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1	HOUSE BILL NO. 923
2 3	Offered January 8, 2014
3	Prefiled January 8, 2014
4	A BILL to amend and reenact § 54.1-2523 of the Code of Virginia, relating to the Prescription
5	Monitoring Program; method of disclosure of information to recipient.
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_	Patron—Peace
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8	Referred to Committee on Health, Welfare and Institutions
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10	Be it enacted by the General Assembly of Virginia:
11 12	1. That § 54.1-2523 of the Code of Virginia is amended and reenacted as follows: § 54.1-2523. Confidentiality of data; disclosure of information; discretionary authority of
12	Director.
13 14	A. All data, records, and reports relating to the prescribing and dispensing of covered substances to
15	recipients and any abstracts from such data, records, and reports that are in the possession of the
16	Prescription Monitoring Program pursuant to this chapter and any material relating to the operation or
17	security of the program shall be confidential and shall be exempt from the Virginia Freedom of
18	Information Act (§ 2.2-3700 et seq.) pursuant to subdivision 15 of § 2.2-3705.5. Further, the Director
19	shall only have discretion to disclose any such information as provided in subsections B and C.
20	B. Upon receiving a request for information in accordance with the Department's regulations and in
21	compliance with applicable federal law and regulations, the Director shall disclose the following:
22	1. Information relevant to a specific investigation of a specific recipient or of a specific dispenser or
23	prescriber to an agent who has completed the Virginia State Police Drug Diversion School designated by
24	the superintendent of the Department of State Police or designated by the chief law-enforcement officer
25	of any county, city, or town or campus police department to conduct drug diversion investigations
26	pursuant to § 54.1-3405.
27	2. Information relevant to an investigation or inspection of or allegation of misconduct by a specific
28	person licensed, certified, or registered by or an applicant for licensure, certification, or registration by a
29	health regulatory board; information relevant to a disciplinary proceeding before a health regulatory
30 31	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'
31 32	Monitoring Program pursuant to Chapter 25.1 (§ 54.1-2515 et seq.).
33	3. Information relevant to the proceedings of any investigatory grand jury or special grand jury that
34	has been properly impaneled in accordance with the provisions of Chapter 13 (§ 19.2-191 et seq.) of
35	Title 19.2.
36	4. Information relevant to a specific investigation of a specific recipient, dispenser, or prescriber to
37	an agent of a federal law-enforcement agency with authority to conduct drug diversion investigations.
38	C. In accordance with the Department's regulations and applicable federal law and regulations, the
39	Director may, in his discretion, disclose:
40	1. Information in the possession of the program concerning a recipient who is over the age of 18 to
41	that recipient. The information shall be mailed to the street or mailing address indicated on the recipient
42	request form.
43	2. Information on a specific recipient to a prescriber, as defined in this chapter, for the purpose of
44 45	establishing the treatment history of the specific recipient when such recipient is either under care and
45 46	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
47	requested by the prescriber from the Prescription Monitoring Program.
48	3. Information on a specific recipient to a dispenser for the purpose of establishing a prescription
49	history to assist the dispenser in determining the validity of a prescription in accordance with
50	§ 54.1-3303 when the recipient is seeking a covered substance from the dispenser or the facility in
51	which the dispenser practices. In a manner specified by the Director in regulation, notice shall be given
52	to patients that information may be requested by the dispenser from the Prescription Monitoring
53	Program.
54	4. Information relevant to an investigation or regulatory proceeding of a specific dispenser or
55	prescriber to other regulatory authorities concerned with granting, limiting or denying licenses,
56	certificates or registrations to practice a health profession when such regulatory authority licenses such
57	dispenser or prescriber or such dispenser or prescriber is seeking licensure by such other regulatory
58	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

64 appropriate.

65 6. Information relevant to determination of the cause of death of a specific recipient to the designated 66 employees of the Office of the Chief Medical Examiner.

67 7. Information for the purpose of bona fide research or education to qualified personnel; however,
68 data elements that would reasonably identify a specific recipient, prescriber, or dispenser shall be deleted
69 or redacted from such information prior to disclosure. Further, release of the information shall only be
70 made pursuant to a written agreement between such qualified personnel and the Director in order to
71 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.

D. The Director may enter into agreements for mutual exchange of information among prescription
 monitoring programs in other jurisdictions, which shall only use the information for purposes allowed by
 this chapter.

E. This section shall not be construed to supersede the provisions of § 54.1-3406 concerning the divulging of confidential records relating to investigative information.

F. Confidential information that has been received, maintained or developed by any board or disclosed by the board pursuant to subsection A shall not, under any circumstances, 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. However, this subsection shall not be construed to inhibit any investigation or prosecution conducted pursuant to Article 1 (§ 18.2-247)

84 et seq.) of Chapter 7 of Title 18.2.