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"CDS registration" means registration with the Division of Consumer Affairs to manufacture, distribute, dispense, or conduct research with controlled dangerous substances issued pursuant to section 11 of P.L.1970, c.226 (C.24:21-11).

"Certified medical assistant" means a person who is a graduate of a post-secondary medical assisting educational program accredited by the American Medical Association's Committee on Allied Health Education and Accreditation (CAHEA), or its successor, the Accrediting Bureau of Health Education Schools (ABHES), or its successor, or any accrediting agency recognized by the U.S. Department of Education, which educational program includes, at a minimum, 600 clock hours of instruction, and encompasses training in the administration of intramuscular and subcutaneous injections, as well as instruction and demonstration in: pertinent anatomy and physiology appropriate to injection procedures; choice of equipment; proper technique, including sterile technique; hazards and complications; and emergency procedures; and who maintains current certification or registration, as appropriate, from the Certifying Board of the American Association of Medical Assistants (AAMA), the National Center for Competency Testing (NCCT), the American Medical Technologists (AMT), or any other recognized certifying body approved by the Board of Medical Examiners.

"Controlled dangerous substance" means any substance that is listed in Schedules II, III, and IV of the schedules provided under the "New Jersey Controlled Dangerous Substances Act," P.L.1970, c.226 (C.24:21-1 et seq.). Controlled dangerous substance also means any substance that is listed in Schedule V under the "New Jersey Controlled Dangerous Substances Act" when the director has determined that reporting Schedule V substances is required by federal law, regulation, or funding eligibility.

"Dental resident" means a person who practices dentistry as a resident pursuant to R.S.45:6-20 and, pursuant to N.J.A.C.13:30-1.3, is a graduate of a dental school approved by the Commission on Dental Accreditation and has passed Part I and Part II of the National Board Dental examination and obtained a resident permit from the New Jersey Board of Dentistry.

"Director" means the Director of the Division of Consumer Affairs in the Department of Law and Public Safety.

"Division" means the Division of Consumer Affairs in the Department of Law and Public Safety.

"Licensed health care professional" means a registered nurse, licensed practical nurse, advanced practice nurse, physician assistant, or dental hygienist licensed pursuant to Title 45 of the Revised Statutes.

"Licensed pharmacist" means a pharmacist licensed pursuant to P.L.2003, c.280 (C.45:14-40 et seq.).

"Medical resident" means a graduate physician who is authorized to practice medicine and surgery by means of a valid permit issued by the State Board of Medical Examiners to a person authorized to engage in the practice of medicine and surgery while in the second year or beyond of a graduate medical education program pursuant to N.J.A.C.13:35-1.5.

"Mental health practitioner" means a clinical social worker, marriage and family therapist, alcohol and drug counselor, professional counselor, psychologist, or psychoanalyst licensed or otherwise authorized to practice pursuant to Title 45 of the Revised Statutes.
"Pharmacy permit holder" means an individual or business entity that holds a permit to operate a pharmacy practice site pursuant to P.L.2003, c.280 (C.45:14-40 et seq.)

"Practitioner" means an individual currently licensed, registered, or otherwise authorized by this State or another state to prescribe drugs in the course of professional practice.

"Registered dental assistant" is a person who has fulfilled the requirements for registration established by "The Dental Auxiliaries Act," P.L.1979, c.46 (C.45:6-48 et al.) and works under the direct supervision of a licensed dentist.

"Ultimate user" means a person who has obtained from a dispenser and possesses for the person's own use, or for the use of a member of the person's household or an animal owned by the person or by a member of the person's household, a controlled dangerous substance.

L.2007, c.244, s.24; amended 2015, c.74, s.2.

45:1-45 Prescription Monitoring Program; requirements.

25. Prescription Monitoring Program; requirements.

a. There is established the Prescription Monitoring Program in the Division of Consumer Affairs in the Department of Law and Public Safety. The program shall consist of an electronic system for monitoring controlled dangerous substances that are dispensed in or into the State by a pharmacist in an outpatient setting.

b. Each pharmacy permit holder shall submit, or cause to be submitted, to the division, by electronic means in a format and at such intervals as are specified by the director, information about each prescription for a controlled dangerous substance dispensed by the pharmacy that includes:

(1) The surname, first name, and date of birth of the patient for whom the medication is intended;

(2) The street address and telephone number of the patient;

(3) The date that the medication is dispensed;

(4) The number or designation identifying the prescription and the National Drug Code of the drug dispensed;

