

Adopted	Rejected
---------	----------

## COMMITTEE REPORT

YES:	10
NO:	0

### MR. SPEAKER:

*Your Committee on Public Health, to which was referred Senate Bill 590, has had the same under consideration and begs leave to report the same back to the House with the recommendation that said bill **be amended** as follows:*

- 1           Page 1, between the enacting clause and line 1, begin a new
- 2           paragraph and insert:
- 3           "SECTION 1. IC 10-13-3-38.5 IS AMENDED TO READ AS
- 4           FOLLOWS [EFFECTIVE JULY 1, 2005]: Sec. 38.5. (a) Under federal
- 5           P.L.92-544 (86 Stat. 1115), the department may use an individual's
- 6           fingerprints submitted by the individual for the following purposes:
- 7           (1) Determining the individual's suitability for employment with
- 8           the state, or as an employee of a contractor of the state, in a
- 9           position:
- 10           (A) that has a job description that includes contact with, care
- 11           of, or supervision over a person less than eighteen (18) years
- 12           of age;
- 13           (B) that has a job description that includes contact with, care
- 14           of, or supervision over an endangered adult (as defined in
- 15           IC 12-10-3-2), except the individual is not required to meet the

- 1 standard for harmed or threatened with harm set forth in  
 2 IC 12-10-3-2(a)(3);  
 3 (C) at a state institution managed by the office of the secretary  
 4 of family and social services or state department of health;  
 5 (D) at the Indiana School for the Deaf established by  
 6 IC 20-16-2-1;  
 7 (E) at the Indiana School for the Blind established by  
 8 IC 20-15-2-1;  
 9 (F) at a juvenile detention facility;  
 10 (G) with the gaming commission under IC 4-33-3-16;  
 11 (H) with the department of financial institutions under  
 12 IC 28-11-2-3; or  
 13 (I) that has a job description that includes access to or  
 14 supervision over state financial or personnel data, including  
 15 state warrants, banking codes, or payroll information  
 16 pertaining to state employees.
- 17 (2) Identification in a request related to an application for a  
 18 teacher's license submitted to the professional standards board  
 19 established under IC 20-1-1.4.
- 20 **(3) Use by the Indiana board of pharmacy in determining the**  
 21 **individual's suitability for a position or employment with a**  
 22 **wholesale drug distributor, as specified in IC 25-26-14-16(b),**  
 23 **IC 25-26-14-16.5(b), IC 25-26-14-17.8(c), and IC 25-26-14-20.**
- 24 An applicant shall submit the fingerprints in an appropriate format or  
 25 on forms provided for the employment or license application. The  
 26 department shall charge each applicant the fee established under section  
 27 28 of this chapter and by federal authorities to defray the costs  
 28 associated with a search for and classification of the applicant's  
 29 fingerprints. The department may forward fingerprints submitted by an  
 30 applicant to the Federal Bureau of Investigation or any other agency for  
 31 processing. The state personnel department or the agency to which the  
 32 applicant is applying for employment or a license may receive the  
 33 results of all fingerprint investigations.
- 34 (b) An applicant who is an employee of the state may not be charged  
 35 under subsection (a).
- 36 (c) Subsection (a)(1) does not apply to an employee of a contractor  
 37 of the state if the contract involves the construction or repair of a capital  
 38 project or other public works project of the state."

1 Page 2, line 35, after "writing" insert "**or is entered into an**  
2 **electronic format**".

3 Page 2, line 35, delete "pharmacist." and insert "pharmacist **or**  
4 **pharmacist intern (as defined by IC 25-26-13-2)**".

5 Page 7, line 25, after "written" insert "**or electronically**  
6 **transmitted**".

7 Page 13, line 39, delete "computer," and insert "**computer**".

8 Page 17, line 3, delete "pharmacist" and insert "**pharmacy**".

9 Page 17, between lines 5 and 6, begin a new paragraph and insert:  
10 "SECTION 19. IC 25-26-14-1 IS AMENDED TO READ AS  
11 FOLLOWS [EFFECTIVE JULY 1, 2005]: Sec. 1. **(a)** This chapter  
12 applies to any individual, partnership, limited liability company,  
13 corporation, or business firm:

14 **(1) located within or outside Indiana; and**

15 **(2) engaging in the wholesale distribution of legend drugs ~~within~~**  
16 **in Indiana.**

17 **(b) Except as required by federal law or regulation, the**  
18 **requirements of this chapter do not apply to a manufacturer that**  
19 **is approved by the federal Food and Drug Administration.**  
20 **However, the board may adopt rules concerning manufacturers**  
21 **that the board considers appropriate and necessary.**

22 SECTION 20. IC 25-26-14-1.5 IS ADDED TO THE INDIANA  
23 CODE AS A NEW SECTION TO READ AS FOLLOWS  
24 [EFFECTIVE JULY 1, 2005]: Sec. 1.5. **As used in this chapter,**  
25 **"adulterated" refers to a drug that:**

26 **(1) consists in whole or in part of a filthy, putrid, or**  
27 **decomposed substance;**

28 **(2) has been produced, prepared, packed, or held under**  
29 **unsanitary conditions and may have been contaminated or**  
30 **rendered injurious to health;**

31 **(3) has been subjected to conditions in the manufacture,**  
32 **processing, packing, or holding of the drug that do not**  
33 **conform to current standards of manufacturing to ensure that**  
34 **the drug is safe for use and possesses the identity, strength,**  
35 **quality, and purity characteristics that the drug is represented**  
36 **to possess;**

37 **(4) is contained in a container composed of a poisonous or**  
38 **deleterious substance that may render the drug injurious to**

- 1           **health;**  
 2           **(5) bears or contains, for purposes of coloring only, a color**  
 3           **additive that is unsafe;**  
 4           **(6) is of a different strength, quality, or purity from the official**  
 5           **compendium standard for the drug; or**  
 6           **(7) does not meet the considerations of the federal Food, Drug,**  
 7           **and Cosmetic Act.**

8           SECTION 21. IC 25-26-14-1.7 IS ADDED TO THE INDIANA  
 9           CODE AS A **NEW** SECTION TO READ AS FOLLOWS  
 10          [EFFECTIVE JULY 1, 2005]: **Sec. 1.7. As used in this chapter,**  
 11          **"authenticate" means to affirmatively verify before distribution**  
 12          **occurs that each transaction that is listed on:**

- 13               **(1) the pedigree of a drug; and**  
 14               **(2) other accompanying documentation for a drug;**  
 15          **has occurred.**

16          SECTION 22. IC 25-26-14-1.8 IS ADDED TO THE INDIANA  
 17          CODE AS A **NEW** SECTION TO READ AS FOLLOWS  
 18          [EFFECTIVE JULY 1, 2005]: **Sec. 1.8. As used in this chapter,**  
 19          **"authorized distributor" means a wholesale drug distributor with**  
 20          **which a manufacturer has established an ongoing relationship to**  
 21          **distribute the manufacturer's products. For purposes of this**  
 22          **section, an ongoing relationship exists between a wholesale drug**  
 23          **distributor, including any affiliated group (as defined in Section**  
 24          **1504 of the Internal Revenue Code) of which the wholesale**  
 25          **distributor is a member, and a manufacturer if the wholesale drug**  
 26          **distributor:**

- 27               **(1) has a written agreement currently in effect with the**  
 28               **manufacturer evidencing an ongoing relationship;**  
 29               **(2) is listed on the manufacturer's current monthly updated**  
 30               **list of authorized distributors; or**  
 31               **(3) has a verifiable account with the manufacturer and a**  
 32               **minimal transaction or volume requirement limit of:**  
 33                       **(A) five thousand (5,000) units per company in the previous**  
 34                       **twelve (12) months; or**  
 35                       **(B) twelve (12) purchases at the manufacturer's minimum**  
 36                       **purchasing requirement per invoice in the previous twelve**  
 37                       **(12) months.**

38          SECTION 23. IC 25-26-14-4.1 IS ADDED TO THE INDIANA

1 CODE AS A NEW SECTION TO READ AS FOLLOWS  
2 [EFFECTIVE JULY 1, 2005]: **Sec. 4.1. As used in this chapter,**  
3 **"compendium" refers to:**

- 4 (1) **the United States Pharmacopoeia;**
- 5 (2) **the Homeopathic Pharmacopoeia of the United States;**
- 6 (3) **the National Formulary;**
- 7 (4) **a drug approved by the federal Food and Drug**  
8 **Administration; or**
- 9 (5) **a supplement to a document specified in subdivision (1),**  
10 **(2), or (3).**

