



Sen. Melinda Bush

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1 AMENDMENT TO SENATE BILL 3024

2 AMENDMENT NO. \_\_\_\_\_. Amend Senate Bill 3024 by replacing  
3 everything after the enacting clause with the following:

4 "Section 5. The Illinois Controlled Substances Act is  
5 amended by changing Section 318 as follows:

6 (720 ILCS 570/318)

7 Sec. 318. Confidentiality of information.

8 (a) Information received by the central repository under  
9 Section 316 and former Section 321 is confidential.

10 (a-1) To ensure the federal Health Insurance Portability  
11 and Accountability Act and confidentiality of substance use  
12 disorder patient records rules that mandate the privacy of an  
13 individual's prescription data reported to the Prescription  
14 Monitoring Program received from a retail dispenser under this  
15 Act, and in order to execute the duties and responsibilities  
16 under Section 316 of this Act and rules for disclosure under

1 this Section, the Clinical Director of the Prescription  
2 Monitoring Program or his or her designee shall maintain  
3 direct access to all Prescription Monitoring Program data. Any  
4 request for Prescription Monitoring Program data from any  
5 other department or agency must be approved in writing by the  
6 Clinical Director of the Prescription Monitoring Program or  
7 his or her designee unless otherwise permitted by law.  
8 Prescription Monitoring Program data shall only be disclosed  
9 as permitted by law.

10 (a-2) As an active step to address the current opioid  
11 crisis in this State and to prevent and reduce addiction  
12 resulting from a sports injury or an accident, the  
13 Prescription Monitoring Program and the Department of Public  
14 Health shall coordinate a continuous review of the  
15 Prescription Monitoring Program and the Department of Public  
16 Health data to determine if a patient may be at risk of opioid  
17 addiction. Each patient discharged from any medical facility  
18 with an International Classification of Disease, 10th edition  
19 code related to a sport or accident injury shall be subject to  
20 the data review. If the discharged patient is dispensed a  
21 controlled substance, the Prescription Monitoring Program  
22 shall alert the patient's prescriber as to the addiction risk  
23 and urge each to follow the Centers for Disease Control and  
24 Prevention guidelines or his or her respective profession's  
25 treatment guidelines related to the patient's injury. This  
26 subsection (a-2), other than this sentence, is inoperative on

1 or after January 1, 2024.

2 (b) The Department must carry out a program to protect the  
3 confidentiality of the information described in subsection  
4 (a). The Department may disclose the information to another  
5 person only under subsection (c), (d), or (f) and may charge a  
6 fee not to exceed the actual cost of furnishing the  
7 information.

8 (c) The Department may disclose confidential information  
9 described in subsection (a) to any person who is engaged in  
10 receiving, processing, or storing the information.

11 (d) The Department may release confidential information  
12 described in subsection (a) to the following persons:

13 (1) A governing body that licenses practitioners and  
14 is engaged in an investigation, an adjudication, or a  
15 prosecution of a violation under any State or federal law  
16 that involves a controlled substance.

17 (2) An investigator for the Consumer Protection  
18 Division of the office of the Attorney General, a  
19 prosecuting attorney, the Attorney General, a deputy  
20 Attorney General, or an investigator from the office of  
21 the Attorney General, who is engaged in any of the  
22 following activities involving controlled substances:

23 (A) an investigation;

24 (B) an adjudication; or

25 (C) a prosecution of a violation under any State  
26 or federal law that involves a controlled substance.

1 (3) A law enforcement officer who is:

2 (A) authorized by the Illinois State Police or the  
3 office of a county sheriff or State's Attorney or  
4 municipal police department of Illinois to receive  
5 information of the type requested for the purpose of  
6 investigations involving controlled substances; or

7 (B) approved by the Department to receive  
8 information of the type requested for the purpose of  
9 investigations involving controlled substances; and

10 (C) engaged in the investigation or prosecution of  
11 a violation under any State or federal law that  
12 involves a controlled substance.

13 (4) Select representatives of the Department of  
14 Children and Family Services through the indirect online  
15 request process. Access shall be established by an  
16 intergovernmental agreement between the Department of  
17 Children and Family Services and the Department of Human  
18 Services.

19 (e) Before the Department releases confidential  
20 information under subsection (d), the applicant must  
21 demonstrate in writing to the Department that:

22 (1) the applicant has reason to believe that a  
23 violation under any State or federal law that involves a  
24 controlled substance has occurred; and

25 (2) the requested information is reasonably related to  
26 the investigation, adjudication, or prosecution of the

1 violation described in subdivision (1).