(5) The pharmacy permit number of the dispensing pharmacy;

(6) The prescribing practitioner's name and Drug Enforcement Administration registration number;

(7) The name, strength, and quantity of the drug dispensed, the number of refills ordered, and whether the drug was dispensed as a refill or a new prescription;

(8) The date that the prescription was issued by the practitioner;

(9) The source of payment for the drug dispensed;
(10) Identifying information for any individual, other than the patient for whom the prescription was written, who picks up a prescription, if the pharmacist has a reasonable belief that the person picking up the prescription may be seeking a controlled dangerous substance, in whole or in part, for any reason other than delivering the substance to the patient for the treatment of an existing medical condition; and

(11) Such other information, not inconsistent with federal law, regulation, or funding eligibility requirements, as the director determines necessary.

The pharmacy permit holder shall submit the information to the division with respect to the prescriptions dispensed during the reporting period not less frequently than every seven days.

c. The division may grant a waiver of electronic submission to any pharmacy permit holder for good cause, including financial hardship, as determined by the director. The waiver shall state the format in which the pharmacy permit holder shall submit the required information.

d. The requirements of this act shall not apply to: the direct administration of a controlled dangerous substance to the body of an ultimate user; or the administration or dispensing of a controlled dangerous substance that is otherwise exempted as determined by the Secretary of Health and Human Services pursuant to the "National All Schedules Prescription Electronic Reporting Act of 2005," Pub.L.109-60.

e. The provisions of paragraph (10) of subsection b. of this section shall not take effect until the director determines that the Prescription Monitoring Program has the technical capacity to accept the information required by that paragraph.

L.2007, c.244, s.25; amended 2015, c.74, s.3.

45:1-45.1 Information required for monitoring; rules, regulations.

11. a. A physician who provides a certification or written instruction for the medical use of marijuana to a qualifying patient pursuant to P.L.2009, c.307 (C.24:6I-1 et al.) and any alternative treatment center shall furnish to the Director of the Division of Consumer Affairs in the Department of Law and Public Safety such information, in such a format and at such intervals, as the director shall prescribe by regulation, for inclusion in a system established to monitor the dispensation of marijuana in this State for medical use as authorized by the provisions of P.L.2009, c.307 (C.24:6I-1 et al.), which system shall serve the same purpose as, and be cross-referenced with, the electronic system for monitoring controlled dangerous substances established pursuant to section 25 of P.L.2007, c.244 (C.45:1-45).

b. The Director of the Division of Consumer Affairs, pursuant to the "Administrative Procedure Act," P.L.1968, c.410 (C.52:14B-1 et seq.), and in consultation with the Commissioner of Health and Senior Services, shall adopt rules and regulations to effectuate the purposes of subsection a. of this section.

c. Notwithstanding any provision of P.L.1968, c.410 to the contrary, the Director of the Division of Consumer Affairs shall adopt, immediately upon filing with the Office of Administrative Law and no later than the 90th day after the effective date of P.L.2009, c.307 (C.24:6I-1 et al.), such regulations as the director deems necessary to implement the provisions of subsection a. of this section. Regulations adopted pursuant to this subsection shall be effective until the adoption of rules and

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regulations pursuant to subsection b. of this section and may be amended, adopted, or readopted by the director in accordance with the requirements of P.L.1968, c.410.

L.2009, c.307, s.11.


a. The division shall maintain procedures to ensure privacy and confidentiality of patients and that patient information collected, recorded, transmitted, and maintained is not disclosed, except as permitted in this section, including, but not limited to, the use of a password-protected system for maintaining this information and permitting access thereto as authorized under sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50), and a requirement that a person as listed in subsection h. or i. of this section provide affirmation of the person's intent to comply with the provisions of sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50) as a condition of accessing the information.

b. The prescription monitoring information submitted to the division shall be confidential and not be subject to public disclosure under P.L.1963, c.73 (C.47:1A-1 et seq.), or P.L.2001, c.404 (C.47:1A-5 et al.).

c. The division shall review the prescription monitoring information provided by a pharmacy permit holder pursuant to sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50). The review shall include, but not be limited to:

(1) a review to identify whether any person is obtaining a prescription in a manner that may be indicative of misuse, abuse, or diversion of a controlled dangerous substance. The director shall establish guidelines regarding the terms "misuse," "abuse," and "diversion" for the purposes of this review. When an evaluation of the information indicates that a person may be obtaining a prescription for the same or a similar controlled dangerous substance from multiple practitioners or pharmacists during the same time period, the division may provide prescription monitoring information about the person to practitioners and pharmacists; and