11 SECTION 24. IC 25-26-14-4.2 IS ADDED TO THE INDIANA  
12 CODE AS A NEW SECTION TO READ AS FOLLOWS  
13 [EFFECTIVE JULY 1, 2005]: **Sec. 4.2. As used in this chapter,**  
14 **"contraband" refers to a drug:**

- 15 (1) **that is counterfeit;**
- 16 (2) **that is stolen;**
- 17 (3) **that is misbranded;**
- 18 (4) **that is obtained by fraud;**
- 19 (5) **that is purchased by a nonprofit institution for the**  
20 **nonprofit institution's own use and placed in commerce in**  
21 **violation of the own use agreement for the drug;**
- 22 (6) **for which a required pedigree does not exist; or**  
23 (7) **for which a pedigree in existence:**
  - 24 (A) **has been forged, counterfeited, or falsely created; or**
  - 25 (B) **contains any altered, false, or misrepresented**  
26 **information.**

27 SECTION 25. IC 25-26-14-4.3 IS ADDED TO THE INDIANA  
28 CODE AS A NEW SECTION TO READ AS FOLLOWS  
29 [EFFECTIVE JULY 1, 2005]: **Sec. 4.3. As used in this chapter,**  
30 **"counterfeit" refers to a drug, or the container, seal, or labeling of**  
31 **a drug, that, without authorization, bears the trademark, trade**  
32 **name, or other identifying mark or imprint of a manufacturer,**  
33 **processor, packer, or distributor other than the person that**  
34 **manufactured, processed, packed, or distributed the drug.**

35 SECTION 26. IC 25-26-14-4.4 IS ADDED TO THE INDIANA  
36 CODE AS A NEW SECTION TO READ AS FOLLOWS  
37 [EFFECTIVE JULY 1, 2005]: **Sec. 4.4. As used in this chapter,**  
38 **"deliver" means the actual, constructive, or attempted transfer of**

1 **a drug from one (1) person to another.**

2 SECTION 27. IC 25-26-14-4.5 IS ADDED TO THE INDIANA  
3 CODE AS A NEW SECTION TO READ AS FOLLOWS  
4 [EFFECTIVE JULY 1, 2005]: **Sec. 4.5. As used in this chapter,**  
5 **"designated representative" means an individual who:**

- 6 (1) **is designated by a wholesale drug distributor;**  
7 (2) **serves as the wholesale drug distributor's responsible**  
8 **individual with the board; and**  
9 (3) **is actively involved in and aware of the actual daily**  
10 **operation of the wholesale drug distributor.**

11 SECTION 28. IC 25-26-14-4.7 IS ADDED TO THE INDIANA  
12 CODE AS A NEW SECTION TO READ AS FOLLOWS  
13 [EFFECTIVE JULY 1, 2005]: **Sec. 4.7. As used in this chapter,**  
14 **"distribute" means to sell, offer to sell, deliver, offer to deliver,**  
15 **broker, give away, or transfer a legend drug, whether by passage**  
16 **of title or physical movement, or both. The term does not include**  
17 **the following:**

- 18 (1) **Dispensing or administering a legend drug.**  
19 (2) **Delivering or offering to deliver a legend drug by a**  
20 **common carrier in the usual course of business as a common**  
21 **carrier.**  
22 (3) **The provision of a drug sample to a patient by a:**  
23 (A) **practitioner;**  
24 (B) **health care professional acting at the direction and**  
25 **under the supervision of a practitioner; or**  
26 (C) **hospital's or other health care entity's pharmacy that**  
27 **received the drug sample in accordance with this chapter**  
28 **and other applicable law to administer or dispense and that**  
29 **is acting at the direction of a practitioner;**  
30 **licensed to prescribe the legend drug.**

31 SECTION 29. IC 25-26-14-4.9 IS ADDED TO THE INDIANA  
32 CODE AS A NEW SECTION TO READ AS FOLLOWS  
33 [EFFECTIVE JULY 1, 2005]: **Sec. 4.9. As used in this chapter,**  
34 **"drug" means any of the following:**

- 35 (1) **Articles recognized in an official compendium and**  
36 **designated by the board for use in the diagnosis, cure,**  
37 **mitigation, treatment, or prevention of disease in humans or**  
38 **animals.**

1           **(2) Articles intended for use in the diagnosis, cure, mitigation,**  
 2           **treatment, or prevention of disease in humans or animals.**

3           **(3) Articles other than food intended to affect the structure or**  
 4           **function of the body of humans or animals.**

5           **(4) Articles intended for use as a component of an article**  
 6           **specified in subdivision (1), (2), or (3).**

7           **The term does not include a device or a device component, part, or**  
 8           **accessory.**

9           SECTION 30. IC 25-26-14-6 IS AMENDED TO READ AS  
 10          FOLLOWS [EFFECTIVE JULY 1, 2005]: Sec. 6. As used in this  
 11          chapter, "health care entity" means any organization or business that  
 12          provides diagnostic, medical, surgical, dental treatment, or  
 13          rehabilitative care. **The term does not include a pharmacy or**  
 14          **wholesale drug distributor.**

15          SECTION 31. IC 25-26-14-6.5 IS ADDED TO THE INDIANA  
 16          CODE AS A NEW SECTION TO READ AS FOLLOWS  
 17          [EFFECTIVE JULY 1, 2005]: **Sec. 6.5. As used in this chapter,**  
 18          **"label" means a display of written, printed, or graphic matter on**  
 19          **the immediate container of a legend drug.**

20          SECTION 32. IC 25-26-14-6.6 IS ADDED TO THE INDIANA  
 21          CODE AS A NEW SECTION TO READ AS FOLLOWS  
 22          [EFFECTIVE JULY 1, 2005]: **Sec. 6.6. As used in this chapter,**  
 23          **"labeling" means labels and other written, printed, or graphic**  
 24          **matter:**

25                 **(1) on a legend drug or a legend drug's container or wrapper;**  
 26                 **or**

27                 **(2) accompanying a legend drug.**

28          SECTION 33. IC 25-26-14-8.3 IS ADDED TO THE INDIANA  
 29          CODE AS A NEW SECTION TO READ AS FOLLOWS  
 30          [EFFECTIVE JULY 1, 2005]: **Sec. 8.3. As used in this chapter,**  
 31          **"misbranded" means that a legend drug's label:**

32                 **(1) is false or misleading;**

33                 **(2) does not bear the name and address of the manufacturer,**  
 34                 **packer, or distributor or does not contain an accurate**  
 35                 **statement of the quantities of active ingredients of the legend**  
 36                 **drug;**

37                 **(3) does not show an accurate monograph for the legend drug;**  
 38                 **or**

1           **(4) does not comply with any other requirements of the federal**  
 2           **Food, Drug and Cosmetic Act.**

3           SECTION 34. IC 25-26-14-8.7 IS ADDED TO THE INDIANA  
 4           CODE AS A NEW SECTION TO READ AS FOLLOWS  
 5           [EFFECTIVE JULY 1, 2005]: **Sec. 8.7. As used in this chapter,**  
 6           **"pedigree" means a statement or record in a written or an**  
 7           **electronic form that is approved by the board, that records each**  
 8           **distribution of a legend drug from the sale by the manufacturer or,**  
 9           **except for drugs on the specified list of susceptible products, from**  
 10           **the last authorized distributor of record through acquisition and**  
 11           **sale by each wholesale drug distributor, and that includes the**  
 12           **following information for each transaction:**

13           **(1) The source of the legend drug, including the name and**  
 14           **principal address of the seller.**

15           **(2) The:**

16                   **(A) amount and dosage form and strength;**

17                   **(B) date of purchase;**

18                   **(C) sales invoice number;**

19                   **(D) container size;**

20                   **(E) number of containers;**

21                   **(F) lot number; and**

22                   **(G) proprietary and established name;**

23           **of the legend drug.**

24           **(3) The:**

25                   **(A) business name and address of each owner of the legend**  
 26                   **drug; and**

27                   **(B) legend drug's shipping information, including the name**  
 28                   **and address of the facility of each person certifying**  
 29                   **delivery or receipt of the legend drug.**

30           **(4) Information that states that the wholesale drug distributor**  
 31           **has acted with due diligence as required under this chapter**  
 32           **with respect to another wholesale drug distributor from which**  
 33           **the wholesale drug distributor purchased or may have**  
 34           **purchased the legend drug.**

35           **(5) A certification from the designated representative of the**  
 36           **wholesale drug distributor that the information contained in**  
 37           **the document is true and accurate under penalty of perjury.**

