1	STATE OF OKLAHOMA
2	2nd Session of the 58th Legislature (2022)
3	COMMITTEE SUBSTITUTE FOR
4	SENATE BILL 1151 By: Standridge
5	
6	
7	COMMITTEE SUBSTITUTE
8	An Act relating to the Anti-Drug Diversion Act; amending 63 O.S. 2021, Sections 2-309B and 2-309D,
9	which relate to definitions and central repository information; modifying definition; prohibiting and
10	allowing certain disclosures; providing for confidentiality of certain records; updating
11	statutory language; and declaring an emergency.
12	
13	
14	BE IT ENACTED BY THE PEOPLE OF THE STATE OF OKLAHOMA:
15	SECTION 1. AMENDATORY 63 O.S. 2021, Section 2-309B, is
16	amended to read as follows:
17	Section 2-309B. For the purposes of the Anti-Drug Diversion
18	Act:
19	1. "Bureau" means the Oklahoma State Bureau of Narcotics and
20	Dangerous Drugs Control;
21	2. "Dispenser" means a person who distributes a Schedule II
22	controlled dangerous substance, but does not include a licensed
23	hospital pharmacy or a licensed nurse or medication aide who
24	

Req. No. 3498

1 administers such a substance at the direction of a licensed
2 physician;

3 3. "Dispenser's registration number" means the dispenser's
4 Oklahoma State Bureau of Narcotics and Dangerous Drugs Control
5 registration number or, in the case of a pharmacist, the National
6 Association of Boards of Pharmacy number for the pharmacy where the
7 dispensation is made;

8 4. "Exception report" means an output of data indicating
9 Schedule II controlled dangerous substance dispensation which is
10 outside expected norms for a prescriber practicing a particular
11 specialty or field of health care, for a dispenser doing business in
12 a particular location, or for a recipient;

13 5. "Recipient" means the person for whom a prescription is14 prescribed and who is the lawful intended ultimate user;

6. "Recipient's agent" means a person who is authorized by the ultimate user to pick up the recipient's medication and deliver it to the recipient or a person who claims a prescription other than the person to whom the medication is prescribed;

19 7. "Recipient's identification number" and "recipient's agent's 20 identification number" means the unique number contained on a valid 21 passport, military identification card, driver license, or

22 identification card issued to a recipient pursuant to Section 6-105
23 of Title 47 of the Oklahoma Statutes or similar statute of another
24 state if the recipient is not a resident of the State of Oklahoma,

Req. No. 3498

1	or, if the recipient is less than eighteen (18) years old and has no
2	such identification, the unique number contained on a valid
3	passport, military identification card, driver license, or
4	identification card issued to the recipient's parent or guardian
5	pursuant to Section 6-105 of Title 47 of the Oklahoma Statutes or
6	similar statute of another state if the parent or guardian is not a
7	resident of the State of Oklahoma, or, if the controlled dangerous
8	substance is obtained for an animal, the unique number contained on
9	the animal owner's valid driver license or identification card
10	issued pursuant to Section 6-105 of Title 47 of the Oklahoma
11	Statutes or similar statute of another state if the owner is not a
12	resident of the State of Oklahoma. Nonresident drug outlets
13	registered pursuant to the Oklahoma Pharmacy Act and resident drug
14	outlets defined in Section 353.1 of Title 59 of the Oklahoma
15	Statutes are exempt from the picture identification requirement if
16	the nonresident and resident drug outlets have obtained the
17	identification of the patient through the prescription benefit plan
18	of the patient forms of identification listed in 8 CFR
19	274a.2(b)(1)(v)(A) and (B);
20	8. "Registrant" means a person, persons, corporation or other
21	entity who has been issued by the Director of the Oklahoma State
22	Bureau of Narcotics and Dangerous Drugs Control a registration
23	pursuant to Section 2-302 of this title; and
~ .	

24

Req. No. 3498

1 9. "State" means any state, territory, or possession of the United States, the District of Columbia, or foreign nation. 2 SECTION 2. AMENDATORY 63 O.S. 2021, Section 2-309D, is 3 amended to read as follows: 4 5 Section 2-309D. A. The information collected at the central repository pursuant to the Anti-Drug Diversion Act shall be 6 confidential and shall not be open to the public. Access to the 7 information shall be limited to: 8 9 1. Peace officers certified pursuant to Section 3311 of Title 10 70 of the Oklahoma Statutes who are employed as investigative agents of the Oklahoma State Bureau of Narcotics and Dangerous Drugs 11 12 Control: 2. The United States Drug Enforcement Administration Diversion 13 Group Supervisor; 14 The executive director or chief investigator, as designated 15 3. by each board, of the following state boards: 16 Board of Podiatric Medical Examiners, 17 a. b. Board of Dentistry, 18 State Board of Pharmacy, 19 с. d. State Board of Medical Licensure and Supervision, 20 State Board of Osteopathic Examiners, 21 e. f. State Board of Veterinary Medical Examiners, 22 Oklahoma Health Care Authority, 23 g. 24