2 (f) The Department may receive and release prescription  
3 record information under Section 316 and former Section 321  
4 to:

5 (1) a governing body that licenses practitioners;

6 (2) an investigator for the Consumer Protection  
7 Division of the office of the Attorney General, a  
8 prosecuting attorney, the Attorney General, a deputy  
9 Attorney General, or an investigator from the office of  
10 the Attorney General;

11 (3) any Illinois law enforcement officer who is:

12 (A) authorized to receive the type of information  
13 released; and

14 (B) approved by the Department to receive the type  
15 of information released; or

16 (4) prescription monitoring entities in other states  
17 per the provisions outlined in subsection (g) and (h)  
18 below;

19 confidential prescription record information collected under  
20 Sections 316 and 321 (now repealed) that identifies vendors or  
21 practitioners, or both, who are prescribing or dispensing  
22 large quantities of Schedule II, III, IV, or V controlled  
23 substances outside the scope of their practice, pharmacy, or  
24 business, as determined by the Advisory Committee created by  
25 Section 320.

26 (f-5) In accordance with a confidentiality agreement

1 entered into with the Department, a medical director, or a  
2 public health administrator and their delegated analysts, of a  
3 county or municipal health department or the Department of  
4 Public Health shall have access to data from the system for any  
5 of the following purposes:

6 (1) developing education programs or public health  
7 interventions relating to prescribing trends and  
8 controlled substance use; or

9 (2) conducting analyses and publish reports on  
10 prescribing trends in their respective jurisdictions.

11 At a minimum, the confidentiality agreement entered into  
12 with the Department shall:

13 (i) prohibit analysis and reports produced under  
14 subparagraph (2) from including information that  
15 identifies, by name, license, or address, any  
16 practitioner, dispenser, ultimate user, or other person  
17 administering a controlled substance; and

18 (ii) specify the appropriate technical and physical  
19 safeguards that the county or municipal health department  
20 must implement to ensure the privacy and security of data  
21 obtained from the system. The data from the system shall  
22 not be admissible as evidence, nor discoverable in any  
23 action of any kind in any court or before any tribunal,  
24 board, agency, or person. The disclosure of any such  
25 information or data, whether proper or improper, shall not  
26 wave or have any effect upon its confidentiality,

1           non-discoverability, or non-admissibility.

2           (g) The information described in subsection (f) may not be  
3 released until it has been reviewed by an employee of the  
4 Department who is licensed as a prescriber or a dispenser and  
5 until that employee has certified that further investigation  
6 is warranted. However, failure to comply with this subsection  
7 (g) does not invalidate the use of any evidence that is  
8 otherwise admissible in a proceeding described in subsection  
9 (h).

10           (h) An investigator or a law enforcement officer receiving  
11 confidential information under subsection (c), (d), or (f) may  
12 disclose the information to a law enforcement officer or an  
13 attorney for the office of the Attorney General for use as  
14 evidence in the following:

15           (1) A proceeding under any State or federal law that  
16 involves a controlled substance.

17           (2) A criminal proceeding or a proceeding in juvenile  
18 court that involves a controlled substance.

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

24           (j) Based upon federal, initial and maintenance funding, a  
25 prescriber and dispenser inquiry system shall be developed to  
26 assist the health care community in its goal of effective

1 clinical practice and to prevent patients from diverting or  
2 abusing medications.

3 (1) An inquirer shall have read-only access to a  
4 stand-alone database which shall contain records for the  
5 previous 12 months.

6 (2) Dispensers may, upon positive and secure  
7 identification, make an inquiry on a patient or customer  
8 solely for a medical purpose as delineated within the  
9 federal HIPAA law.

10 (3) The Department shall provide a one-to-one secure  
11 link and encrypted software necessary to establish the  
12 link between an inquirer and the Department. Technical  
13 assistance shall also be provided.

14 (4) Written inquiries are acceptable but must include  
15 the fee and the requester's ~~requestor's~~ Drug Enforcement  
16 Administration license number and submitted upon the  
17 requester's ~~requestor's~~ business stationery.

18 (5) As directed by the Prescription Monitoring Program  
19 Advisory Committee and the Clinical Director for the  
20 Prescription Monitoring Program, aggregate data that does  
21 not indicate any prescriber, practitioner, dispenser, or  
22 patient may be used for clinical studies.

23 (6) Tracking analysis shall be established and used  
24 per administrative rule.

25 (7) Nothing in this Act or Illinois law shall be  
26 construed to require a prescriber or dispenser to make use



1 of this inquiry system.

2 (8) If there is an adverse outcome because of a  
3 prescriber or dispenser making an inquiry, which is  
4 initiated in good faith, the prescriber or dispenser shall  
5 be held harmless from any civil liability.

6 (k) The Department shall establish, by rule, the process  
7 by which to evaluate possible erroneous association of  
8 prescriptions to any licensed prescriber or end user of the  
9 Illinois Prescription Information Library (PIL).

10 (l) The Prescription Monitoring Program Advisory Committee  
11 is authorized to evaluate the need for and method of  
12 establishing a patient specific identifier.

13 (m) Patients who identify prescriptions attributed to them  
14 that were not obtained by them shall be given access to their  
15 personal prescription history pursuant to the validation  
16 process as set forth by administrative rule.

17 (n) The Prescription Monitoring Program is authorized to  
18 develop operational push reports to entities with compatible  
19 electronic medical records. The process shall be covered  
20 within administrative rule established by the Department.