38           SECTION 35. IC 25-26-14-9 IS AMENDED TO READ AS



1       FOLLOWS [EFFECTIVE JULY 1, 2005]: Sec. 9. As used in this  
 2       chapter, "person" means an individual, a partnership, a business firm,  
 3       a limited liability company, or a corporation, **or another entity,**  
 4       **including a governmental entity.**

5       SECTION 36. IC 25-26-14-9.2 IS ADDED TO THE INDIANA  
 6       CODE AS A **NEW** SECTION TO READ AS FOLLOWS  
 7       [EFFECTIVE JULY 1, 2005]: **Sec. 9.2. As used in this chapter,**  
 8       **"practitioner" has the meaning set forth in IC 16-42-19-5.**

9       SECTION 37. IC 25-26-14-9.3 IS ADDED TO THE INDIANA  
 10       CODE AS A **NEW** SECTION TO READ AS FOLLOWS  
 11       [EFFECTIVE JULY 1, 2005]: **Sec. 9.3. As used in this chapter,**  
 12       **"repackage" means changing the container, wrapper, quantity, or**  
 13       **labeling of a legend drug to further the distribution of the legend**  
 14       **drug.**

15       SECTION 38. IC 25-26-14-10.5 IS ADDED TO THE INDIANA  
 16       CODE AS A **NEW** SECTION TO READ AS FOLLOWS  
 17       [EFFECTIVE JULY 1, 2005]: **Sec. 10.5. As used in this chapter,**  
 18       **"specified list of susceptible products" means a specific list of**  
 19       **legend drugs established by the board, the board's agent, or a third**  
 20       **party approved by the board, as:**

- 21               **(1) susceptible to adulteration, counterfeiting, or diversion;**  
 22               **and**  
 23               **(2) posing the potential for a particular public health risk.**

24       SECTION 39. IC 25-26-14-11 IS AMENDED TO READ AS  
 25       FOLLOWS [EFFECTIVE JULY 1, 2005]: Sec. 11. As used in this  
 26       chapter, "wholesale distribution" means ~~distribution of~~ **to distribute**  
 27       legend drugs to persons other than a consumer or patient. The term does  
 28       not include:

- 29               (1) a sale **or transfer** between a division, a subsidiary, a parent,  
 30               an affiliated, or a related company under the common ownership  
 31               and control of a corporate entity;  
 32               (2) the purchase or acquisition by a hospital or other health care  
 33               entity that is a member of a group purchasing organization of a  
 34               drug for the hospital's or health care entity's own use from the  
 35               group purchasing organization or from other hospitals or health  
 36               care entities that are members of the organization;  
 37               (3) the sale of a drug by a charitable organization described in  
 38               Section 501(c)(3) of the Internal Revenue Code, to a nonprofit

- 1 affiliate of the organization to the extent otherwise permitted by  
 2 law;
- 3 (4) the sale of a drug among hospitals or other health care entities  
 4 that are under common control;
- 5 (5) the sale of a drug for emergency medical reasons, including  
 6 transfers of legend drugs by a retail pharmacy to another retail  
 7 pharmacy to alleviate a temporary shortage, if the gross dollar  
 8 value of the transfers does not exceed five percent (5%) of the  
 9 total legend drug sales revenue of either the transferor or  
 10 transferee pharmacy during any twelve (12) consecutive month  
 11 period;
- 12 (6) the sale of a drug or the dispensing of a drug pursuant to a  
 13 prescription;
- 14 (7) the distribution of drug samples by manufacturers'  
 15 representatives or distributors' representatives;
- 16 (8) the sale of blood and blood components intended for  
 17 transfusion;
- 18 (9) the sale of a drug by a retail pharmacy to a practitioner (as  
 19 defined in IC 25-26-13-2) for office use, if the gross dollar value  
 20 of the transfers does not exceed five percent (5%) of the retail  
 21 pharmacy's total legend drug sales during any twelve (12)  
 22 consecutive months; ~~or~~
- 23 (10) the sale of a drug by a retail pharmacy that is ending its  
 24 business and liquidating its inventory to another retail pharmacy;
- 25 **(11) drug returns by a hospital, health care entity, or**  
 26 **charitable institution conducted under 21 CFR 203.23;**
- 27 **(12) the sale of minimal quantities of drugs by retail**  
 28 **pharmacies to licensed practitioners for office use; or**
- 29 **(13) the distribution of prescription drugs by the original**  
 30 **manufacturer of the finished form of the prescription drug or**  
 31 **the distribution of the prescription drugs by a co-promoting**  
 32 **partner of the original manufacturer of the finished form of**  
 33 **the prescription drug.**
- 34 SECTION 40. IC 25-26-14-12 IS AMENDED TO READ AS  
 35 FOLLOWS [EFFECTIVE JULY 1, 2005]: Sec. 12. As used in this  
 36 chapter, "wholesale drug distributor" means a person engaged in  
 37 wholesale distribution of legend drugs, including:
- 38 (1) manufacturers;

- 1 (2) repackers;
- 2 (3) own-label distributors;
- 3 (4) private-label distributors;
- 4 (5) jobbers;
- 5 (6) brokers;
- 6 (7) warehouses, including manufacturers' and distributors'
- 7 warehouses, chain drug warehouses, and wholesale drug
- 8 warehouses;
- 9 (8) independent wholesale drug traders; and
- 10 (9) retail and hospital pharmacies that conduct wholesale
- 11 distributions.

12 The term does not include a common carrier or person hired solely to  
 13 transport prescription drugs.

14 SECTION 41. IC 25-26-14-14 IS AMENDED TO READ AS  
 15 FOLLOWS [EFFECTIVE JULY 1, 2005]: Sec. 14. (a) ~~After September~~  
 16 ~~14, 1992,~~ A person may not engage in wholesale distributions of legend  
 17 drugs without: ~~having~~

18 **(1) obtaining and maintaining accreditation or certification**  
 19 **from an accreditation body approved by the board under**  
 20 **subsection (g);**

21 **(2) obtaining and maintaining** a license ~~from~~ **issued by** the  
 22 board; and

23 **(3) paying** any reasonable fee required by the board.

24 (b) The board may not issue or renew the license of a wholesale  
 25 drug distributor that does not comply with this chapter.

26 (c) The board ~~may~~ **shall** require a separate license for  
 27 ~~(1) each facility directly or indirectly owned or operated by the~~  
 28 ~~same business in Indiana; or~~  
 29 ~~(2) a parent entity with divisions, subsidiaries, or affiliate~~  
 30 ~~companies in Indiana when operations are conducted at more than~~  
 31 ~~one (1) location and there exists joint ownership and control~~  
 32 ~~among all the entities; or location where wholesale distribution~~  
 33 **operations are conducted.**

34 (d) An agent or employee of any licensed wholesale drug distributor  
 35 does not need a license and may lawfully possess pharmaceutical drugs  
 36 when acting in the usual course of business or employment.

37 (e) The issuance of a license under this chapter does not affect tax  
 38 liability imposed by the department of state revenue or the department

1 of local government finance on any wholesale drug distributor.

2 (f) The board may adopt rules that permit out-of-state wholesale  
3 drug distributors to obtain a license on the basis of reciprocity if:

4 (1) an out-of-state wholesale drug distributor possesses a valid  
5 license granted by another state and the legal standards for  
6 licensure in the other state are comparable to the standards under  
7 this chapter; and

8 (2) the other state extends reciprocity to wholesale drug  
9 distributors licensed in Indiana.

10 **However, if the requirements for licensure under this chapter are**  
11 **more restrictive than the standards of the other state, the**  
12 **out-of-state wholesale drug distributor must comply with the**  
13 **additional requirements of this chapter to obtain a license under**  
14 **this chapter.**

15 (g) The board shall adopt rules under IC 4-22-2 to approve an  
16 accreditation body to:

17 (1) evaluate a wholesale drug distributor's operations to  
18 determine compliance with:

19 (A) professional standards;

20 (B) this chapter; and

21 (C) any other applicable law; and

22 (2) perform inspections of each facility and location where  
23 wholesale distribution operations are conducted by the  
24 wholesale drug distributor.

25 SECTION 42. IC 25-26-14-14.5 IS ADDED TO THE INDIANA  
26 CODE AS A NEW SECTION TO READ AS FOLLOWS  
27 [EFFECTIVE JULY 1, 2005]: **Sec. 14.5. After June 30, 2006, a**  
28 **wholesale drug distributor may not accept or deliver a legend drug**  
29 **without a current, accompanying pedigree.**

30 SECTION 43. IC 25-26-14-15 IS AMENDED TO READ AS  
31 FOLLOWS [EFFECTIVE JULY 1, 2005]: Sec. 15. (a) The board shall  
32 require the following minimum information from each wholesale drug  
33 distributor as part of the license described in section 14 of this chapter  
34 and as part of any renewal of such license:

35 (1) The name, full business address, and telephone number of the  
36 licensee.

