PROPOSED REGULATION OF THE

STATE BOARD OF PHARMACY

LCB File No. R047-15

September 15, 2015

EXPLANATION - Matter in *italics* is new; matter in brackets [omitted material] is material to be omitted.

AUTHORITY: §1, NRS 639.070.

A REGULATION relating to pharmacy; revising provisions relating to the transmission of information regarding the dispensing of controlled substances to certain persons; and providing other matters properly relating thereto.

Legislative Counsel's Digest:

Under existing regulations, the State Board of Pharmacy requires each pharmacy that uses a computerized system to record information concerning prescriptions and that dispenses to certain persons a controlled substance that is listed in schedule II, III or IV to transmit certain information concerning the controlled substance to the Board or its agent on a weekly basis. (NAC 639.926) Certain practitioners who dispense controlled substances are also subject to those requirements. (NAC 639.745) This regulation requires such a pharmacy or practitioner to transmit that information not later than the next business day after dispensing the controlled substance. This regulation also requires such a pharmacy or practitioner that does not dispense such a controlled substance to transmit to the Board or its agent a zero report stating that the pharmacy or practitioner did not dispense such a controlled substance on the immediately preceding business day. Finally, this regulation revises the methods that a pharmacy or practitioner is required to use to transmit the information or zero report.

Section 1. NAC 639.926 is hereby amended to read as follows:

639.926 1. Each pharmacy that uses a computerized system to record information concerning prescriptions and that dispenses a controlled substance that is listed in schedule II, III or IV to a person who is not an inpatient of a hospital, correctional institution or nursing facility shall transmit to the Board or its agent the following information, as applicable, set forth in the 2011 ASAP Version 4.2 Standard for Prescription Monitoring Programs published by the American Society for Automation in Pharmacy. The following Segments and the accompanying

Data Elements of the Implementation Guide for the 2011 ASAP Version 4.2 Standard for Prescription Monitoring Programs are hereby adopted by reference:

(a) The Segment entitled "TH Transaction Header" and the following Data Elements:
(1) Version/Release Number;
(2) Transaction Control Number;
(3) Transaction Type;
(4) Response ID;
(5) Creation Date;
(6) Creation Time;
(7) File Type; and
(8) Segment Terminator Character;
(b) The Segment entitled "IS Information Source" and the following Data Elements:
(1) Unique Information Source ID;
(2) Information Source Entity Name; and
(3) Message;
(c) The Segment entitled "PHA Pharmacy Header" and the following Data Elements:
(1) National Provider Identifier (NPI);
(2) DEA Number;
(3) Pharmacy or Dispensing Prescriber Name;
(4) Phone Number;
(5) Contact Name; and
(6) Chain Site ID;
(d) The Segment entitled "PAT Patient Information" and the following Data Elements

(1)	Last Name;
(2)	First Name;
(3)	Address Information - 1;
(4)	City Address;
(5)	State Address;
(6)	ZIP Code Address;
(7)	Phone Number;
(8)	Date of Birth; and
(9)	Gender Code;
(e) The	e Segment entitled "DSP Dispensing Record" and the following Data Elements:
(1)	Reporting Status;
(2)	Prescription Number;
(3)	Date Written;
(4)	Refills Authorized;
(5)	Date Filled;
(6)	Refill Number;
(7)	Product ID Qualifier;
(8)	Product ID;
(9)	Quantity Dispensed;
(10)	Days Supply;
(11)	Transmission Form of Rx Origin Code;
(12)	Classification Code for Payment Type; and
(13)	Date Sold;

(1) National Provider Identifier (NPI); (2) DEA Number; (3) DEA Number Suffix; (4) Last Name; (5) First Name; and (6) Phone Number; (g) The Segment entitled "CDI Compound Drug Ingredient Detail" and the following Data Elements: (1) Compound Drug Ingredient Sequence Number; (2) Product ID Qualifier; (3) Product ID; (4) Component Ingredient Quantity; and (5) Compound Drug Dosage Units Code; (h) The Segment entitled "TP Pharmacy Trailer" and the Data Element Detail Segment Count; and (i) The Segment entitled "TT Transaction Trailer" and the following Data Elements: (1) Transaction Control Number; and (2) Segment Count. 2. A copy of the publication may be obtained from the American Society for Automation in Pharmacy at the Internet address http://www.asapnet.org, or by telephone at (610) 825-7783,

(f) The Segment entitled "PRE Prescriber Information" and the following Data Elements:

for the price of \$175 for members and \$770 for nonmembers.

- 3. [The] A pharmacy that dispenses a controlled substance and is required to transmit information to the Board or its agent pursuant to subsection 1 shall transmit the information [required pursuant to this section] not later than [each Wednesday for the prescriptions filled from the immediately preceding Sunday through Saturday. If a Wednesday falls on a legal holiday, then the information must be reported on the next business day that is not a legal holiday.] the end of the next business day after dispensing the controlled substance. A pharmacy that does not dispense a controlled substance as specified in subsection 1 shall transmit to the Board or its agent a zero report stating that the pharmacy did not dispense such a controlled substance on the immediately preceding business day.
- 4. The information *required pursuant to this section or a zero report* must be transmitted by means of [a form of electronic data transmission approved by the Board, including, without limitation, a computer modem that can transmit information at the rate of 2400 baud or more.]:
 - (a) A secure file transfer protocol;
 - (b) An upload from an Internet web portal; or
 - (c) A manual entry.