

THE GENERAL ASSEMBLY OF PENNSYLVANIA

HOUSE BILL

No. 2334 Session of 2015

INTRODUCED BY GAINEY, DERMODY, MILLARD, TAYLOR, DEAN, DRISCOLL, DAVIS, SNYDER, BULLOCK, REGAN, NEILSON, MAHONEY, YOUNGBLOOD AND GINGRICH, SEPTEMBER 14, 2016

REFERRED TO COMMITTEE ON JUDICIARY, SEPTEMBER 14, 2016

AN ACT

1 Amending the act of April 14, 1972 (P.L.233, No.64), entitled
 2 "An act relating to the manufacture, sale and possession of
 3 controlled substances, other drugs, devices and cosmetics;
 4 conferring powers on the courts and the secretary and
 5 Department of Health, and a newly created Pennsylvania Drug,
 6 Device and Cosmetic Board; establishing schedules of
 7 controlled substances; providing penalties; requiring
 8 registration of persons engaged in the drug trade and for the
 9 revocation or suspension of certain licenses and
 10 registrations; and repealing an act," further providing for
 11 professional prescription, administration and dispensing.

12 The General Assembly of the Commonwealth of Pennsylvania
 13 hereby enacts as follows:

14 Section 1. Section 11 of the act of April 14, 1972 (P.L.233,
 15 No.64), known as The Controlled Substance, Drug, Device and
 16 Cosmetic Act, is amended by adding a subsection to read:

17 Section 11. Professional Prescription, Administration, and
 18 Dispensing.--* * *

19 (a.1) (1) Notwithstanding any provision of this act or any
 20 other law or regulation to the contrary and except as provided
 21 in subparagraph (iii), the following shall apply:

22 (i) When issuing a prescription for an extended release

1 long-acting opioid analgesic to an adult patient for the first
2 time, a practitioner shall not issue a prescription for more
3 than a seven-day supply.

4 (ii) In the case of a minor, a practitioner shall not issue
5 a prescription for an extended release long-acting opioid
6 analgesic for more than a seven-day supply at any time and shall
7 discuss with the parent or guardian the risk of abuse potential
8 associated with the use of extended release long-acting opioid
9 analgesic.

10 (iii) If, in the professional medical judgment of a
11 practitioner, more than a seven-day supply of an extended
12 release long-acting opioid analgesic is required to stabilize an
13 adult patient's emergency medical condition, or the extended
14 release long-acting opioid analgesic is prescribed for chronic
15 pain management, pain associated with a cancer diagnosis or for
16 palliative care, then the practitioner may issue a prescription
17 only for the quantity needed to stabilize the patient's
18 condition.

19 (iv) If a patient's medical condition requires the issuance
20 of a prescription for an opioid analgesic of more than a seven-
21 day supply in accordance with subparagraph (iii), the condition
22 triggering the prescription shall be documented in the patient's
23 medical record and the practitioner shall indicate that a
24 nonopioid alternative, including an opioid analgesic with abuse
25 deterrent properties, was either not appropriate or not
26 available to medically address the patient's medical condition.

27 (2) As used in this section, the following words and phrases
28 shall have the meanings given to them in this section unless the
29 context clearly indicates otherwise:

30 (i) "Extended release long-acting opioid analgesic in a non-

1 abuse deterrent form" means an opioid analgesic which meets all
2 of the following:

3 (A) Is subject to the United States Food and Drug
4 Administration's Extended Release and Long-Acting Opioid
5 Analgesics Risk Evaluation and Mitigation Strategy.

6 (B) Is an opioid analgesic approved for medical use but does
7 not meet the requirements for listing as an opioid analgesic
8 with abuse-deterrent properties pursuant to guidelines published
9 by the United States Food and Drug Administration.

10 (C) Has been identified as an opioid analgesic that poses a
11 heightened risk to public health and safety.

12 (ii) "Nonabuse deterrent opioid" means an opioid analgesic
13 that is approved for medical use but does not meet the
14 requirements for listing as an opioid analgesic with abuse-
15 deterrent properties.

16 (iii) "Opioid analgesic" means a Schedule I, Schedule II or
17 Schedule III controlled substance, which is an opiate-like
18 compound derived from the opium poppy or from synthetic or
19 partially synthetic formulas, as approved by the United States
20 Food and Drug Administration for the alleviation or treatment of
21 pain.

22 (iv) "Opioid analgesic with abuse-deterrent properties"
23 means a formulation or reformulation of an opioid analgesic with
24 physical and chemical properties that make abuse and misuse more
25 difficult.

26 * * *

27 Section 2. This act shall take effect in 60 days.