37 (2) All trade or business names used by the licensee.

38 (3) Addresses, telephone numbers, and the names of contact

- 1 persons for all facilities used by the licensee for the storage,  
 2 handling, and distribution of legend drugs.
- 3 (4) The type of ownership of operation.
- 4 (5) The name of each owner and operator of the licensee,  
 5 including:
- 6 (A) if an individual, the name, **address, Social Security**  
 7 **number, and date of birth** of the individual;
- 8 (B) if a partnership, the name, **address, Social Security**  
 9 **number, and date of birth** of each partner, and the name of  
 10 the partnership **and federal employer identification number**;
- 11 (C) if a corporation:
- 12 (i) the name, **address, Social Security number, date of**  
 13 **birth**, and title of each corporate officer and director;
- 14 (ii) the corporate names, ~~and~~ the name of the state of  
 15 incorporation, **the federal employer identification**  
 16 **number, and the name of the parent company, if any;**  
 17 **and**
- 18 (iii) **the name, address, and Social Security number of**  
 19 **each shareholder owning ten percent (10%) or more of**  
 20 **the voting stock of the corporation, unless the stock is**  
 21 **traded on a major stock exchange and not traded over**  
 22 **the counter;**
- 23 (D) if a limited liability company, the name of each manager  
 24 and member, the name **and federal identification number** of  
 25 the limited liability company, and the name of the state where  
 26 organized; and
- 27 (E) if a sole proprietorship, the full name, **address, Social**  
 28 **Security number, and date of birth** of the sole proprietor and  
 29 the name **and federal employer identification number** of the  
 30 business entity.
- 31 (6) The name, **address, and telephone number** of the ~~person~~  
 32 ~~designated by the licensee as responsible for the operation~~  
 33 **representative of the facilities: each facility.**
- 34 (7) **Additional information concerning record keeping**  
 35 **required under this chapter.**
- 36 (b) **The board shall require a wholesale drug distributor to post**  
 37 **a surety bond of at least one hundred thousand dollars (\$100,000),**  
 38 **or an equivalent means of security acceptable to the board,**

1 including insurance, an irrevocable letter of credit, or funds  
 2 deposited in a trust account or financial institution, to secure  
 3 payment of any administrative penalties that may be imposed by  
 4 the board and any fees and costs that may be incurred by the board  
 5 and that:

- 6 (1) are related to a license held by the wholesale drug  
 7 distributor;
- 8 (2) are authorized under Indiana law; and
- 9 (3) the wholesale drug distributor fails to pay less than thirty  
 10 (30) days after the penalties, fees, or costs become final.

11 However, a separate surety bond or an equivalent means of  
 12 security is not required for a separate location or a company of the  
 13 wholesale drug distributor.

14 (c) The board may make a claim against a bond or security  
 15 posted under subsection (b) within one (1) year after the wholesale  
 16 drug distributor's license is no longer valid or sixty (60) days after  
 17 the conclusion of:

- 18 (1) an administrative or legal proceeding before or on behalf  
 19 of the board that involves the wholesale drug distributor and  
 20 results in penalties, fees, or costs described in subsection (b);  
 21 or
- 22 (2) an appeal of a proceeding described in subdivision (1);

23 whichever occurs later.

24 (d) The board shall inspect each facility where wholesale  
 25 distribution operations are conducted before initial licensure and  
 26 periodically thereafter in accordance with a schedule determined  
 27 by the board, but at least one (1) time in each three (3) year period.

28 (e) A wholesale drug distributor must publicly display or have  
 29 readily available all licenses and the most recent inspection report  
 30 administered by the board.

31 ~~(b)~~ (f) A material change in any information in ~~subsection (a)~~ of this  
 32 section must be submitted to the board at the time of license renewal or  
 33 within thirty (30) days from the date of the change, whichever occurs  
 34 first.

35 SECTION 44. IC 25-26-14-15.5 IS ADDED TO THE INDIANA  
 36 CODE AS A NEW SECTION TO READ AS FOLLOWS  
 37 [EFFECTIVE JULY 1, 2005]: Sec. 15.5. (a) A wholesale drug  
 38 distributor that is an authorized distributor of a manufacturer is

1 **not considered to be an authorized distributor of the manufacturer**  
2 **under this chapter unless:**

3 **(1) the manufacturer files the manufacturer's monthly**  
4 **updated list of authorized distributors with the board;**

5 **(2) the list is available from the manufacturer upon request or**  
6 **on the Internet; and**

7 **(3) the manufacturer notifies the board of any change to the**  
8 **list within ten (10) days after the change.**

9 **(b) The board shall make available on the board's Internet web**  
10 **site a manufacturer's list of authorized distributors filed as**  
11 **described in subsection (a).**

12 SECTION 45. IC 25-26-14-16 IS AMENDED TO READ AS  
13 FOLLOWS [EFFECTIVE JULY 1, 2005]: Sec. 16. **(a)** In reviewing,  
14 **for purposes of licensure or renewal of a license under this chapter,**  
15 the qualifications of persons who engage in wholesale distribution of  
16 legend drugs within in Indiana, the board shall consider the following  
17 factors:

18 (1) ~~A conviction of the applicant relating to drug samples;~~  
19 ~~wholesale or retail drug distribution; or distribution of controlled~~  
20 ~~substances: finding by the board that the applicant has:~~

21 **(A) violated a law; or**

22 **(B) been disciplined by a regulatory agency for violating a**  
23 **law;**

24 **related to drug distribution in any state.**

25 (2) A ~~felony criminal~~ conviction of the applicant.

26 (3) The applicant's past experience in the manufacture or  
27 distribution of legend drugs, including controlled substances.

28 (4) The furnishing by the applicant of false or fraudulent material  
29 in any application made in connection with drug manufacturing or  
30 distribution.

31 (5) Suspension or revocation **of any license held by the**  
32 **applicant or the applicant's owner or the imposition of**  
33 **sanctions against the applicant or the applicant's owner** by the  
34 federal or a state **or local** government ~~of any license held by the~~  
35 ~~applicant~~ for the manufacture or distribution of any drugs,  
36 including controlled substances.

37 (6) Compliance with licensing requirements under previously  
38 granted licenses.

1 (7) Compliance with requirements to maintain and make available  
 2 to the board or to federal, state, or local law enforcement officials  
 3 those records required under this chapter.

4 (8) Any other factors or qualifications the board considers relevant  
 5 to the public health and safety, including whether the granting of  
 6 the license would not be in the public interest.

7 **(b) In reviewing an application for licensure or renewal of a**  
 8 **license under this chapter, the board shall consider the results of a**  
 9 **national criminal history background check (as defined in**  
 10 **IC 10-13-3-12) for:**

11 (1) the applicant;

12 (2) all personnel involved in the operations of the wholesale  
 13 drug distributor;

14 (3) the most senior individual responsible for facility  
 15 operations, purchasing, and inventory control, and the  
 16 individual to whom the senior individual reports;

17 (4) company officers;

18 (5) key management personnel;

19 (6) principals; and

20 (7) owners with at least a ten percent (10%) interest in the  
 21 wholesale drug distributor, if the wholesale drug distributor  
 22 is a nonpublicly held company.

