420-7-2-.12 Prescription Drug Monitoring Program Reporting to Database by Dispensers

(1) Entities and practitioners that dispense controlled substances, Class II-V, shall report controlled substances prescription information to the Prescription Drug Monitoring Program database. These entities and practitioners include but are not limited to:

(a) Licensed pharmacies;

(b) Mail order pharmacies or pharmacy benefit programs filling prescriptions for or dispensing controlled substances to residents of Alabama; and

(c) Licensed physicians, dentists, podiatrists, optometrists, and veterinarians who dispense controlled substances.

(2) The reporting requirement in this rule does not apply to a controlled substance dispensed:

(a) By a pharmacy of a hospital, nursing home, or other inpatient health care facility if administered and used by a patient on the facility’s premises;

(b) By a practitioner if administered during the course of a patient’s treatment by injection, topical application, suppository administration, or oral administration; or

(c) By a practitioner as an appropriately labeled sample medication.

(3) Entities and practitioners shall submit reports as follows:

(a) Entities and nonveterinary practitioners shall submit reports at least once daily by 11:59 p.m.

1. If an entity or practitioner does not dispense a controlled substance on a specific day, the entity or practitioner shall report that zero controlled substances were dispensed.

2. The daily reporting requirement does not apply on days that the entity or practitioner’s business is closed and no controlled substances are dispensed.

(b) Veterinary practitioners shall submit reports at least once monthly by 11:59 p.m. on the last business day of the month. If a veterinary practitioner does not dispense a controlled substance in a specific month, the veterinary practitioner shall report that zero controlled substances were dispensed.

(c) Reports must be in electronic format according to American Society for Automation in Pharmacy Standards using the U.S. Postal Service’s Postal Addressing Standards.

1. If electronic transmission is not feasible, an entity or practitioner may request a waiver.
2. An entity or practitioner who receives a waiver may submit prescription information in an alternate format approved by the Prescription Drug Monitoring Program.

3. Entities and practitioners shall submit waiver requests and reports formatted pursuant to a valid waiver to:

Alabama Department of Public Health
Prescription Drug Monitoring Program
The RSA Tower, Suite 1010
P.O. Box 303017
Montgomery, AL 36130-3017
Fax: (334) 206-5663

4. Penalties for noncompliance/non-reporting:

(a) On a monthly basis or as designated by the Prescription Drug Monitoring Program, licensing boards shall supply an electronic listing to the Prescription Drug Monitoring Program of entities and practitioners required to report controlled substances.

(b) The Prescription Drug Monitoring Program will monitor the list of entities and practitioners provided by the licensing boards for compliance in reporting to the database.

(c) The Department will notify the appropriate licensing board of an entity or practitioner’s failure to report. Upon notification of a non-reporting entity or practitioner, the relevant licensing board shall investigate and report to the Department the outcome.

Author: Charles C. Thomas, R. Ph., State Pharmacy Director