

1 ENGROSSED

2 COMMITTEE SUBSTITUTE

3 FOR

4 **Senate Bill No. 588**

5 (By Senators Palumbo, Stollings, Plymale, Jenkins and Barnes)

6 \_\_\_\_\_  
7 [Originating in the Committee on the Judiciary;  
8 reported February 24, 2012.]  
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11  
12 A BILL to repeal §60A-8-4 of the Code of West Virginia, 1931, as  
13 amended; to amend and reenact §60A-8-3, §60A-8-5 and §60A-8-7  
14 of said code; and to amend said code by adding thereto three  
15 new sections, designated §60A-8-14, §60A-8-15 and §60A-8-16,  
16 all relating generally to wholesale drug distributors licensed  
17 by Board of Pharmacy; specifying purpose of article; defining  
18 terms; specifying wholesale drug distributor licensing  
19 requirements; specifying powers of Board of Pharmacy;  
20 increasing licensing fees; requiring updates when material  
21 changes occur to a licensee; authorizing board to take certain  
22 disciplinary action against licensees, including revocation or  
23 suspension of licenses, refusal to renew license and civil  
24 penalties; providing for register of wholesale and pharmacy

1 distributors of prescription drugs; and providing for the  
2 disposition of fees.

3 *Be it enacted by the Legislature of West Virginia:*

4 That §60A-8-4 of the Code of West Virginia, 1931, as amended,  
5 be repealed; that §60A-8-3, §60A-8-5 and §60A-8-7 of said code be  
6 amended and reenacted; and that said code be amended by adding  
7 thereto three new sections, designated §60A-8-14, §60A-8-15 and  
8 §60A-8-16, all to read as follows:

9 **ARTICLE 8. WHOLESALE DRUG DISTRIBUTION LICENSING ACT OF 1991.**

10 **§60A-8-3. Purpose.**

11 The purpose of this article is to protect the health, safety  
12 and general welfare of residents of this state and to implement the  
13 federal Prescription Drug Marketing Act of 1987 ("PDMA"), U. S.  
14 Public Law 100-293, 102 Stat. 95, codified at 21 U. S. Code §321;  
15 and particularly PDMA requirements that no person or entity may  
16 engage in the wholesale distribution of human prescription drugs in  
17 any state unless such person or entity is licensed by such state in  
18 accordance with federally-prescribed minimum standards, terms and  
19 conditions as set forth in guidelines issued by United States food  
20 and drug administration (FDA) regulations pursuant to 21 U. S. Code  
21 §353(e) (2) (A) and (B); and such regulations as are set forth in 21  
22 C. F. R. Part 205.

23 **§60A-8-5. Definitions.**

24 As used in this article:

1 (a) "Wholesale distribution" and "wholesale distributions"  
2 mean distribution of prescription drugs, including directly or  
3 through the use of a third-party logistics provider or any other  
4 situation in which title, ownership or control over the  
5 prescription drug remains with one person or entity but the  
6 prescription drug is brought into this state by another person or  
7 entity on his, her or its behalf, to persons other than a consumer  
8 or patient, but does not include:

9 (1) Intracompany sales, being defined as any transaction, ~~or~~  
10 transfer or delivery into or within this state between any  
11 division, subsidiary, parent and/or affiliated or related company  
12 under the common ownership and control of a corporate entity;

13 (2) The purchase or other acquisition by a hospital or other  
14 health care entity that is a member of a group purchasing  
15 organization of a drug for its own use from the group purchasing  
16 organization or from other hospitals or health care entities that  
17 are members of such organizations;

18 (3) The sale, purchase or trade of a drug or an offer to sell,  
19 purchase or trade a drug by a charitable organization described in  
20 section 501(c)(3) of the United States Internal Revenue Code of  
21 ~~1954~~ 1986 to a nonprofit affiliate of the organization to the  
22 extent otherwise permitted by law;

23 (4) The sale, purchase or trade of a drug or an offer to sell,  
24 purchase or trade a drug among hospitals or other health care

1 entities that are under common control. For purposes of this  
2 article, "common control" means the power to direct or cause the  
3 direction of the management and policies of a person or an  
4 organization, whether by ownership of stock, voting rights, by  
5 contract, or otherwise;

6 (5) The sale, purchase or trade of a drug or an offer to sell,  
7 purchase or trade a drug for "emergency medical reasons" for  
8 purposes of this article includes transfers of prescription drugs  
9 by a retail pharmacy to another retail pharmacy to alleviate a  
10 temporary shortage, except that the gross dollar value of such  
11 transfers shall not exceed five percent of the total prescription  
12 drug sales revenue of either the transferor or ~~transferee~~ transferee  
13 pharmacy during any twelve consecutive month period;

14 (6) The sale, purchase or trade of a drug, an offer to sell,  
15 purchase, or trade a drug or the dispensing of a drug pursuant to  
16 a prescription;

17 (7) The distribution of drug samples by manufacturers'  
18 representatives or distributors' representatives, if the  
19 distribution is permitted under federal law [21 U. S. C. 353(d)];  
20 or

21 (8) The sale, purchase or trade of blood and blood components  
22 intended for transfusion.

