SUMMARY ANALYSIS

CS/CS/HB 557 passed the House on April 5, 2017. The bill was amended in the Senate on May 1, 2017, and was returned to the House on May 1, 2017. The House concurred in the Senate amendment as amended by the House on May 3, 2017. The Senate concurred in the House amendment to the Senate amendment, and passed the Senate as amended on May 5, 2017.

In 2009, the Legislature created the Prescription Drug Monitoring Program (PDMP) within the Department of Health (DOH). The PDMP employs a database to monitor the prescribing and dispensing of certain controlled substances. Dispensers of controlled substances listed in Schedule II, III, or IV must report certain information to the PDMP database, including the name of the prescriber, the date the prescription is filled and dispensed, and the name, address, and date of birth of the person to whom the controlled substance is dispensed. Currently, dispensers must report dispensing controlled substances to the database within seven days of dispensing the controlled substances via the internet or other DOH-approved format, such as on a disc or regular mail.

Beginning January 1, 2018, the bill reduces the amount of time a dispenser has to report the dispensing of a controlled substance to the PDMP database from seven days to the close of the next business day after the controlled substance is dispensed. The bill requires PDMP reporting to be completed via the DOH-approved electronic system, and eliminates DOH authority to approve other options for submission.

Dispensing and administering controlled substances are exempt from PDMP reporting in certain health care settings where the risk of controlled substances being overprescribed or diverted is low. These health care settings include a licensed hospital, nursing home, ambulatory surgical center, hospice, intermediate care facility for the developmentally disabled, rehabilitative hospital, and assisted living facility.

The bill requires the patient to be present and receiving care for the reporting exemption for a rehabilitation hospital, assisted living facility, or nursing home to apply.

The bill authorizes certain health care employees of the U.S. Veterans’ Administration to access the PDMP database in manner established by DOH. Such access is limited to the authorized employee’s review of his or her patient’s controlled substance prescription history.

The bill may have an insignificant, negative fiscal impact on DOH that can be absorbed with existing resources and has no fiscal impact on local governments.

Subject to the Governor’s veto powers, the effective date of this bill is July 1, 2017.
I. SUBSTANTIVE INFORMATION

A. EFFECT OF CHANGES:

Prescription Drug Monitoring Program

Prescription Drug Monitoring Programs (PDMPs) are state-run electronic databases used to track the prescribing and dispensing of certain controlled prescription drugs to patients. Prescription Drug Monitoring Programs (PDMPs) are designed to monitor this information for suspected abuse or diversion and provide prescribers and pharmacists with critical information regarding a patient’s controlled substance prescription history. As of September 2015, 49 states had an operational PDMP database.

Chapter 2009-197, Laws of Fla., established Florida’s PDMP within the Department of Health (DOH), and is codified in s. 893.055, F.S. The PDMP uses an electronic database system to monitor the prescribing and dispensing of certain controlled substances. The PDMP database became operational in September of 2011, when it began receiving prescription data from pharmacies and dispensing practitioners.

PDMP Reporting Requirements

Dispensers of controlled substances listed in Schedule II, III, or IV of the Florida Comprehensive Drug Abuse Prevention and Control Act must report specified information to the PDMP database:

- The name of the prescribing practitioner, the practitioner’s federal Drug Enforcement Administration (DEA) registration number, the practitioner’s National Provider Identification (NPI) or other appropriate identifier, and the date of the prescription;
- The date the prescription was filled and the method of payment, such as cash by an individual or third-party payment;
- The full name, address, and date of birth of the person for whom the prescription was written;
- The name, national drug code, quantity, and strength of the controlled substance dispensed;
- The full name, federal DEA registration number, and address of the pharmacy, other location, or other practitioner from which the controlled substance was dispensed;
- The name of the pharmacy or practitioner, other than a pharmacist, dispensing the controlled substance and the practitioner’s NPI; and
- Other appropriate identifying information as determined by DOH rule.