21 (o) Hospital emergency departments and freestanding  
22 healthcare facilities providing healthcare to walk-in patients  
23 may obtain, for the purpose of improving patient care, a  
24 unique identifier for each shift to utilize the PIL system.

25 (p) The Prescription Monitoring Program shall  
26 automatically create a log-in to the inquiry system when a

1 prescriber or dispenser obtains or renews his or her  
2 controlled substance license. The Department of Financial and  
3 Professional Regulation must provide the Prescription  
4 Monitoring Program with electronic access to the license  
5 information of a prescriber or dispenser to facilitate the  
6 creation of this profile. The Prescription Monitoring Program  
7 shall send the prescriber or dispenser information regarding  
8 the inquiry system, including instructions on how to log into  
9 the system, instructions on how to use the system to promote  
10 effective clinical practice, and opportunities for continuing  
11 education for the prescribing of controlled substances. The  
12 Prescription Monitoring Program shall also send to all  
13 enrolled prescribers, dispensers, and designees information  
14 regarding the unsolicited reports produced pursuant to Section  
15 314.5 of this Act.

16 (q) A prescriber or dispenser may authorize a designee to  
17 consult the inquiry system established by the Department under  
18 this subsection on his or her behalf, provided that all the  
19 following conditions are met:

20 (1) the designee so authorized is employed by the same  
21 hospital or health care system; is employed by the same  
22 professional practice; or is under contract with such  
23 practice, hospital, or health care system;

24 (2) the prescriber or dispenser takes reasonable steps  
25 to ensure that such designee is sufficiently competent in  
26 the use of the inquiry system;

1           (3) the prescriber or dispenser remains responsible  
2 for ensuring that access to the inquiry system by the  
3 designee is limited to authorized purposes and occurs in a  
4 manner that protects the confidentiality of the  
5 information obtained from the inquiry system, and remains  
6 responsible for any breach of confidentiality; and

7           (4) the ultimate decision as to whether or not to  
8 prescribe or dispense a controlled substance remains with  
9 the prescriber or dispenser.

10          The Prescription Monitoring Program shall send to  
11 registered designees information regarding the inquiry system,  
12 including instructions on how to log onto the system.

13          (r) The Prescription Monitoring Program shall maintain an  
14 Internet website in conjunction with its prescriber and  
15 dispenser inquiry system. This website shall include, at a  
16 minimum, the following information:

17           (1) current clinical guidelines developed by health  
18 care professional organizations on the prescribing of  
19 opioids or other controlled substances as determined by  
20 the Advisory Committee;

21           (2) accredited continuing education programs related  
22 to prescribing of controlled substances;

23           (3) programs or information developed by health care  
24 professionals that may be used to assess patients or help  
25 ensure compliance with prescriptions;

26           (4) updates from the Food and Drug Administration, the

1 Centers for Disease Control and Prevention, and other  
2 public and private organizations which are relevant to  
3 prescribing;

4 (5) relevant medical studies related to prescribing;

5 (6) other information regarding the prescription of  
6 controlled substances; and

7 (7) information regarding prescription drug disposal  
8 events, including take-back programs or other disposal  
9 options or events.

10 The content of the Internet website shall be periodically  
11 reviewed by the Prescription Monitoring Program Advisory  
12 Committee as set forth in Section 320 and updated in  
13 accordance with the recommendation of the advisory committee.

14 (s) The Prescription Monitoring Program shall regularly  
15 send electronic updates to the registered users of the  
16 Program. The Prescription Monitoring Program Advisory  
17 Committee shall review any communications sent to registered  
18 users and also make recommendations for communications as set  
19 forth in Section 320. These updates shall include the  
20 following information:

21 (1) opportunities for accredited continuing education  
22 programs related to prescribing of controlled substances;

23 (2) current clinical guidelines developed by health  
24 care professional organizations on the prescribing of  
25 opioids or other drugs as determined by the Advisory  
26 Committee;

1 (3) programs or information developed by health care  
2 professionals that may be used to assess patients or help  
3 ensure compliance with prescriptions;

4 (4) updates from the Food and Drug Administration, the  
5 Centers for Disease Control and Prevention, and other  
6 public and private organizations which are relevant to  
7 prescribing;

8 (5) relevant medical studies related to prescribing;

9 (6) other information regarding prescribing of  
10 controlled substances;

11 (7) information regarding prescription drug disposal  
12 events, including take-back programs or other disposal  
13 options or events; and

14 (8) reminders that the Prescription Monitoring Program  
15 is a useful clinical tool.

16 (t) Notwithstanding any other provision of this Act,  
17 neither the Prescription Monitoring Program nor any other  
18 person shall disclose any information in violation of the  
19 restrictions and requirements of paragraph (3.5) of subsection  
20 (a) of Section 316 as implemented under Public Act 102-527.

21 (Source: P.A. 99-480, eff. 9-9-15; 100-125, eff. 1-1-18;  
22 100-1093, eff. 8-26-18.)".