educational, research or public information purposes; and

(vi) Subject to the rules of evidence, information obtained from the program is admissible in a criminal proceeding or an administrative proceeding involving professional licensing.

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