Dispensers must report dispensing a specified controlled substance to the PDMP database within seven days. As of June 30, 2016, approximately 96 percent of pharmacies required to report data to the PDMP had uploaded information into the system within the seven-day statutory limit. Of those, 66 percent reported the information within 24 hours. The time in which a dispenser must submit

2 Id.
4 Section 893.055(2)(a), F.S.
6 Section 893.055(3), F.S.; controlled substances listed in Schedule II, III, or IV can be found in s. 893.03(2)-(4), F.S.
7 Id.
8 Section 893.055(4), F.S.
9 Supra, FN 5.
10 Id.
information to the PDMP varies across the nation. As indicated below, some states require the 
dispenser to submit data within 24 hours, others (like Florida) allow up to 7 days, and two allow the 
dispenser up to a month to submit the data:¹¹

Prescription Drug Monitoring Programs (PDMP): 
Data Submission Interval and Mandatory Use Requirements

In Florida, more than 6,500 dispensers have reported to the PDMP creating the more than 198 million 
dispensing records that are maintained in the PDMP system.¹²

Exemptions from PDMP Reporting Requirements

The purpose of the PDMP is to track the dispensing of prescribed controlled substances to provide 
information to subsequent prescribing physicians and prevent the overprescribing and diversion of such 
substances. However, there are some circumstances in which there is inherently a low risk of controlled 
substances being overprescribed or diverted, and in those circumstances, the law exempts 
practitioners from having to report the dispensing of controlled substances. Specifically, the following 
acts are not required to be reported:

- A health care practitioner administering a controlled substance directly to a patient if the amount 
of the controlled substance is adequate to treat the patient during that particular treatment 
  session;
- A pharmacist or health care practitioner administering a controlled substance to a patient or 
  resident receiving care as a patient at a hospital, nursing home, ambulatory surgical center, 
  hospice, or intermediate care facility for the developmentally disabled which is licensed in this 
  state;
- A practitioner administering or dispensing a controlled substance in the health care system of 
  the Department of Corrections;

¹¹ National Conference of State Legislatures, “Prescription Drug Monitoring Programs,” (June 1, 2016), available at 
¹² Supra, FN 5.
• A practitioner administering a controlled Substance in the emergency room of a licensed hospital;
• A health care practitioner administering or dispensing a controlled Substance to a person under the age of 16;
• A pharmacist or a dispensing practitioner dispensing a one-time, 72-hour emergency resupply of a controlled Substance to a patient; and
• A rehabilitative hospital, assisted living facility, or nursing home dispensing a certain dosage of a controlled Substance, as needed, to a patient as ordered by the patient’s treating physician.13

Access to PDMP Data

Direct access to the PDMP database is presently limited by law to a pharmacy, prescriber, or dispenser.14 A pharmacy, prescriber, or dispenser has access to information in the PDMP database that relates to a patient of that pharmacy, prescriber, or dispenser, as needed, for reviewing the patient’s controlled Substance prescription history.15 Currently, the only prescribers authorized to access the PDMP database are Florida-licensed health care practitioners.16 Health care practitioners who work for the United States Department of Veterans Affairs (VA) in one of its facilities in Florida are not required to be licensed in Florida. The VA requires that a health care practitioner have an active, unrestricted license to practice from any state to practice at any one of its facilities nationwide.17 Therefore, a health care practitioner practicing in a VA facility in Florida who is licensed in another state would not have access to the PDMP database.

Health care practitioners began accessing the PDMP database on October 17, 2011.18 As of June 30 2016, 36,718 health care practitioners, or 23.7 percent of all licensed health care practitioners, were registered to use the PDMP Database.19 Pharmacists have had the highest utilization rate of the PDMP; from July 1, 2015, to June 30, 2016, 54.5 percent of pharmacists were registered to use the PDMP and 90.1 percent of pharmacists registered to use the PDMP had queried it.20 From July 1, 2015, to June 30, 2016, in-state prescribers issued 37,048,030 controlled Substance prescriptions to 7,387,884 Florida residents.21 During that same timeframe, 28,984 registered health care practitioners queried the PDMP database 27,501,266 times.22

In Florida, indirect access to the PDMP database is provided to:

• DOH and its relevant health care regulatory boards;
• The Attorney General for Medicaid fraud cases involving prescribed controlled substances;
• A law enforcement agency during active investigations regarding potential criminal activity, fraud, or theft regarding prescribed controlled substances; and
• A patient or the legal guardian or designated health care surrogate of an incapacitated patient, for verifying the accuracy of database information.23

