

# HOUSE BILL 1296

J3, J1, J2

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By: **Delegates Kach and Olszewski**

Introduced and read first time: February 7, 2014

Assigned to: Health and Government Operations

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## A BILL ENTITLED

1 AN ACT concerning

2 **Prescription Drug Monitoring Program – Review and Reporting of Possible**  
3 **Misuse or Abuse of Monitored Prescription Drugs**

4 FOR the purpose of authorizing the Prescription Drug Monitoring Program, in  
5 accordance with certain regulations, to review prescription monitoring data for  
6 a certain purpose and, under certain circumstances, report possible misuse or  
7 abuse of a monitored prescription drug to a certain prescriber or dispenser;  
8 requiring the Program, before reporting the possible misuse or abuse of a  
9 monitored prescription drug, to obtain from the technical advisory committee to  
10 the Program certain clinical guidance and interpretation; requiring the  
11 Secretary of Health and Mental Hygiene to adopt regulations that specify the  
12 process for the Program’s review of prescription monitoring data and reporting  
13 of possible misuse or abuse of a monitored prescription drug; altering the  
14 purpose of the technical advisory committee; making a stylistic change; and  
15 generally relating to the Prescription Drug Monitoring Program and the review  
16 of prescription monitoring data and reporting of possible misuse or abuse of a  
17 monitored prescription drug.

18 BY repealing and reenacting, without amendments,  
19 Article – Health – General  
20 Section 21–2A–02(a)  
21 Annotated Code of Maryland  
22 (2009 Replacement Volume and 2013 Supplement)

23 BY repealing and reenacting, with amendments,  
24 Article – Health – General  
25 Section 21–2A–04, 21–2A–06, and 21–2A–07  
26 Annotated Code of Maryland  
27 (2009 Replacement Volume and 2013 Supplement)

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EXPLANATION: CAPITALS INDICATE MATTER ADDED TO EXISTING LAW.

[Brackets] indicate matter deleted from existing law.



1 SECTION 1. BE IT ENACTED BY THE GENERAL ASSEMBLY OF  
2 MARYLAND, That the Laws of Maryland read as follows:

3 **Article – Health – General**

4 21–2A–02.

5 (a) There is a Prescription Drug Monitoring Program in the Department.

6 21–2A–04.

7 (a) The Secretary, in consultation with the Board, shall adopt regulations to  
8 carry out this subtitle.

9 (b) The regulations adopted by the Secretary shall:

10 (1) Specify the prescription monitoring data required to be submitted  
11 under § 21–2A–03 of this subtitle;

12 (2) Specify the electronic or other means by which information is to be  
13 submitted:

14 (i) Without unduly increasing the workload and expense on  
15 dispensers; and

16 (ii) In a manner as compatible as possible with existing data  
17 submission practices of dispensers;

18 (3) Specify that the Program:

19 (i) Shall provide the information technology software to  
20 dispensers necessary to upload prescription drug monitoring data to the Program; and

21 (ii) May not impose any fees or other assessments on prescribers  
22 or dispensers to support the operation of the Program;

23 (4) Specify that a prescriber or dispenser is not required or obligated  
24 to access or use prescription monitoring data available under the Program;

25 (5) Identify the mechanism by which prescription monitoring data are  
26 disclosed to a person, in accordance with § 21–2A–06 of this subtitle;

27 (6) Identify the circumstances under which a person may disclose  
28 prescription monitoring data received under the Program;

29 **(7) SPECIFY THE PROCESS FOR THE PROGRAM’S REVIEW OF**  
30 **PRESCRIPTION MONITORING DATA AND REPORTING OF POSSIBLE MISUSE OR**

1 ABUSE OF A MONITORED PRESCRIPTION DRUG UNDER § 21-2A-06(C) OF THIS  
2 SUBTITLE;

3           ~~[(7)] (8)~~ Establish requirements for Program retention of  
4 prescription monitoring data for 3 years; and

5           ~~[(8)] (9)~~ Require that:

6                   (i) Confidential or privileged patient information be kept  
7 confidential; and

8                   (ii) Records or information protected by a privilege between a  
9 health care provider and a patient, or otherwise required by law to be held  
10 confidential, be filed in a manner that, except as otherwise provided in § 21-2A-06 of  
11 this subtitle, does not disclose the identity of the person protected.

12 21-2A-06.

13           (a) Prescription monitoring data:

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

16                   (2) Are not public records; and

17                   (3) Except as provided in subsections ~~[(b) and (d)] (B), (C), AND (E)~~ of  
18 this section or as otherwise provided by law, may not be disclosed to any person.