1 h. Department of Mental Health and Substance Abuse 2 Services, Board of Examiners in Optometry, 3 i. j. Oklahoma Board of Nursing, 4 5 k. Office of the Chief Medical Examiner, and State Board of Health; 6 1. A multicounty grand jury properly convened pursuant to the 7 4. Multicounty Grand Jury Act; 8 9 5. Medical practitioners employed by the United States Department of Veterans Affairs, the United States Military, or other 10 federal agencies treating patients in this state; 11 6. At the discretion of the Director of the Oklahoma State 12 Bureau of Narcotics and Dangerous Drugs Control, medical 13 practitioners and their staff $_{\tau}$ including those employed by the 14 federal government in this state; and 15 7. The members of the Opioid Overdose Fatality Review Board for 16 the purpose of carrying out the duties prescribed by Section 2-1001 17 of this title. 18 This section shall not prevent access, at the discretion of 19 в. the Director of the Oklahoma State Bureau of Narcotics and Dangerous 20 Drugs Control, to investigative information by peace officers and 21 investigative agents of federal, state, tribal, county or municipal 22 law enforcement agencies, district attorneys and the Attorney 23 General in furtherance of criminal, civil or administrative 24

investigations or prosecutions within their respective
 jurisdictions, designated legal, communications, and analytical
 employees of the Bureau, and to registrants in furtherance of
 efforts to guard against the diversion of controlled dangerous
 substances.

С. This section shall not prevent the disclosure, at the 6 discretion of the Director of the Oklahoma State Bureau of Narcotics 7 and Dangerous Drugs Control, of statistical information gathered 8 9 from the central repository to the general public which shall be 10 limited to types and quantities of controlled substances dispensed 11 and the county where dispensed for statistical, research, substance abuse prevention, or educational purposes, provided that consumer 12 13 confidentiality is not compromised.

D. 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 prescription-monitoring-program
information to prescription-monitoring programs of other states
provided a reciprocal data-sharing agreement is in place.

E. The Department of Mental Health and Substance Abuse Services
and the State Department of Health may utilize the information in
the central repository for statistical, research, substance abuse
prevention, or educational purposes, provided that consumer
confidentiality is not compromised.

24

F. Any unauthorized disclosure of any information collected at the central repository provided by the Anti-Drug Diversion Act shall be a misdemeanor. Violation of the provisions of this section shall be deemed willful neglect of duty and shall be grounds for removal from office.

1. Registrants shall have access to the central repository 6 G. for the purposes of patient treatment and to aid in the 7 determination in prescribing or screening new patients. 8 The 9 physician or designee shall provide, upon request by the patient, the history of the patient or the query history of the patient. 10 2. Prior to prescribing or authorizing for refill, if one 11 a. hundred eighty (180) days have elapsed prior to the 12 previous access and check, of opiates, synthetic 13 opiates, semisynthetic opiates, benzodiazepine or 14 carisoprodol to a patient of record, registrants or 15 members of their medical or administrative staff shall 16 be required to access the information in the central 17 repository to assess medical necessity and the 18 possibility that the patient may be unlawfully 19 obtaining prescription drugs in violation of the 20 Uniform Controlled Dangerous Substances Act. The duty 21 to access and check shall not alter or otherwise amend 22 appropriate medical standards of care. The registrant 23 or medical provider shall note in the patient file 24

1	that the central repository has been checked and may
2	maintain a copy of the information.
3	b. The requirements set forth in subparagraph a of this
4	paragraph shall not apply:
5	(1) to medical practitioners who prescribe the
6	controlled substances set forth in subparagraph a
7	of this paragraph for hospice or end-of-life
8	care, or
9	(2) for a prescription of a controlled substance set
10	forth in subparagraph a of this paragraph that is
11	issued by a practitioner for a patient residing
12	in a nursing facility as defined by Section 1-
13	1902 of this title, provided that the
14	prescription is issued to a resident of such
15	facility.
16	3. Registrants shall not be liable to any person for any claim
17	of damages as a result of accessing or failing to access the
18	information in the central repository and no lawsuit may be
19	predicated thereon.
20	4. The failure of a registrant to access and check the central
21	repository as required under state or federal law or regulation may,
22	after investigation, be grounds for the licensing board of the
23	registrant to take disciplinary action against the registrant.