23 (b) "Wholesale drug distributor" or "wholesale distributor"  
24 means any person or entity engaged in wholesale distribution of

1 prescription drugs, including, but not limited to, manufacturers,  
2 repackers, own-label distributors, jobbers, private-label  
3 distributors, brokers, warehouses, including manufacturers' and  
4 distributors' warehouses, chain drug warehouses and wholesale drug  
5 warehouses, independent wholesale drug traders, prescription drug  
6 repackagers, physicians, dentists, veterinarians, birth control and  
7 other clinics, individuals, hospitals, nursing homes and/or their  
8 providers, health maintenance organizations and other health care  
9 providers, and retail and hospital pharmacies that conduct  
10 wholesale distributions, including, but not limited to, any  
11 pharmacy distributor as defined in this section. A wholesale drug  
12 distributor shall not include any for hire carrier or person or  
13 entity hired solely to transport prescription drugs.

14 (c) "Pharmacy distributor" means any pharmacy licensed in this  
15 state or hospital pharmacy which is engaged in the delivery or  
16 distribution of prescription drugs either to any other pharmacy  
17 licensed in this state or to any other person or entity, including,  
18 but not limited to, a wholesale drug distributor as defined in  
19 subdivision (b) of this section engaged in the delivery or  
20 distribution of prescription drugs and who is involved in the  
21 actual, constructive or attempted transfer of a drug in this state  
22 to other than the ultimate consumer except as otherwise provided  
23 for by law.

24 (d) "Manufacturer" means ~~anyone~~ any person who is engaged in

1 manufacturing, preparing, propagating, compounding, processing,  
2 packaging, repackaging or labeling of a prescription drug, whether  
3 within or outside this state.

4 (e) "West Virginia Board of Pharmacy", "Board of Pharmacy" or  
5 "board" means the agency of this state authorized to license  
6 wholesale drug distribution except where otherwise provided.

7 (f) "Prescription drug" means any human drug required by  
8 federal law or regulation to be dispensed only by prescription,  
9 including finished dosage forms and active ingredients subject to  
10 section 503(b) of the federal food, drug and cosmetic act.

11 (g) "Blood" means whole blood collected from a single donor  
12 and processed either for transfusion or further manufacturing.

13 (h) "Blood component" means that part of blood separated by  
14 physical or mechanical means.

15 (i) "Drug sample" means a unit of a prescription drug that is  
16 not intended to be sold and is intended to promote the sale of the  
17 drug.

18 (j) "Person" means any individual, partnership, association,  
19 limited liability company, corporation or other entity.

20 (k) "Key person" means any of the following:

21 (1) An officer, director, trustee, partner, principal or  
22 proprietor of a person that has applied for or holds a license  
23 issued under this article or an affiliate or holding company that  
24 has control of a person that has applied for or holds a license

1 under this article.

2 (2) A person that holds a combined direct, indirect or  
3 attributed debt or equity interest of more than five percent in a  
4 person that has applied for or holds a license under this article;

5 (3) A person that holds a combined direct, indirect or  
6 attributed equity interest of more than five percent in a person  
7 that has a controlling interest in a person that has applied for or  
8 holds license under this article;

9 (4) A managerial employee of a person that has applied for or  
10 holds a license under this article or a managerial employee of an  
11 affiliate or holding company that has control of a person that has  
12 applied for or holds a license under this article, who performs the  
13 function of principal executive officer, principal operating  
14 officer, principal accounting officer or an equivalent officer;

15 (5) A managerial employee of a person that has applied for or  
16 holds a license under this article or a managerial employee of an  
17 affiliate or holding company that has control of a person that has  
18 applied for or holds a license under this article who will perform  
19 or performs the function of an operations manager or will exercise  
20 or exercises management, supervisory or policy-making authority  
21 over the distribution of prescription drugs.

22 (1) "Third-party logistics provider" means a person who  
23 contracts with a prescription drug manufacturer to provide or  
24 coordinate warehousing, distribution or other services on behalf of

1 a manufacturer, but does not take title to the prescription drug or  
2 have general responsibility to direct the prescription drug's sale  
3 or disposition. A third-party logistics provider must be licensed  
4 as a wholesale distributor under this article and, in order to be  
5 considered part of the normal distribution channel, must also be an  
6 authorized distributor of record.