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

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

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

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

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

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

1                   (6)    A patient with respect to prescription monitoring data about the  
2 patient;

3                   (7)    Subject to subsection [(g)] **(H)** of this section, the authorized  
4 administrator of another state's prescription drug monitoring program;

5                   (8)    The following units of the Department, on approval of the  
6 Secretary, for the purpose of furthering an existing bona fide individual investigation:

7                   (i)     The Office of the Chief Medical Examiner;

8                   (ii)    The Maryland Medical Assistance Program;

9                   (iii)   The Office of the Inspector General;

10                  (iv)    The Office of Health Care Quality; and

11                  (v)     The Division of Drug Control; or

12                  (9)    The technical advisory committee established under § 21-2A-07 of  
13 this subtitle for the purposes set forth in [subsection (c)] **SUBSECTIONS (C) AND (D)**  
14 of this section.

15                  **(C) (1) IN ACCORDANCE WITH REGULATIONS ADOPTED BY THE**  
16 **SECRETARY:**

17                               **(i) THE PROGRAM MAY REVIEW PRESCRIPTION**  
18 **MONITORING DATA FOR INDICATIONS OF POSSIBLE MISUSE OR ABUSE OF A**  
19 **MONITORED PRESCRIPTION DRUG; AND**

20                               **(ii) IF THE PROGRAM'S REVIEW OF PRESCRIPTION**  
21 **MONITORING DATA INDICATES POSSIBLE MISUSE OR ABUSE OF A MONITORED**  
22 **PRESCRIPTION DRUG, THE PROGRAM MAY REPORT THE POSSIBLE MISUSE OR**  
23 **ABUSE TO THE PRESCRIBER OR DISPENSER OF THE MONITORED PRESCRIPTION**  
24 **DRUG.**

25                               **(2) BEFORE THE PROGRAM REPORTS THE POSSIBLE MISUSE OR**  
26 **ABUSE OF A MONITORED PRESCRIPTION DRUG TO A PRESCRIBER OR DISPENSER**  
27 **UNDER THIS SUBSECTION, THE PROGRAM SHALL OBTAIN FROM THE TECHNICAL**  
28 **ADVISORY COMMITTEE:**

29                               **(i) CLINICAL GUIDANCE REGARDING INDICATIONS OF**  
30 **POSSIBLE MISUSE OR ABUSE; AND**

1                   **(II) INTERPRETATION OF THE PRESCRIPTION MONITORING**  
2 **DATA THAT INDICATES POSSIBLE MISUSE OR ABUSE.**

3           **[(c)] (D)**     Before the Program discloses information under subsection (b)(3),  
4 (4), (5), (7), or (8) of this section, the technical advisory committee [to the Program]  
5 shall:

6                   (1)     Review the requests for information;

7                   (2)     Provide clinical guidance and interpretation of the information  
8 requested to the Secretary to assist in the Secretary's decision on how to respond to a  
9 judicial subpoena, administrative subpoena, or other request; and

10                  (3)     Provide clinical guidance and interpretation of the information  
11 requested to the authorized recipient of the information.

12           **[(d)] (E)**     Except as provided by regulations adopted by the Secretary, a  
13 person who receives prescription monitoring data from the Program may not disclose  
14 the data.

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

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

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

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

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

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

31           **[(h)] (I)**     The Program may:

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

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

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

10 [(j)] (K) Prescription monitoring data may not be used as the basis for  
11 imposing clinical practice standards.

12 21-2A-07.

13 (a) There is a technical advisory committee to the Program.

14 (b) The purpose of the technical advisory committee is to [review]:

15 (1) REVIEW requests for information from the Program under §  
16 21-2A-06(b)(3), (4), (5), (7), and (8) of this subtitle; AND

17 (2) PROVIDE CLINICAL GUIDANCE AND INTERPRETATION TO THE  
18 PROGRAM REGARDING INDICATIONS OF POSSIBLE MISUSE OR ABUSE OF A  
19 MONITORED PRESCRIPTION DRUG UNDER § 21-2A-06(C)(3) OF THIS SUBTITLE.

20 (c) The technical advisory committee consists of the following members,  
21 appointed by the Secretary:

22 (1) A board certified anesthesiologist licensed and practicing in the  
23 State, nominated by the Maryland Society of Anesthesiologists;

24 (2) A certified addiction medicine specialist licensed and practicing in  
25 the State, nominated by the Maryland Society for Addiction Medicine;

26 (3) A pharmacist licensed and practicing in the State;

27 (4) A medical professional, licensed and practicing in the State, who is  
28 treating cancer patients; and

29 (5) A board certified physician specializing in the treatment of  
30 patients with pain, licensed and practicing in the State, nominated by the Maryland  
31 Society of Physical Medicine and Rehabilitation.

1           SECTION 2. AND BE IT FURTHER ENACTED, That this Act shall take effect  
2    October 1, 2014.