24

1 Η. The State Board of Podiatric Medical Examiners, the State 2 Board of Dentistry, the State Board of Medical Licensure and Supervision, the State Board of Examiners in Optometry, the State 3 Oklahoma Board of Nursing, the State Board of Osteopathic Examiners 4 5 and the State Board of Veterinary Medical Examiners shall have the sole responsibility for enforcement of the provisions of subsection 6 G of this section. Nothing in this section shall be construed so as 7 to permit the Director of the State Bureau of Narcotics and 8 9 Dangerous Drugs Control to assess administrative fines provided for in Section 2-304 of this title. 10

The Director of the Oklahoma State Bureau of Narcotics and 11 I. 12 Dangerous Drugs Control, or a designee thereof, shall provide a 13 monthly list to the Directors of the State Board of Podiatric Medical Examiners, the State Board of Dentistry, the State Board of 14 Medical Licensure and Supervision, the State Board of Examiners in 15 Optometry, the State Oklahoma Board of Nursing, the State Board of 16 17 Osteopathic Examiners and the State Board of Veterinary Medical Examiners of the top twenty prescribers of controlled dangerous 18 substances within their respective areas of jurisdiction. Upon 19 discovering that a registrant is prescribing outside the limitations 20 of his or her licensure or outside of drug registration rules or 21 applicable state laws, the respective licensing board shall be 22 notified by the Bureau in writing. Such notifications may be 23 considered complaints for the purpose of investigations or other 24

Req. No. 3498

actions by the respective licensing board. Licensing boards shall
 have exclusive jurisdiction to take action against a licensee for a
 violation of subsection G of this section.

Information regarding fatal and nonfatal overdoses, other 4 J. 5 than statistical information as required by Section 2-106 of this title, shall be completely confidential. Access to this information 6 shall be strictly limited to the Director of the Oklahoma State 7 Bureau of Narcotics and Dangerous Drugs Control or designee, the 8 9 Chief Medical Examiner, state agencies and boards provided in subsection A of this section, and the registrant that enters the 10 information. Registrants shall not be liable to any person for a 11 12 claim of damages for information reported pursuant to the provisions 13 of Section 2-105 of this title.

14 K. The Director of the Oklahoma State Bureau of Narcotics and
15 Dangerous Drugs Control shall provide adequate means and procedures
16 allowing access to central repository information for registrants
17 lacking direct computer access.

L. Upon completion of an investigation in which it is determined that a death was caused by an overdose, either intentionally or unintentionally, of a controlled dangerous substance, the medical examiner shall be required to report the decedent's name and date of birth to the Oklahoma State Bureau of Narcotics and Dangerous Drugs Control. The Oklahoma State Bureau of Narcotics and Dangerous Drugs Control shall be required to maintain

Req. No. 3498

a database containing the classification of medical practitioners
 who prescribed or authorized controlled dangerous substances
 pursuant to this subsection.

The Oklahoma State Bureau of Narcotics and Dangerous Drugs 4 Μ. 5 Control is authorized to provide unsolicited notification to the licensing board of a pharmacist or practitioner if a patient has 6 received one or more prescriptions for controlled substances in 7 quantities or with a frequency inconsistent with generally 8 9 recognized standards of safe practice. An unsolicited notification 10 to the licensing board of the practitioner pursuant to this section: 1. Is confidential; 11

May not disclose information that is confidential pursuant
 to this section; and

14 3. May be in a summary form sufficient to provide notice of the 15 basis for the unsolicited notification.

16 <u>N. Except as otherwise provided for in subsections A and B of</u> 17 <u>this section, any information collected at the central repository,</u> 18 as outlined in Section 2-309C of this title, shall:

- 19 1. Be confidential by law and privileged;
- 20 2. Not be subject to the Oklahoma Open Records Act;
- 21 3. Not be subject to subpoena; and
- 22 <u>4. Not be subject to discovery or admissible in evidence in any</u> 23 private civil action.
- 24

1	SECTION 3. It being immediately necessary for the preservation
2	of the public peace, health or safety, an emergency is hereby
3	declared to exist, by reason whereof this act shall take effect and
4	be in full force from and after its passage and approval.
5	
6	58-2-3498 JES 2/28/2022 3:03:50 PM
7	
8	
9	
10	
11	
12	
13	
14	
15	
16	
17	
18	
19	
20	
21	
22	
23	
24	