7 **§60A-8-7. Wholesale drug distributor licensing requirements.**

8 (a) Every applicant for a license under this article shall  
9 provide the board with the following as part of the application for  
10 a license and as part of any renewal of such license:

11 (1) The name, full business address and telephone number of  
12 the licensee;

13 (2) All trade or business names used by the licensee;

14 (3) Addresses, telephone numbers and the names of contact  
15 persons for all facilities used by the licensee for the storage,  
16 handling and distribution of prescription drugs;

17 (4) The type of ownership or operation (i.e., partnership,  
18 corporation or sole proprietorship);

19 (5) The name(s) of the owner and operator, or both, of the  
20 licensee, including:

21 (A) If a person, the name of the person;

22 (B) If a partnership, the name of each partner and the name of  
23 the partnership;

24 (C) If a corporation, the name and title of each corporate



1 officer and director, the corporate names and the name of the state  
2 of incorporation; and

3 (D) If a sole proprietorship, the full name of the sole  
4 proprietor and the name of the business entity; and

5 (6) Any other information or documentation that the board may  
6 require.

7 (b) All wholesale distributors and pharmacy distributors shall  
8 be subject to the following requirements:

9 (a) (1) No person or distribution outlet may act as a  
10 wholesale drug distributor without first obtaining a license to do  
11 so from the Board of Pharmacy and paying any reasonable fee  
12 required by the Board of Pharmacy, such fee not to exceed four  
13 hundred dollars per year: *Provided, That for licenses that are*  
14 effective on and after July 1, 2012, the annual fee shall be \$750  
15 per license until modified by legislative rule.

16 (b) (2) The Board of Pharmacy may grant a temporary license  
17 when a wholesale drug distributor first applies to the board for a  
18 ~~wholesale drug distributor's license to operate within this state~~  
19 and the temporary license shall remain valid until the Board of  
20 Pharmacy finds that the applicant meets or fails to meet the  
21 requirements for regular licensure, except that no temporary  
22 license shall be valid for more than ninety days from the date of  
23 issuance. Any temporary license issued pursuant to this subdivision  
24 shall be renewable for a similar period of time not to exceed

1 ninety days pursuant to policies and procedures to be prescribed by  
2 the Board of Pharmacy.

3 ~~(c)~~ (3) No license may be issued or renewed for a wholesale  
4 drug distributor to operate unless the distributor operates in a  
5 manner prescribed by law and according to the rules promulgated by  
6 the Board of Pharmacy with respect thereto.

7 ~~(d)~~ (4) The Board of Pharmacy may require a separate license  
8 for each facility directly or indirectly owned or operated by the  
9 same business entity within this state, or for a parent entity with  
10 divisions, subsidiaries, or affiliate companies within this state  
11 when operations are conducted at more than one location and there  
12 exists joint ownership and control among all the entities.

13 ~~(e)~~ (c) The minimum qualifications for licensure are set forth  
14 in this section as follows:

15 (1) As a condition for receiving and retaining any wholesale  
16 drug distributor license issued pursuant to this article, each  
17 applicant shall satisfy the Board of Pharmacy that it has and will  
18 continuously maintain:

19 (A) Acceptable storage and handling conditions plus facilities  
20 standards;

21 (B) Minimum liability and other insurance as may be required  
22 under any applicable federal or state law;

23 (C) A security system which includes after hours central alarm  
24 or comparable entry detection capability, restricted premises

1 access, adequate outside perimeter lighting, comprehensive  
2 employment applicant screening and safeguards against employee  
3 theft;

4 (D) An electronic, manual or any other reasonable system of  
5 records describing all wholesale distributor activities governed by  
6 this article for the two-year period following disposition of each  
7 product and being reasonably accessible as defined by Board of  
8 Pharmacy regulations during any inspection authorized by the Board  
9 of Pharmacy;

10 (E) Officers, directors, managers and other persons in charge  
11 of wholesale drug distribution, storage and handling, who must at  
12 all times demonstrate and maintain their capability of conducting  
13 business according to sound financial practices as well as state  
14 and federal law;

15 (F) Complete, updated information to be provided to the Board  
16 of Pharmacy as a condition for obtaining and retaining a license  
17 about each wholesale distributor to be licensed under this article  
18 including all pertinent licensee ownership and other key personnel  
19 and facilities information determined necessary for enforcement of  
20 this article; ~~with any changes in the information to be submitted~~  
21 ~~at the time of license renewal or within twelve months from the~~  
22 ~~date of the change, whichever occurs first;~~

