

THE GENERAL ASSEMBLY OF PENNSYLVANIA

SENATE BILL

No. 514 Session of
2015INTRODUCED BY VANCE, KITCHEN, DINNIMAN, BAKER, VULAKOVICH,
BREWSTER, MENSCH, HUGHES AND AUMENT, FEBRUARY 19, 2015

SENATOR VANCE, PUBLIC HEALTH AND WELFARE, FEBRUARY 25, 2015

AN ACT

1 Amending the act of November 24, 1976 (P.L.1163, No.259),
2 entitled "An act relating to the prescribing and dispensing
3 of generic equivalent drugs," further providing for
4 definitions, for substitutions, for posting requirements, for
5 powers and duties of Department of Health and for immunity of
6 pharmacists under certain circumstances.

7 The General Assembly of the Commonwealth of Pennsylvania
8 hereby enacts as follows:

9 Section 1. Section 2 of the act of November 24, 1976
10 (P.L.1163, No.259), referred to as the Generic Equivalent Drug
11 Law, is amended by adding definitions to read:

12 Section 2. As used in this act:

13 "Biological product" shall have the same meaning as
14 "biological product" in the Public Health Service Act (58 Stat.
15 682, 42 U.S.C. § 207 et seq.).

16 * * *

17 "Interchangeable biological product" means a biological
18 product licensed by the United States Food and Drug
19 Administration and determined to meet the safety standards for
20 interchangeability pursuant to the Public Health Service Act (58

Stat. 682, 42 U.S.C. § 207 et seq.) or a biological product
determined by the United States Food and Drug Administration to
be therapeutically equivalent as set forth in the latest edition <--
or supplement of the United States Food and Drug Administration
Approved Drug Products with Therapeutic Equivalence Evaluations,
sometimes referred to as the "Orange Book." TO A PRESCRIBED <--
BIOLOGICAL PRODUCT.

* * *

Section 2. Section 3(c) and (d) of the act are amended and
the section is amended by adding subsections to read:

Section 3. * * *

(a.1) A pharmacist may substitute a AN INTERCHANGEABLE <--
biological product for a prescribed biological product only if:

(1) the biological product has been determined by the United
States Food and Drug Administration to be interchangeable with
the prescribed product;

(2) the prescriber does not designate verbally or in writing
on the prescription that substitution is prohibited; and

(3) the person presenting the prescription receives
notification of such substitution in the same manner provided in
subsection (b).

(a.2) Within a reasonable time following the dispensing of a <--
AN INTERCHANGEABLE biological product, the dispensing pharmacist <--
or the pharmacist's designee shall communicate to the prescriber
the specific product provided to the patient, including the name
of the product and the manufacturer. The communication shall be
conveyed by making an entry in the electronic health record of
the patient, as defined in the act of July 5, 2012 (P.L.1042,
No.121), known as the "Pennsylvania eHealth Information
Technology Act," or through an electronic prescribing technology

1 or a pharmacy record that is electronically accessible by the
2 prescriber. Otherwise, the pharmacist shall communicate the
3 INTERCHANGEABLE biological product dispensed to the prescriber, <--
4 using facsimile, telephone, electronic transmission or other
5 prevailing means, provided that the communication may not be
6 required where:

7 (1) there is no United States Food and Drug Administration-
8 approved interchangeable biological product for the biological
9 product prescribed; or

10 (2) it is a refill prescription where the biological product
11 dispensed is the same biological product which was dispensed at
12 the prior filling of the prescription and the prescriber was <--
13 notified of the previous substitution.

14 (a.3) Subsections (a.1) and (a.2) may not apply to a
15 biological product which may be dispensed without a
16 prescription.

17 * * *

18 (c) Any pharmacist substituting a less expensive drug
19 product or interchangeable biological product shall charge the
20 purchaser the regular and customary retail price for the
21 generically equivalent drug or interchangeable biological
22 product.

23 (d) Each pharmacist shall maintain a record of any
24 substitution of a generically equivalent drug product or
25 interchangeable biological product for a prescribed brand name
26 drug.

27 * * *

28 Section 3. Sections 4 and 5(a) and (b) of the act, amended
29 July 11, 1990 (P.L.509, No.121), are amended to read:

30 Section 4. (a) Every pharmacy shall post in a prominent

place that is in clear and unobstructed public view, at or near the place where prescriptions are dispensed, a sign which shall read: "Pennsylvania law permits pharmacists to substitute a less expensive generically equivalent drug or interchangeable biological product for a brand name drug unless you or your physician direct otherwise."

(b) Every pharmacy shall post in a conspicuous place, easily accessible to the general public, a list of commonly used generically equivalent drugs and interchangeable biological products containing the generic names and brand names where applicable.

(c) Each pharmacy shall have available to the public a price listing of brand name and generic equivalent drug products and interchangeable biological products available at the pharmacy for selection by the purchaser.

Section 5. (a) The Department of Health shall have the power and its duty shall be to:

(1) Administer and enforce the provisions of this act.

(2) Adopt necessary regulations consistent with this act.

(3) Publicize the provisions of this act.

(4) Publish by notice in the Pennsylvania Bulletin the addition or deletion of generically equivalent drugs and interchangeable biological products and any determination by the secretary to not recognize a generically equivalent drug or interchangeable biological product in accordance with subsection (b). The department shall also provide notice that a complete list of generically equivalent drugs and interchangeable biological products may be obtained from the United States Food and Drug Administration. This notice shall be published at least every three months.

(b) The secretary, with the advice of the Pennsylvania Drug, Device and Cosmetic Board, may determine that a drug shall not be recognized as a generically equivalent drug or interchangeable biological product for purposes of substitution in Pennsylvania and the time after which recognition shall be restored.

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Section 4. Section 6(a) and (b) of the act are amended to read:

Section 6. (a) No pharmacist complying with the provisions of this act shall be liable in any way for the dispensing of a generically equivalent drug or interchangeable biological product unless the generically equivalent drug or interchangeable biological product was incorrectly substituted.

(b) In no event when a pharmacist substitutes a drug or interchangeable biological product shall the prescriber be liable in any action for loss, damage, injury or death or any person occasioned by or arising from the use of the substituted drug or interchangeable biological product unless the original drug was incorrectly prescribed.

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~~Section 5. The addition of section 3(a.2) of the act shall expire five years from the effective date of this act.~~ <--

~~Section 6 5. This act shall take effect as follows:~~ <--

~~(1) The addition of section 3(a.2) of the act shall take effect January 1, 2017, or immediately, whichever is later.~~

~~(2) The remainder of this act shall take effect in 60 days.~~