(2) a review to identify whether a violation of law or regulation or a breach of the applicable standards of practice by any person may have occurred, including, but not limited to, diversion of a controlled dangerous substance. If the division determines that such a violation or breach may have occurred, the division shall notify the appropriate law enforcement agency or professional licensing board, and provide the prescription monitoring information required for an investigation.

d. (Deleted by amendment, P.L.2015, c.74)

e. (Deleted by amendment, P.L.2015, c.74)

f. (Deleted by amendment, P.L.2015, c.74)

g. (Deleted by amendment, P.L.2015, c.74)

h. (1) The division shall register a practitioner to access prescription monitoring information upon issuance or renewal of the practitioner's CDS registration.
(2) The division shall provide to a pharmacist who is employed by a current pharmacy permit holder online access to prescription monitoring information for the purpose of providing health care to a current patient or verifying information with respect to a patient or a prescriber.

(3) The division shall provide to a practitioner who has a current CDS registration online access to prescription monitoring information for the purpose of providing health care to a current patient or verifying information with respect to a patient or a prescriber. The division shall also grant online access to prescription monitoring information to as many licensed health care professionals as are authorized by a practitioner to access that information and for whom the practitioner is responsible for the use or misuse of that information, subject to a limit on the number of such health care professionals as deemed appropriate by the division for that particular type and size of professional practice, in order to minimize the burden to practitioners to the extent practicable while protecting the confidentiality of the prescription monitoring information obtained. The director shall establish, by regulation, the terms and conditions under which a practitioner may delegate that authorization, including procedures for authorization and termination of authorization, provisions for maintaining confidentiality, and such other matters as the division may deem appropriate.

(4) The division shall provide online access to prescription monitoring information to as many medical or dental residents as are authorized by a faculty member of a medical or dental teaching facility to access that information and for whom the practitioner is responsible for the use or misuse of that information. The director shall establish, by regulation, the terms and conditions under which a faculty member of a medical or dental teaching facility may delegate that authorization, including procedures for authorization and termination of authorization, provisions for maintaining confidentiality, provisions regarding the duration of a medical or dental resident's authorization to access prescription monitoring information, and such other matters as the division may deem appropriate.

(5) The division shall provide online access to prescription monitoring information to as many certified medical assistants as are authorized by a practitioner to access that information and for whom the practitioner is responsible for the use or misuse of that information. The director shall establish, by regulation, the terms and conditions under which a practitioner may delegate that authorization, including procedures for authorization and termination of authorization, provisions for maintaining confidentiality, provisions regarding the duration of a certified medical assistant's authorization to access prescription monitoring information, and such other matters as the division may deem appropriate.

(6) The division shall provide online access to prescription monitoring information to as many registered dental assistants as are authorized by a licensed dentist to access that information and for whom the licensed dentist is responsible for the use or misuse of that information. The director shall establish, by regulation, the terms and conditions under which a licensed dentist may delegate that authorization, including procedures for authorization and termination of authorization, provisions for maintaining confidentiality, provisions regarding the duration of a registered dental assistant's authorization to access prescription monitoring information, and such other matters as the division may deem appropriate.

(7) A person listed in this subsection, as a condition of accessing prescription monitoring information pursuant thereto, shall certify that the request is for the purpose of providing health care to a current patient or verifying information with respect to a patient or practitioner. Such certification shall be furnished through means of an online statement or alternate means.
authorized by the director, in a form and manner prescribed by rule or regulation adopted by the director.

i. The division may provide online access to prescription monitoring information, or may provide access to prescription monitoring information through any other means deemed appropriate by the director, to the following persons:

(1) authorized personnel of the division or a vendor or contractor responsible for maintaining the Prescription Monitoring Program;

(2) authorized personnel of the division responsible for administration of the provisions of P.L.1970, c.226 (C.24:21-1 et seq.);

(3) the State Medical Examiner, a county medical examiner, a deputy or assistant county medical examiner, or a qualified designated assistant thereof, who certifies that the request is for the purpose of investigating a death pursuant to P.L.1967, c.234 (C.52:17B-78 et seq.);

(4) a controlled dangerous substance monitoring program in another state with which the division has established an interoperability agreement, or which participates with the division in a system that facilitates the secure sharing of information between states;