23 (G) Written policies and procedures which assure reasonable  
24 wholesale distributor preparation for protection against and

1 handling of any facility security or operation problems, including,  
2 but not limited to, those caused by natural disaster or government  
3 emergency, inventory inaccuracies or product shipping and  
4 receiving, outdated product or other unauthorized product control,  
5 appropriate disposition of returned goods and product recalls;

6 (H) Sufficient inspection procedures for all incoming and  
7 outgoing product shipments; and

8 (I) Operations in compliance with all federal legal  
9 requirements applicable to wholesale drug distribution.

10 (2) The board of pharmacy shall consider, at a minimum, the  
11 following factors in reviewing the qualifications of persons who  
12 ~~engage in wholesale distribution of prescription drugs with this~~  
13 ~~state~~ apply for a wholesale distributor license under this section  
14 or for renewal of that license:

15 (A) Any conviction of the applicant under any federal, state  
16 or local laws relating to drug samples, wholesale or retail drug  
17 distribution or distribution of controlled substances;

18 (B) Any felony convictions of the applicant or any key person  
19 under federal, state or local laws;

20 (C) The applicant's past experience in the manufacture or  
21 distribution of prescription drugs, including, but not limited to,  
22 controlled substances;

23 (D) The furnishing by the applicant of false or fraudulent  
24 material in any application made in connection with drug

1 manufacturing or distribution;

2 (E) Suspension or revocation by federal, state or local  
3 government of any license currently or previously held by the  
4 applicant for the manufacture or distribution of any drug,  
5 including, but not limited to, controlled substances;

6 (F) Compliance with licensing requirements under previously  
7 granted licenses, if any;

8 (G) Whether personnel employed by the applicant in wholesale  
9 drug distribution have appropriate education or experience, or both  
10 education and experience, to assume responsibility for positions  
11 related to compliance with the requirements of this article;

12 ~~(G)~~ (H) Compliance with requirements to maintain and make  
13 available to the Board of Pharmacy or to federal, state or local  
14 law-enforcement officials those records required by this article;  
15 and

16 ~~(H)~~ (I) Any other factors or qualifications the Board of  
17 Pharmacy considers relevant to and consistent with the public  
18 health and safety, including whether the granting of the license  
19 would not be in the public interest.

20 (3) All requirements set forth in this subsection shall  
21 conform to wholesale drug distributor licensing guidelines formally  
22 adopted by the United States Food and Drug Administration (FDA);  
23 and in case of conflict between any wholesale drug distributor  
24 licensing requirement imposed by the Board of Pharmacy pursuant to

1 this subsection and any food and drug administration wholesale drug  
2 distributor licensing guideline, the latter shall control.

3 ~~(f)~~ (d) An ~~agent or~~ employee of any licensed wholesale drug  
4 distributor need not seek licensure under this section and may  
5 lawfully possess pharmaceutical drugs when the ~~agent or~~ employee is  
6 acting in the usual course of business or employment.

7 ~~(g)~~ (e) The issuance of a license pursuant to this article  
8 does not change or affect tax liability imposed by this state's  
9 Department of Tax and Revenue on any wholesale drug distributor.

10 (f) An applicant who is awarded a license or renewal of a  
11 license shall give the board written notification of any material  
12 change in the information previously submitted in, or with the  
13 application for the license or for renewal thereof, whichever is  
14 the most recent document filed with the board, within thirty days  
15 after the material change occurs or the licensee becomes aware of  
16 the material change, whichever event occurs last. Material changes  
17 include, but are not limited to:

18 (1) A change of the physical address or mailing address;

19 (2) A change of the responsible individual, compliance officer  
20 or other executive officers or board members;

21 (3) A change of the licensee's name or trade name;

22 (4) A change in the location where the records of the licensee  
23 are retained;

24 (5) The felony conviction of a key person of the licensee; and

1       (6) Any other material change that the board may specify by  
2 rule.

3       (g) The board may deny a license to an applicant for a license  
4 or for renewal of a license if the board determines that the  
5 granting of the license would not be in the public interest.

6       (h) The licensing of any person as a wholesale drug  
7 distributor subjects the person and the person's agents and  
8 employees to the jurisdiction of the board and to the laws of this  
9 state for the purpose of the enforcement of this article, article  
10 five, chapter thirty of this code and the rules of the board.  
11 However, the filing of an application for a license as a wholesale  
12 drug distributor by, or on behalf of, any person or the licensing  
13 of any person as a wholesale drug distributor may not, of itself,  
14 constitute evidence that the person is doing business within this  
15 state.