23 **The national criminal history background check must be**  
 24 **conducted at the applicant's expense and must include all states of**  
 25 **residence since the applicant became eighteen (18) years of age.**

26 **(c) An applicant shall provide and attest to:**

27 (1) an affirmation that the applicant has not been involved in  
 28 or convicted of any criminal or prohibited acts; or

29 (2) a statement providing a complete disclosure of the  
 30 applicant's past criminal convictions and violations of state  
 31 and federal laws;

32 **regarding drugs.**

33 SECTION 46. IC 25-26-14-16.5 IS ADDED TO THE INDIANA  
 34 CODE AS A NEW SECTION TO READ AS FOLLOWS  
 35 [EFFECTIVE JULY 1, 2005]: **Sec. 16.5. (a) A wholesale drug**  
 36 **distributor shall designate in writing on a form prescribed by the**  
 37 **board a designated representative for each of the wholesale drug**  
 38 **distributor's facilities licensed under this chapter.**



1           **(b) A designated representative shall submit to the board an**  
2 **application prescribed by the board and provide to the board the**  
3 **following:**

4           **(1) A set of the designated representative's fingerprints, under**  
5 **procedures specified by the board and according to**  
6 **requirements of the state police department under**  
7 **IC 10-13-3-38.5, with the payment of the amount equal to the**  
8 **costs of a national criminal history background check (as**  
9 **defined in IC 10-13-3-12) of the designated representative to**  
10 **be obtained by the state police department.**

11           **(2) The date and place of birth of the designated**  
12 **representative.**

13           **(3) A list of the occupations, positions of employment, and**  
14 **offices held by the designated representative during the**  
15 **immediately preceding seven (7) years, including the principal**  
16 **business and address of the organization with which the**  
17 **occupation, position, or office was associated.**

18           **(4) A statement concerning whether the designated**  
19 **representative, during the immediately preceding seven (7)**  
20 **years, has been temporarily or permanently enjoined by a**  
21 **court from violating a state or federal law regulating the**  
22 **possession, control, or distribution of drugs, including details**  
23 **of related events.**

24           **(5) A description of any involvement by the designated**  
25 **representative with a business that:**

26           **(A) manufactured, administered, prescribed, distributed,**  
27 **or stored drugs; and**

28           **(B) was named as a party in a lawsuit;**

29 **during the immediately preceding seven (7) years, including**  
30 **investments other than the ownership of stock in a publicly**  
31 **traded company or mutual fund.**

32           **(6) A description of any criminal offense of which the**  
33 **designated representative has been convicted, regardless of**  
34 **whether adjudication of guilt was withheld or whether the**  
35 **designated representative pleaded nolo contendere. If the**  
36 **designated representative indicates that a criminal conviction**  
37 **is under appeal, the designated representative shall submit to**  
38 **the board:**

- 1                   **(A) a copy of the notice of appeal; and**
- 2                   **(B) a copy of the final written order of disposition.**
- 3                   **(7) A photograph of the designated representative taken**
- 4                   **within the immediately preceding thirty (30) days under**
- 5                   **procedures specified by the board.**
- 6                   **(8) A list of the name, address, occupation, and date and place**
- 7                   **of birth of each member of the designated representative's**
- 8                   **immediate family, including the designated representative's**
- 9                   **spouse, children, parents, and siblings, and the spouses of the**
- 10                   **designated representative's children and siblings. Information**
- 11                   **collected under this subdivision is confidential.**
- 12                   **(9) Any other information required by the board.**
- 13                   **(c) A designated representative must have at least two (2) years**
- 14                   **of verifiable full-time managerial or supervisory experience in a**
- 15                   **pharmacy or with a wholesale drug distributor licensed under this**
- 16                   **chapter or in another state. The designated representative's**
- 17                   **responsibilities must have included record keeping, storage, and**
- 18                   **shipment of legend drugs.**
- 19                   **(d) A designated representative shall not serve as the designated**
- 20                   **representative for more than one (1) wholesale drug distributor**
- 21                   **facility at any one (1) time.**
- 22                   **(e) A designated representative shall be actively involved and**
- 23                   **aware of the actual daily operations of the wholesale drug**
- 24                   **distributor as follows:**
- 25                   **(1) Be employed full time in a managerial position by the**
- 26                   **wholesale drug distributor.**
- 27                   **(2) Be physically present at the wholesale drug distributor's**
- 28                   **facility during normal business hours, except when absent due**
- 29                   **to illness, family illness or death, scheduled vacation, or**
- 30                   **another authorized absence.**
- 31                   **(3) Be aware of and knowledgeable about all policies and**
- 32                   **procedures pertaining to the operations of the wholesale drug**
- 33                   **distributor.**
- 34                   **(f) A designated representative must complete continuing**
- 35                   **education programs specified by the board regarding state and**
- 36                   **federal law relevant to the distribution, handling, and storage of**
- 37                   **legend drugs.**

38                   SECTION 47. IC 25-26-14-16.6 IS ADDED TO THE INDIANA

1 CODE AS A NEW SECTION TO READ AS FOLLOWS  
 2 [EFFECTIVE JULY 1, 2005]: **Sec. 16.6. (a) A wholesale drug**  
 3 **distributor that:**

- 4 (1) **is licensed under this chapter;**  
 5 (2) **is located outside Indiana; and**  
 6 (3) **distributes legend drugs in Indiana;**

7 **shall designate an agent in Indiana for service of process.**

8 (b) **A wholesale drug distributor that does not designate an**  
 9 **agent under subsection (a) is considered to have designated the**  
 10 **secretary of state to be the wholesale drug distributor's true and**  
 11 **lawful attorney, upon whom legal process may be served in an**  
 12 **action or a proceeding against the wholesale drug distributor**  
 13 **arising from the wholesale drug distributor's wholesale distribution**  
 14 **operations.**

15 (c) **The board shall mail a copy of any service of process to a**  
 16 **wholesale drug distributor by certified mail, return receipt**  
 17 **requested, postage prepaid, at the address designated by the**  
 18 **wholesale drug distributor on the application for licensure**  
 19 **submitted under this chapter.**

20 (d) **Service of process on the secretary of state is sufficient in an**  
 21 **action or a proceeding against a wholesale drug distributor that is**  
 22 **not licensed under this chapter.**

23 SECTION 48. IC 25-26-14-17 IS AMENDED TO READ AS  
 24 FOLLOWS [EFFECTIVE JULY 1, 2005]: **Sec. 17. As a condition for**  
 25 **receiving and retaining any a wholesale drug distributor license issued**  
 26 **under to this chapter, each an applicant must satisfy the board that the**  
 27 **applicant has and will continuously maintain the following:**

- 28 (1) **Acceptable storage and handling conditions and facilities**  
 29 **standards for each facility at which legend drugs are received,**  
 30 **stored, warehoused, handled, held, offered, marketed, or**  
 31 **displayed, or from which legend drugs are transported,**  
 32 **including:**

33 (A) **suitable construction of the facility and appropriate**  
 34 **monitoring equipment to ensure that legend drugs in the**  
 35 **facility are maintained in accordance with labeling or in**  
 36 **compliance with official compendium standards;**

37 (B) **suitable size and construction to facilitate cleaning,**  
 38 **maintenance, and proper wholesale distribution**

- 1                   **operations;**
- 2                   **(C) adequate storage areas to provide appropriate lighting,**
- 3                   **ventilation, temperature, sanitation, humidity, space,**
- 4                   **equipment, and security conditions;**
- 5                   **(D) a quarantine area for separate storage of legend drugs**
- 6                   **that are outdated, damaged, deteriorated, misbranded,**
- 7                   **adulterated, counterfeit, suspected counterfeit, otherwise**
- 8                   **unfit for distribution, or contained in immediate or sealed**
- 9                   **secondary containers that have been opened;**
- 10                  **(E) maintenance of the facility in a clean and orderly**
- 11                  **condition;**
- 12                  **(F) maintenance of the facility in a commercial,**
- 13                  **nonresidential building; and**
- 14                  **(G) freedom of the facility from infestation.**
- 15                  **(2) Security of each facility from unauthorized entry as**
- 16                  **follows:**
- 17                   **(A) Entry into areas where legend drugs are held is limited**
- 18                   **to authorized personnel.**
- 19                   **(B) Each facility is equipped with a security system that**
- 20                   **includes:**
- 21                    ~~(A)~~ **(i) an after hours central alarm or a comparable entry**
- 22                    **detection capability;**
- 23                    ~~(B)~~ **(ii) restricted premises access;**
- 24                    ~~(C)~~ **(iii) adequate outside perimeter lighting; and**
- 25                    ~~(D)~~ **(iv) safeguards against theft and diversion, including**
- 26                    **employee theft and theft or diversion facilitated or hidden**
- 27                    **by tampering with computers or electronic records; and**
- 28                    **(v) a means of protecting the integrity and confidentiality**
- 29                    **of data and documents and of making the data and**
- 30                    **documents readily available to the board and other state**
- 31                    **and federal law enforcement officials.**
- 32                  **(3) A reasonable system of record keeping that as follows:**
- 33                    **(A) The system describes all the wholesale distributor's**
- 34                    **activities governed by this chapter for the two ~~(2)~~ three (3)**
- 35                    **year period after the disposition of each product and all**
- 36                    **records are maintained for at least three (3) years after**
- 37                    **disposition of the legend drug to which the record applies.**
- 38                    **(B) The system is reasonably accessible as determined by**