(5) a designated representative of the State Board of Medical Examiners, New Jersey State Board of Dentistry, State Board of Nursing, New Jersey State Board of Optometrists, State Board of Pharmacy, State Board of Veterinary Medical Examiners, or any other board in this State or another state that regulates the practice of persons who are authorized to prescribe or dispense controlled dangerous substances, as applicable, who certifies that the representative is engaged in a bona fide specific investigation of a designated practitioner or pharmacist whose professional practice was or is regulated by that board;

(6) a State, federal, or municipal law enforcement officer who is acting pursuant to a court order and certifies that the officer is engaged in a bona fide specific investigation of a designated practitioner, pharmacist, or patient. A law enforcement agency that obtains prescription monitoring information shall comply with security protocols established by the director by regulation;

(7) a designated representative of a state Medicaid or other program who certifies that the representative is engaged in a bona fide investigation of a designated practitioner, pharmacist, or patient;

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

(9) a licensed mental health practitioner providing treatment for substance abuse to patients at a residential or outpatient substance abuse treatment center licensed by the Division of Mental Health and Addiction Services in the Department of Human Services, who certifies that the request is for the purpose of providing health care to a current patient or verifying information with respect to a patient or practitioner, and who furnishes the division with the written consent of the patient for the mental health practitioner to obtain prescription monitoring information about the patient. The director shall establish, by regulation, the terms and conditions under which a mental health practitioner may request and receive prescription monitoring information. Nothing in sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50) shall be construed to require or obligate a mental health practitioner to
access or check the prescription monitoring information in the course of treatment beyond that which may be required as part of the mental health practitioner's professional practice.

j. A person listed in subsection i. of this section, as a condition of obtaining prescription monitoring information pursuant thereto, shall certify the reasons for seeking to obtain that information. Such certification shall be furnished through means of an online statement or alternate means authorized by the director, in a form and manner prescribed by rule or regulation adopted by the director.

k. The division shall offer an online tutorial for those persons listed in subsections h. and i. of this section, which shall, at a minimum, include: how to access prescription monitoring information; the rights of persons who are the subject of this information; the responsibilities of persons who access this information; a summary of the other provisions of sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50) and the regulations adopted pursuant thereto, regarding the permitted uses of that information and penalties for violations thereof; and a summary of the requirements of the federal health privacy rule set forth at 45 CFR Parts 160 and 164 and a hypertext link to the federal Department of Health and Human Services website for further information about the specific provisions of the privacy rule.

l. The division may request and receive prescription monitoring information from prescription monitoring programs in other states and may use that information for the purposes of sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50). When sharing data with programs in another state, the division shall not be required to obtain a memorandum of understanding unless required by the other state.

m. The director may provide nonidentifying prescription drug monitoring information to public or private entities for statistical, research, or educational purposes, in accordance with the provisions of sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50).

n. Nothing shall be construed to prohibit the division from obtaining unsolicited automated reports from the program or disseminating such reports to pharmacists, practitioners, mental health care practitioners, and other licensed health care professionals.

o. (1) A current patient of a practitioner may request from that practitioner that patient's own prescription monitoring information that has been submitted to the division pursuant to sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50). A parent or legal guardian of a child who is a current patient of a practitioner may request from that practitioner the child's prescription monitoring information that has been submitted to the division pursuant to sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50).

(2) Upon receipt of a request pursuant to paragraph (1) of this subsection, a practitioner or health care professional authorized by that practitioner may provide the current patient or parent or legal guardian, as the case may be, with access to or a copy of the prescription monitoring information pertaining to that patient or child.

(3) The division shall establish a process by which a patient, or the parent or legal guardian of a child who is a patient, may request a pharmacy permit holder that submitted prescription monitoring information concerning a prescription for controlled dangerous substances for that patient or child to the division pursuant to sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50) to correct information that the person believes to have been inaccurately
entered into that patient's or child's prescription profile. Upon confirmation of the inaccuracy of any such entry into a patient's or child's prescription profile, the pharmacy permit holder shall be authorized to correct any such inaccuracies by submitting corrected information to the division pursuant to sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50). The process shall provide for review by the Board of Pharmacy of any disputed request for correction, which determination shall be appealable to the director.

p. The division shall take steps to ensure that appropriate channels of communication exist to enable any licensed health care professional, licensed pharmacist, mental health practitioner, pharmacy permit holder, or other practitioner who has online access to the Prescription Monitoring Program pursuant to this section to seek or provide information to the division related to the provisions of this section. (cf: P.L.2007, c.244, s.26)

L.2007, c.244, s.26; amended 2015, c.74, s.4.