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13 Section 893.055(5). F.S.
14 Section 893.055(7)(b), F.S.
15 Id.
19 Supra note 5 at p. 10.
20 Id. at p. 10, 18.
21 Id. at p. 14.
22 Id. at p. 18.
23 Section 893.055(7)(c), F.S.
Entities with indirect access to the PDMP database may request information from the PDMP program manager that is otherwise confidential and exempt from public disclosure under s. 893.0551, F.S.\textsuperscript{24} Prior to release, the PDMP program manager must verify that the request is authentic and authorized with the requesting organization.\textsuperscript{25}

**Public Records Exemption for Information in the PDMP Database**

Section 893.0551, F.S.,\textsuperscript{26} provides that personal information of a patient and certain information concerning health care practitioners contained in the PDMP database are confidential and exempt from s. 119.07(1), F.S., and Art. I, Sec. 24 of the Florida Constitution.\textsuperscript{27} The statute makes confidential and exempt identifying information, including, but not limited to, the name, address, telephone number, insurance plan number, government-issued identification number, provider number, Drug Enforcement Administration number, or any other unique identifying number of a patient, patient’s agent, health care practitioner or practitioner as defined in s. 893.055, F.S., or an employee of the practitioner who is acting on behalf of and at the direction of the practitioner, a pharmacist, or a pharmacy, which is contained in the PDMP database.

Any agency or person that obtains information pursuant to s. 893.0551, F.S., must maintain the confidential and exempt status of that information.\textsuperscript{28}

**Effect of Proposed Changes**

Beginning January 1, 2018, CS/HB 557 reduces the amount of time a pharmacy or dispenser has to report the dispensing of a controlled substance to the PDMP database from seven days after the controlled substance is dispensed to no later than the end of the next business day after the controlled substance is dispensed.

The bill requires the controlled substance reporting by pharmacies or dispensers to be done via the department-approved electronic system, and eliminates the authority of the department to approve other methods of submission, such as submission by disc or by regular mail.

The bill clarifies that the exemption to the reporting required under this section provided to a rehabilitation hospital, assisted living facility, or nursing homes, applies only while the patient is present and receiving care as ordered by the patient’s treating physician.

The bill authorizes employees of the VA who provide health care services and have authority to prescribe controlled substances to access the PDMP database in a manner prescribed by DOH. The access is limited to information related to the patient of authorized VA employee and may only be accessed to review such patient's controlled substance prescription history.

The bill provides an effective date of July 1, 2017.

\textsuperscript{24} Id.
\textsuperscript{25} Id.
\textsuperscript{26} The public records exemption was established in 2009 in conjunction with the PDMP. See s. 1, ch. 2009-197, Laws of Fla. Additionally, the public records exemption was reauthorized in 2014. See .s 1 ch. 2014-156, Laws of Fla.
\textsuperscript{27} Section 893.0551(2), F.S.
\textsuperscript{28} Section 893.0551(6), F.S. However, a law enforcement agency with lawful access to such information is permitted to disclose confidential and exempt information received from DOH to a criminal justice agency as part of an active investigation of a specific violation of law. Section 893.0551(4).
II. FISCAL ANALYSIS & ECONOMIC IMPACT STATEMENT

A. FISCAL IMPACT ON STATE GOVERNMENT:

1. Revenues:
   None.

2. Expenditures:
   DOH may incur insignificant costs associated with rulemaking to amend current rules to align with the statutory changes proposed by the bill. Current budget authority is adequate to absorb such costs.\textsuperscript{29}

B. FISCAL IMPACT ON LOCAL GOVERNMENTS:

1. Revenues:
   None.

2. Expenditures:
   None.

C. DIRECT ECONOMIC IMPACT ON PRIVATE SECTOR:

A pharmacy or dispenser may incur additional costs associated with meeting the new requirement to report the dispensing of a controlled substance by the end of the next business day.

D. FISCAL COMMENTS:

None.

\textsuperscript{29} Department of Health, 2017 \textit{Agency Legislative Bill Analysis: House Bill 557}, January 27, 2017, (on file with the Health and Human Services Committee).