- 1 board rules in any inspection authorized by the board.
- 2 **(C) The system provides a means to establish and maintain**
- 3 **inventories and records of transactions regarding the**
- 4 **receipt and distribution or other disposition of all legend**
- 5 **drugs, including the following:**
- 6 **(i) For legend drugs manufactured by a manufacturer**
- 7 **for which the wholesale drug distributor is an authorized**
- 8 **distributor, a pedigree for each distributed legend drug**
- 9 **that is on the specified list of susceptible products or that**
- 10 **leaves the normal distribution chain of custody from the**
- 11 **manufacturer to a wholesale drug distributor, to a**
- 12 **pharmacy, and to the patient or the patient's agent.**
- 13 **(ii) For legend drugs manufactured by a manufacturer**
- 14 **for which the wholesale drug distributor is not an**
- 15 **authorized distributor, a pedigree for each distributed**
- 16 **legend drug.**
- 17 **(iii) After January 1, 2007, at the board's discretion, for**
- 18 **each legend drug received and distributed by the**
- 19 **wholesale drug distributor, an electronic pedigree**
- 20 **developed in accordance with standards and**
- 21 **requirements of the board to authenticate, track, and**
- 22 **trace legend drugs. The standards and requirements of**
- 23 **the board may indicate the information required to be**
- 24 **part of the electronic pedigree.**
- 25 **(iv) Dates of receipt and distribution or other disposition**
- 26 **of the legend drugs by the wholesale drug distributor.**
- 27 **(v) Availability for inspection and photocopying by any**
- 28 **authorized official of a local, state, or federal**
- 29 **governmental agency for three (3) years after the**
- 30 **creation date of the inventories and records.**
- 31 **(D) Onsite electronic inventories and records are**
- 32 **immediately available for inspection. Records kept at a**
- 33 **central location apart from the inspection site and not**
- 34 **electronically retrievable are available for inspection**
- 35 **within two (2) working days after a request by an**
- 36 **authorized official of a local, state, or federal governmental**
- 37 **agency.**
- 38 **(E) The system maintains an ongoing list of persons with**

- 1           whom the wholesale drug distributor does business.
- 2           **(F) The system provides for reporting counterfeit or**
- 3           **suspected counterfeit legend drugs or counterfeiting or**
- 4           **suspected counterfeiting activities to the board and federal**
- 5           **Food and Drug Administration.**
- 6           **(G) The system provides for mandatory reporting of**
- 7           **significant shortages or losses of legend drugs to the board**
- 8           **and federal Food and Drug Administration if diversion is**
- 9           **known or suspected.**
- 10          (4) Written policies and procedures **to which the wholesale drug**
- 11          **distributor adheres for the receipt, security, storage,**
- 12          **inventory, transport, shipping, and distribution of legend**
- 13          **drugs, and** that assure reasonable wholesale distributor
- 14          preparation for, protection against, and handling of any facility
- 15          security or operation problems, including **the following:**
- 16                (A) ~~those~~ **Facility security or operation problems** caused by
- 17                natural disaster or government emergency.
- 18                (B) **Correction of inventory inaccuracies.** ~~or~~
- 19                (C) **Product shipping and receiving problems.**
- 20                ~~(D)~~ **(D) Quarantine and return to the manufacturer or**
- 21                **destruction in accordance with state and federal law of all**
- 22                outdated ~~product~~ **products and outdated or expired legend**
- 23                **drugs, including appropriate documentation and**
- 24                **witnessing.**
- 25                ~~(E)~~ **(E) Appropriate disposition of returned goods.** ~~and~~
- 26                ~~(F)~~ **(F) Product recalls.**
- 27                **(G) Identifying, recording, and reporting losses or thefts.**
- 28                **(H) Implementation and maintenance of a continuous**
- 29                **quality improvement system.**
- 30                **(I) Recalls and withdrawals of legend drugs due to:**
- 31                    **(i) an action initiated by the federal Food and Drug**
- 32                    **Administration or another federal, state, or local**
- 33                    **governmental agency;**
- 34                    **(ii) a volunteer action by the manufacturer to remove**
- 35                    **defective or potentially defective legend drugs from the**
- 36                    **market; or**
- 37                    **(iii) an action undertaken to promote public health and**
- 38                    **safety by replacing existing merchandise with an**

- 1           **improved product or a new package design.**
- 2           **(J) Disposition and destruction of containers, labels, and**  
 3           **packaging to ensure that the containers, labels, and**  
 4           **packaging are not used in counterfeiting activities,**  
 5           **including necessary documentation and witnessing in**  
 6           **accordance with state and federal law.**
- 7           **(K) Investigation of discrepancies in the inventory**  
 8           **involving counterfeit, suspected counterfeit, contraband, or**  
 9           **suspected contraband legend drugs and reporting of**  
 10           **discrepancies within three (3) business days to the board**  
 11           **and any other appropriate state or federal governmental**  
 12           **agency.**
- 13           **(L) Reporting of criminal or suspected criminal activities**  
 14           **involving the inventory of legend drugs to the board within**  
 15           **three (3) business days.**
- 16           **(M) Conducting for cause authentication and random**  
 17           **authentication as required under sections 17.2, 17.3, and**  
 18           **17.8 of this chapter.**
- 19           **(5) Written policies and procedures and sufficient inspection**  
 20           **procedures for all incoming and outgoing product shipments,**  
 21           **including the following:**
- 22                   **(A) Upon receipt, visual examination of each shipping**  
 23                   **container in a manner adequate to identify the legend**  
 24                   **drugs in the container and to determine whether the legend**  
 25                   **drugs may be outdated, adulterated, misbranded,**  
 26                   **contaminated, contraband, counterfeit, suspected**  
 27                   **counterfeit, damaged, or otherwise unfit for distribution.**
- 28                   **(B) Upon receipt, review of records by the wholesale drug**  
 29                   **distributor for the acquisition of legend drugs for accuracy**  
 30                   **and completeness, considering the:**
- 31                           **(i) total facts and circumstances surrounding each**  
 32                           **transaction involving the legend drugs; and**  
 33                           **(ii) wholesale drug distributors involved.**
- 34                   **(C) Quarantine of a legend drug considered to be outdated,**  
 35                   **adulterated, misbranded, contaminated, contraband,**  
 36                   **counterfeit, suspected counterfeit, damaged, or otherwise**  
 37                   **unfit for distribution until:**
- 38                           **(i) examination and a determination that the legend drug**

- 1           **is not outdated, adulterated, misbranded, contaminated,**
- 2           **contraband, counterfeit, damaged, or otherwise unfit for**
- 3           **distribution; or**
- 4           **(ii) the legend drug is destroyed or returned to the**
- 5           **manufacturer or wholesale drug distributor from which**
- 6           **the legend drug was acquired.**
- 7           **(D) Written policies and procedures to ensure that a legend**
- 8           **drug that was:**
- 9           **(i) ordered in error or in excess of need by the wholesale**
- 10           **drug distributor;**
- 11           **(ii) identified within three (3) business days after receipt**
- 12           **as ordered in error or in excess of need; and**
- 13           **(iii) maintained such that the legend drug's integrity has**
- 14           **not been compromised;**
- 15           **may be returned to the manufacturer or wholesale drug**
- 16           **distributor from which the legend drug was acquired if the**
- 17           **appropriate documentation is completed and necessary**
- 18           **notations are made to a required pedigree.**
- 19           **(E) Written policies and procedures to ensure that if the**
- 20           **wholesale drug distributor determines that a legend drug**
- 21           **is adulterated, misbranded, counterfeit, or suspected**
- 22           **counterfeit, the wholesale drug distributor provides notice**
- 23           **of the adulteration, misbranding, counterfeiting, or**
- 24           **suspected counterfeiting to the board, the federal Food and**
- 25           **Drug Administration, and the manufacturer or wholesale**
- 26           **drug distributor from which the legend drug was acquired**
- 27           **within three (3) business days.**
- 28           **(F) Written policies and procedures to ensure that if the**
- 29           **immediate or sealed outer or secondary container or**
- 30           **labeling of a legend drug is adulterated, misbranded,**
- 31           **counterfeit, or suspected counterfeit, the wholesale drug**
- 32           **distributor:**
- 33           **(i) quarantines the legend drug until the legend drug is**
- 34           **destroyed or returned to the manufacturer or wholesale**
- 35           **drug distributor from which the legend drug was**
- 36           **acquired; and**
- 37           **(ii) provides notice of the adulteration, misbranding,**
- 38           **counterfeiting, or suspected counterfeiting to the board,**



1           the federal Food and Drug Administration, and the  
2           manufacturer or wholesale drug distributor from which  
3           the legend drug was acquired within three (3) business  
4           days.