45:1-46.1  Proper time to access prescription monitoring information; restrictions in dispensing Schedule II controlled dangerous substance; exceptions.

8. a. (1) Except as provided in subsection b. of this section, a practitioner or other person who is authorized by a practitioner to access prescription monitoring information pursuant to subsection h. of section 26 of P.L.2007, c.244 (C.45:1-46) shall access prescription monitoring information the first time the practitioner or other person prescribes a Schedule II controlled dangerous substance to a new patient for acute or chronic pain. In addition, for any prescription of a Schedule II controlled dangerous substance for a new or current patient for acute or chronic pain which is written on or after the effective date of P.L.2015, c.74 (C.45:1-46.1 et al.) a practitioner or other authorized person shall access prescription monitoring information on a quarterly basis during the period of time the patient continues to receive such prescriptions.

(2) (a) A pharmacist shall not dispense a Schedule II controlled dangerous substance to any person without first accessing the prescription monitoring information, as authorized pursuant to subsection h. of section 26 of P.L.2007, c.244 (C.45:1-46), to determine if the person has received other prescriptions that indicate misuse, abuse, or diversion, if the pharmacist has a reasonable belief that the person may be seeking a controlled dangerous substance, in whole or in part, for any purpose other than the treatment of an existing medical condition, such as for purposes of misuse, abuse, or diversion.

(b) A pharmacist shall not dispense a prescription to a person other than the patient for whom the prescription is intended, unless the person picking up the prescription provides personal identification to the pharmacist, and the pharmacist, as required by subsection b. of section 25 of P.L.2007, c.244 (C.45:1-45), inputs that identifying information into the Prescription Monitoring Program if the pharmacist has a reasonable belief that the person may be seeking a controlled dangerous substance, in whole or in part, for any reason other than delivering the substance to the patient for the treatment of an existing medical condition. The provisions of this subparagraph shall not take effect until the director determines that the Prescription Monitoring Program has the technical capacity to accept such information.

b. The provisions of subsection a. of this section shall not apply to:

(1) a veterinarian;
(2) a practitioner or the practitioner’s agent administering methadone, or another controlled dangerous substance designated by the director as appropriate for treatment of a patient with a substance abuse disorder, as interim treatment for a patient on a waiting list for admission to an authorized substance abuse treatment program;

(3) a practitioner administering a controlled dangerous substance directly to a patient;

(4) a practitioner prescribing a controlled dangerous substance to be dispensed by an institutional pharmacy, as defined in N.J.A.C.13:39-9.2;

(5) a practitioner prescribing a controlled dangerous substance in the emergency department of a general hospital, provided that the quantity prescribed does not exceed a five-day supply of the substance;

(6) a practitioner prescribing a controlled dangerous substance to a patient under the care of a hospice;

(7) a situation in which it is not reasonably possible for the practitioner or pharmacist to access the Prescription Monitoring Program in a timely manner, no other individual authorized to access the Prescription Monitoring Program is reasonably available, and the quantity of controlled dangerous substance prescribed or dispensed does not exceed a five-day supply of the substance;

(8) a practitioner or pharmacist acting in compliance with regulations promulgated by the director as to circumstances under which consultation of the Prescription Monitoring Program would result in a patient's inability to obtain a prescription in a timely manner, thereby adversely impacting the medical condition of the patient;

(9) a situation in which the Prescription Monitoring Program is not operational as determined by the division or where it cannot be accessed by the practitioner due to a temporary technological or electrical failure, as set forth in regulation;

(10) a practitioner or pharmacist who has been granted a waiver due to technological limitations that are not reasonably within the control of the practitioner or pharmacist, or other exceptional circumstances demonstrated by the practitioner or pharmacist, pursuant to a process established in regulation, and in the discretion of the director; or

(11) a practitioner who is prescribing a controlled dangerous substance to a patient immediately after the patient has undergone an operation, procedure, or treatment for acute trauma, when less than a 30-day supply is prescribed.

L.2015, c.74, s.8.

45:1-47 Prescription monitoring program; provisions for expansion.