16       ~~(h)~~ (i) The Board of Pharmacy may adopt rules pursuant to  
17 section nine of this article which permit out-of-state wholesale  
18 drug distributors to obtain any license required by this article on  
19 the basis of reciprocity to the extent that: ~~(i)~~ (1) An out-of-  
20 state wholesale drug distributor possesses a valid license granted  
21 by another state pursuant to legal standards comparable to those  
22 which must be met by a wholesale drug distributor of this state as  
23 prerequisites for obtaining a license under the laws of this state;  
24 and ~~(ii)~~ (2) such other state would extend reciprocal treatment

1 under its own laws to a wholesale drug distributor of this state.

2 **§60A-8-14. Disciplinary actions - wholesale drug distributor.**

3 (a) In accordance with article five, chapter thirty of this  
4 code, the Board of Pharmacy may suspend, revoke or refuse to renew  
5 any license issued to a wholesale distributor of prescription drugs  
6 pursuant to this article or may impose a civil money penalty not to  
7 exceed \$1,000, in the discretion of the board for any of the  
8 following causes:

9 (1) Making any false material statements in an application for  
10 a license or for renewal of a license as a wholesale distributor or  
11 pharmacy distributor of prescription drugs;

12 (2) Violating any federal, state or local drug law, any  
13 provision of this article or any rule of the board;

14 (3) Conviction of a felony. For purposes of this subdivision  
15 "felony" means a felony or crime punishable as a felony under the  
16 laws of this state, any other state or the United States;

17 (4) Ceasing to satisfy the qualifications for licensure under  
18 section seven of this article or the rules of the board;

19 (5) The license or registration of a wholesale drug  
20 distributor licensed under this article has been revoked by the  
21 licensing authority of another state, jurisdiction of foreign  
22 nation; or

23 (6) Any reason for which the board may impose disciplinary  
24 sanctions under the provisions of chapter thirty of this code.



1 (b) Upon the suspension or revocation of the license of any  
2 wholesale distributor of prescription drugs, the distributor shall  
3 immediately surrender the license to the board.

4 (c) If the board suspends, revokes or refuses to renew any  
5 license issued to a wholesale distributor of prescription drugs and  
6 determines that there is clear and convincing evidence of a danger  
7 of immediate and serious harm to any person, the board may place  
8 under seal all drugs owned by or in the possession, custody or  
9 control of the affected wholesale distributor. Except as provided  
10 in this article, the board may not dispose of the drugs sealed  
11 under this subsection until the distributor exhausts all of his or  
12 her appeal rights under this article or article five, chapter  
13 thirty of this code. The court involved in the appeal may order the  
14 board, during the pendency of the appeal, to sell sealed dangerous  
15 drugs that are perishable. The board shall deposit the proceeds of  
16 the sale with the court.

17 **§60A-8-15. Maintenance of register and roster of wholesale and**  
18 **pharmacy distributors.**

19 (a) The Executive Director of the Board of Pharmacy shall  
20 maintain a register of the names, addresses and the date the  
21 current license was issued or renewed pursuant to this article for  
22 license years beginning on and after July 1, 2013. The register  
23 shall be the property of the board and shall be open for public  
24 examination and inspection at all reasonable times, as the board

1 may direct.

2 (b) The register shall set forth the names and addresses of:

3 (1) Those persons who are or have been licensed under this  
4 article for the current license year;

5 (2) Those persons whose licenses have been suspended, revoked  
6 or surrendered during the current license year or during the two  
7 preceding license years; and

8 (3) Those persons whose licenses have not been renewed for the  
9 current license year.

10 (c) In lieu of annually publishing a typed or printed register  
11 providing the information required by this subsection, the board  
12 may make the information required to be published available at its  
13 website.

14 (d) A written statement signed and verified by the executive  
15 director of the board, in which it is stated that after diligent  
16 search of the register no record or entry of the issuance of a  
17 license or registration certificate to a person is found, is  
18 admissible in evidence and constitutes presumptive evidence of the  
19 fact that the person is not a licensed as a wholesale drug  
20 distributor under this article.

21 **§60A-8-16. Disposition of fees.**

22 The board shall pay all fees it collects under this article  
23 into the separate fund created in the State Treasury for the board  
24 pursuant to section ten, article one, chapter thirty of this code.

1 The money in this fund shall be used exclusively by the board for  
2 the purposes of administering and enforcement of its duties  
3 pursuant to this article, articles one and five, chapter thirty of  
4 this code, or any other duty of the board prescribed by any other  
5 provision of this code.