5           **(G) Written policies and procedures to ensure that a legend**  
6           **drug that has been opened or used, but is not adulterated,**  
7           **misbranded, counterfeit, or suspected counterfeit, is**  
8           **identified as such and quarantined until the legend drug is**  
9           **destroyed or returned to the manufacturer or wholesale**  
10          **drug distributor from which the legend drug was acquired.**

11          **(H) Written policies and procedures to ensure that:**

12            **(i) a legend drug that will be returned to a manufacturer**  
13            **or wholesale drug distributor is kept under proper**  
14            **conditions for storage, handling, transport, and shipment**  
15            **before the return; and**

16            **(ii) documentation showing that proper conditions were**  
17            **maintained is provided to the manufacturer or wholesale**  
18            **drug distributor to which the legend drug is returned.**

19          **(I) Inspection of each outgoing shipment for identity of the**  
20          **legend drugs and to ensure that the legend drugs have not**  
21          **been damaged in storage or held under improper**  
22          **conditions.**

23          **(J) Written policies and procedures to ensure that if**  
24          **conditions under which a legend drug has been returned to**  
25          **the wholesale drug distributor cast doubt on the legend**  
26          **drug's safety, identity, strength, quality, or purity, the**  
27          **legend drug is destroyed or returned to the manufacturer**  
28          **or wholesale drug distributor from which the legend drug**  
29          **was acquired unless examination, testing, or other**  
30          **investigation proves that the legend drug meets**  
31          **appropriate standards of safety, identity, strength, quality,**  
32          **and purity. In determining whether the conditions under**  
33          **which a legend drug has been returned cast doubt on the**  
34          **legend drug's safety, identity, strength, quality, or purity,**  
35          **the wholesale drug distributor considers the conditions**  
36          **under which the legend drug has been held, stored, or**  
37          **shipped before or during the legend drug's return and the**  
38          **condition of the legend drug and the legend drug's**

- 1 container, carton, or labeling upon receipt of the returned  
2 legend drug.
- 3 **(K) Written policies and procedures to ensure that**  
4 **contraband, counterfeit, or suspected counterfeit legend**  
5 **drugs, other evidence of criminal activity, and**  
6 **accompanying documentation are retained until a**  
7 **disposition is authorized by the board and the federal Food**  
8 **and Drug Administration.**
- 9 **(L) Written policies and procedures to ensure that any**  
10 **shipping, immediate, or sealed outer or secondary**  
11 **container or labeling, and accompanying documentation,**  
12 **suspected of or determined to be counterfeit or fraudulent,**  
13 **are retained until a disposition is authorized by the board**  
14 **and federal Food and Drug Administration.**
- 15 (6) Operations in compliance with all federal legal requirements  
16 applicable to wholesale drug distribution.
- 17 **(7) Written policies and procedures to provide for the secure**  
18 **and confidential storage of information with restricted access**  
19 **and to protect the integrity and confidentiality of the**  
20 **information.**
- 21 **(8) A pedigree as required under this chapter, including an**  
22 **electronic pedigree developed in accordance with standards**  
23 **and requirements of the board under subdivision (3)(C)(iii).**
- 24 **(9) Appropriate inventory management and control systems**  
25 **to:**
- 26 **(A) prevent; and**
- 27 **(B) allow detection and documentation of;**
- 28 **theft, counterfeiting, or diversion of legend drugs.**
- 29 **(10) If the wholesale drug distributor is involved in the**  
30 **distribution of controlled substances, registration with the**  
31 **federal Drug Enforcement Administration and board and**  
32 **compliance with all laws related to the storage, handling,**  
33 **transport, shipment, and distribution of controlled substances.**
- 34 **(11) Isolation of controlled substances from noncontrolled**  
35 **substances and storage of the controlled substances in a secure**  
36 **area in accordance with federal Drug Enforcement**  
37 **Administration security requirements and standards.**
- 38 **(12) Technology and equipment that allow the wholesale drug**

1 distributor to authenticate, track, and trace legend drugs. The  
 2 technology and equipment meets standards set by the board  
 3 and is used as required by the board to conduct for cause and  
 4 random tracking, tracing, and authentication of legend drugs.

5 (13) Employment, training, and documentation of the training  
 6 concerning the proper use of the technology and equipment  
 7 required under subdivision (12).

8 (14) Packaging operations in accordance with an official  
 9 compendium allowing the identification of a compromise in  
 10 the integrity of the legend drugs due to tampering or adverse  
 11 storage conditions.

12 SECTION 49. IC 25-26-14-17.2 IS ADDED TO THE INDIANA  
 13 CODE AS A NEW SECTION TO READ AS FOLLOWS  
 14 [EFFECTIVE JULY 1, 2005]: Sec. 17.2. (a) A wholesale drug  
 15 distributor that purchases legend drugs from another wholesale  
 16 drug distributor and has reason to believe that a legend drug  
 17 purchased from the other wholesale drug distributor is counterfeit,  
 18 suspected counterfeit, misbranded, or adulterated shall conduct a  
 19 for cause authentication of each distribution of the legend drug  
 20 back to the manufacturer.

21 (b) A wholesale drug distributor that has engaged in the  
 22 distribution of a legend drug for which a purchasing wholesale  
 23 drug distributor conducts a for cause authentication under  
 24 subsection (a) shall provide, upon request, detailed information  
 25 regarding the distribution of the legend drug, including the:

- 26 (1) date of purchase of the legend drug;  
 27 (2) lot number of the legend drug;  
 28 (3) sales invoice number of the legend drug; and  
 29 (4) contact information, including name, address, telephone  
 30 number, and electronic mail address of the wholesale drug  
 31 distributor that sold the legend drug.

32 (c) If a wholesale drug distributor conducts a for cause  
 33 authentication under subsection (a) and is unable to authenticate  
 34 each distribution of the legend drug, the wholesale drug distributor  
 35 shall quarantine the legend drug and report the circumstances to  
 36 the board and the federal Food and Drug Administration not more  
 37 than ten (10) business days after completing the attempted  
 38 authentication.

1           **(d) If a wholesale drug distributor authenticates the distribution**  
 2 **of a legend drug back to the manufacturer under subsection (a), the**  
 3 **wholesale drug distributor shall maintain records of the**  
 4 **authentication for three (3) years and shall produce the records for**  
 5 **the board and the federal Food and Drug Administration upon**  
 6 **request.**

7           SECTION 50. IC 25-26-14-17.3 IS ADDED TO THE INDIANA  
 8 CODE AS A NEW SECTION TO READ AS FOLLOWS  
 9 [EFFECTIVE JULY 1, 2005]: **Sec. 17.3. (a) A wholesale drug**  
 10 **distributor that purchases legend drugs from another wholesale**  
 11 **drug distributor shall, at least annually, conduct a random**  
 12 **authentication of a required pedigree on at least ten percent (10%)**  
 13 **of sales units of wholesale distributions of legend drugs purchased**  
 14 **from other wholesale drug distributors.**

15           **(b) If a wholesale drug distributor purchases from another**  
 16 **wholesale drug distributor a legend drug that is on the specified list**  
 17 **of susceptible products, the wholesale drug distributor shall, at**  
 18 **least quarterly, conduct a random authentication of a required**  
 19 **pedigree on at least ninety percent (90%) of sales units of**  
 20 **distributions of legend drugs that are on the specified list of**  
 21 **susceptible products and that were purchased from other wholesale**  
 22 **drug distributors.**

23           **(c) A wholesale drug distributor from whom another wholesale**  
 24 **drug distributor purchases legend drugs shall cooperate with**  
 25 **random authentications of pedigrees described in this section and**  
 26 **provide requested information in a timely manner.**

27           **(d) If a wholesale drug distributor conducts a random**  
 28 **authentication under this section and is unable to authenticate each**  
 29 **distribution of the legend drug, the wholesale drug distributor shall**  
 30 **quarantine the legend drug and report the circumstances to the**  
 31 **board and the federal Food and Drug Administration not more**  
 32 **than ten (10) business days after completing the attempted**  
 33 **authentication.**

34           SECTION 51. IC 25-26-14-17.8 IS ADDED TO THE INDIANA  
 35 CODE AS A NEW SECTION TO READ AS FOLLOWS  
 36 [EFFECTIVE JULY 1, 2005]: **Sec. 17.8. (a) A wholesale drug**  
 37 **distributor licensed under this chapter that purchases legend drugs**  
 38 **from a wholesale drug distributor that is not licensed under this**