27. Prescription Monitoring Program; provisions for expansion.

a. Notwithstanding the provisions of section 25 of P.L.2007, c.244 (C.45:1-45) to the contrary, the director may adopt a regulation to expand the program to include information about each prescription dispensed for a prescription drug that is not a controlled dangerous substance. In determining whether a prescription drug other than a controlled dangerous substance should be
monitored, the director shall consider: the actual or relative potential for abuse; scientific evidence of its pharmacological effect, if known; the state of current scientific knowledge regarding the drug; its history and current pattern of abuse; the scope, duration and significance of abuse; what, if any, risk to the public health; and its psychic or physiological dependence liability. The regulation shall provide that the prescription drug shall be monitored for a period of time. At the conclusion of the monitoring period, the director shall publish and make public the decision of whether inclusion of the prescription drug in the program shall be permanent.

b. At the time the notice to expand the program pursuant to subsection a. is published in the New Jersey Register, the director shall provide a copy of the notice of proposed rule making to the chairpersons of the standing legislative reference committees on health of the Senate and General Assembly.

L.2007, c.244, s.27.

45:1-48 Immunity from liability.

28. Immunity from liability.

a. The division shall be immune from civil liability arising from inaccuracy of any of the information submitted to it pursuant to sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50).

b. A pharmacy permit holder, pharmacist, mental health practitioner, licensed health care professional, or practitioner shall be immune from civil liability arising from compliance with sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50).

L.2007, c.244, s.28; amended 2015, c.74, s.5.

45:1-49 Penalties.

29. Penalties.

a. A pharmacy permit holder, or a person designated by a pharmacy permit holder to be responsible for submitting data required by section 25 of P.L.2007, c.244 (C.45:1-45), who knowingly fails to submit data as required, shall be subject to disciplinary action pursuant to section 8 of P.L.1978, c.73 (C.45:1-21) and may be subject to a civil penalty in an amount not to exceed $1,000 for failure to comply with sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50).

b. (1) A pharmacy permit holder, pharmacist, mental health practitioner, licensed health care professional, or practitioner, or any other person or entity who knowingly obtains or attempts to obtain prescription monitoring information in violation of the provisions of sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50) shall be subject to a civil penalty in an amount not to exceed $10,000.

(2) A pharmacy permit holder, pharmacist, mental health practitioner, licensed health care professional, or practitioner who knowingly discloses or uses prescription monitoring information in violation of the provisions of sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50), shall also be subject to disciplinary action pursuant to section 8 of P.L.1978, c.73 (C.45:1-21).
c. In addition to any other penalty provided by law, a person who is authorized to obtain prescription monitoring information from the Prescription Monitoring Program who knowingly discloses such information in violation of the provisions of sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50) shall be guilty of a crime of the fourth degree and shall be subject to a civil penalty in an amount not to exceed $10,000.

d. In addition to any other penalty provided by law, a person who is authorized to obtain prescription monitoring information from the Prescription Monitoring Program who uses this information in the course of committing, attempting to commit, or conspiring to commit any criminal offense shall be guilty of a crime of the third degree. Notwithstanding the provisions of N.J.S.2C:1-8 or any other provision of law, a conviction under this subsection shall not merge with a conviction of any other offense, nor shall any other conviction merge with a conviction under this subsection. The court shall impose separate sentences upon a conviction under this subsection and any other criminal offense.

e. In addition to any other penalty provided by law, a person who is not authorized to obtain prescription monitoring information from the Prescription Monitoring Program who knowingly obtains or attempts to obtain such information in violation of the provisions of sections 25 through 30 of P.L.2007, c.244 (C.45:1-45 through C.45:1-50), shall be guilty of a crime of the fourth degree.

f. A civil penalty imposed under this section shall be collected by the director pursuant to the "Penalty Enforcement Law of 1999," P.L.1999, c.274 (C.2A:58-10 et seq.).

L.2007, c.244, s.29; amended 2015, c.74, s.6.

45:1-50 Authority to contract.

30. Authority to contract. The division may contract with one or more vendors to establish and maintain the Prescription Monitoring Program pursuant to guidelines established by the director.

L.2007, c.244, s.30.

45:1-50.1 Annual report.

9. The division shall annually submit a report to the Legislature, pursuant to section 2 of P.L.1991, c.164 (C.52:14-19.1), which provides information on the nature and extent of registration with, and utilization of, the Prescription Monitoring Program, as well as recommendations for program improvement.

L.2015, c.74, s.9.

45:1-50.2 Completion of assessment.

10. The division shall complete an assessment regarding the design, implementation requirements, and costs associated with a real time prescription monitoring system, and shall report its assessment and any recommendations to the Legislature, pursuant to section 2 of P.L.1991, c.164 (C.52:14-19.1), within 18 months after the enactment of P.L.2015, c.74 (C.45:1-46.1 et al.).

L.2015, c.74, s.10.