- 1 chapter shall act with due diligence as required under this section.
- 2 (b) Before the initial purchase of legend drugs from the
- 3 unlicensed wholesale drug distributor, the licensed wholesale drug
- 4 distributor shall obtain the following information from the
- 5 unlicensed wholesale drug distributor:
- 6 (1) A list of states in which the unlicensed wholesale drug
- 7 distributor is licensed.
- 8 (2) A list of states into which the unlicensed wholesale drug
- 9 distributor ships legend drugs.
- 10 (3) Copies of all state and federal regulatory licenses and
- 11 registrations held by the unlicensed wholesale drug
- 12 distributor.
- 13 (4) The unlicensed wholesale drug distributor's most recent
- 14 facility inspection reports.
- 15 (5) Information regarding general and product liability
- 16 insurance maintained by the unlicensed wholesale drug
- 17 distributor, including copies of relevant policies.
- 18 (6) A list of other names under which the unlicensed wholesale
- 19 drug distributor does business or has been previously known.
- 20 (7) A list of corporate officers and managerial employees of
- 21 the unlicensed wholesale drug distributor.
- 22 (8) A list of all owners of the unlicensed wholesale drug
- 23 distributor that own more than ten percent (10%) of the
- 24 unlicensed wholesale drug distributor, unless the unlicensed
- 25 wholesale drug distributor is publicly traded.
- 26 (9) A list of all disciplinary actions taken against the
- 27 unlicensed wholesale drug distributor by state and federal
- 28 agencies.
- 29 (10) A description, including the address, dimensions, and
- 30 other relevant information, of each facility used by the
- 31 unlicensed wholesale drug distributor for legend drug storage
- 32 and distribution.
- 33 (11) A description of legend drug import and export activities
- 34 of the unlicensed wholesale drug distributor.
- 35 (12) A description of the unlicensed wholesale drug
- 36 distributor's procedures to ensure compliance with this
- 37 chapter.
- 38 (13) A statement:

1           **(A) as to whether; and**  
2           **(B) of the identity of each manufacturer for which;**  
3           **the unlicensed wholesale drug distributor is an authorized**  
4           **distributor.**

5           **(c) Before the initial purchase of legend drugs from an**  
6           **unlicensed wholesale drug distributor, the licensed wholesale drug**  
7           **distributor shall:**

8               **(1) request that the board obtain and consider the results of a**  
9               **national criminal history background check (as defined in**  
10              **IC 10-13-3-12) through the state police department of all**  
11              **individuals associated with the unlicensed wholesale drug**  
12              **distributor as specified for licensure of a wholesale drug**  
13              **distributor under section 16(b) of this chapter; and**

14              **(2) verify the unlicensed wholesale drug distributor's status as**  
15              **an authorized distributor, if applicable.**

16           **(d) If an unlicensed wholesale drug distributor's facility has not**  
17           **been inspected by the board or the board's agent within three (3)**  
18           **years after a contemplated purchase described in subsection (a),**  
19           **the licensed wholesale drug distributor shall conduct an inspection**  
20           **of the unlicensed wholesale drug distributor's facility:**

21               **(1) before the initial purchase of legend drugs from the**  
22               **unlicensed wholesale drug distributor; and**

23               **(2) at least once every three (3) years unless the unlicensed**  
24               **wholesale drug distributor's facility has been inspected by the**  
25               **board, or the board's agent, during the same period;**

26           **to ensure compliance with applicable laws and regulations relating**  
27           **to the storage and handling of legend drugs. A third party may be**  
28           **engaged to conduct the site inspection on behalf of the licensed**  
29           **wholesale drug distributor.**

30           **(e) At least annually, a licensed wholesale drug distributor that**  
31           **purchases legend drugs from an unlicensed wholesale drug**  
32           **distributor shall ensure that the unlicensed wholesale drug**  
33           **distributor maintains a record keeping system that meets the**  
34           **requirements of section 17(3) of this chapter.**

35           **(f) If a licensed wholesale drug distributor that purchases legend**  
36           **drugs from an unlicensed wholesale drug distributor has reason to**  
37           **believe that a legend drug purchased from the unlicensed wholesale**  
38           **drug distributor is misbranded, adulterated, counterfeit, or**

1       **suspected counterfeit, the licensed wholesale drug distributor shall**  
 2       **conduct a for cause authentication of each distribution of the**  
 3       **legend drug back to the manufacturer.**

4       **(g) An unlicensed wholesale drug distributor that has engaged**  
 5       **in the distribution of a legend drug for which a licensed wholesale**  
 6       **drug distributor conducts a for cause authentication under**  
 7       **subsection (f) shall provide, upon request, detailed information**  
 8       **regarding the distribution of the legend drug, including the:**

9               **(1) date of purchase of the legend drug;**

10              **(2) lot number of the legend drug;**

11              **(3) sales invoice number of the legend drug; and**

12              **(4) contact information, including name, address, telephone**  
 13              **number, and any electronic mail address of the unlicensed**  
 14              **wholesale drug distributor that sold the legend drug.**

15       **(h) If a licensed wholesale drug distributor conducts a for cause**  
 16       **authentication under subsection (f) and is unable to authenticate**  
 17       **each distribution of the legend drug, the licensed wholesale drug**  
 18       **distributor shall quarantine the legend drug and report the**  
 19       **circumstances to the board and the federal Food and Drug**  
 20       **Administration within ten (10) business days after completing the**  
 21       **attempted authentication.**

22       **(i) If a licensed wholesale drug distributor authenticates the**  
 23       **distribution of a legend drug back to the manufacturer under**  
 24       **subsection (f), the licensed wholesale drug distributor shall**  
 25       **maintain records of the authentication for three (3) years and shall**  
 26       **provide the records to the board upon request.**

27       **(j) A licensed wholesale drug distributor that purchases legend**  
 28       **drugs from an unlicensed wholesale drug distributor shall, at least**  
 29       **annually, conduct random authentications of required pedigrees on**  
 30       **at least ten percent (10%) of sales units of distributions of legend**  
 31       **drugs that were purchased from unlicensed wholesale drug**  
 32       **distributors.**

33       **(k) A licensed wholesale drug distributor that has purchased a**  
 34       **legend drug that is on the specified list of susceptible products**  
 35       **shall, at least quarterly, conduct random authentications of**  
 36       **required pedigrees on at least ninety percent (90%) of sales units**  
 37       **of distributions of legend drugs that:**

38              **(1) are on the specified list of susceptible products; and**

1           **(2) were purchased from unlicensed wholesale drug**  
 2           **distributors.**

3           **(l) An unlicensed wholesale drug distributor from which a**  
 4           **licensed wholesale drug distributor has purchased legend drugs**  
 5           **shall cooperate with the random authentications of pedigrees under**  
 6           **this section and provide requested information in a timely manner.**

7           **(m) If a wholesale drug distributor conducts a random**  
 8           **authentication under subsection (j) or (k) and is unable to**  
 9           **authenticate each distribution of the legend drug, the wholesale**  
 10           **drug distributor shall quarantine the legend drug and report the**  
 11           **circumstances to the board and the federal Food and Drug**  
 12           **Administration not more than ten (10) business days after**  
 13           **completing the attempted authentication.**

14           SECTION 52. IC 25-26-14-17.9 IS ADDED TO THE INDIANA  
 15           CODE AS A NEW SECTION TO READ AS FOLLOWS  
 16           [EFFECTIVE JULY 1, 2005]: **Sec. 17.9. A wholesale drug**  
 17           **distributor licensed under this chapter may not use a trade name**  
 18           **or business name identical to a trade name or business name used**  
 19           **by another wholesale drug distributor licensed under this chapter.**

20           SECTION 53. IC 25-26-14-20 IS AMENDED TO READ AS  
 21           FOLLOWS [EFFECTIVE JULY 1, 2005]: **Sec. 20. (a) A person**  
 22           **employed in wholesale distribution must have appropriate education or**  
 23           **experience to assume responsibility for positions related to compliance**  
 24           **with licensing requirements.**

25           **(b) Before employing a person to be engaged in the operation**  
 26           **and handling of legend drugs, a wholesale drug distributor shall**  
 27           **request that the board obtain and consider the results of a national**  
 28           **criminal history background check (as defined in IC 10-13-3-12)**  
 29           **through the state police department for the person.**

30           SECTION 54. IC 25-26-14-21.5 IS ADDED TO THE INDIANA  
 31           CODE AS A NEW SECTION TO READ AS FOLLOWS  
 32           [EFFECTIVE JULY 1, 2005]: **Sec. 21.5. (a) A person may not**  
 33           **perform, cause the performance of, or aid the performance of the**  
 34           **following:**

35           **(1) The manufacture, repackaging, sale, delivery, holding, or**  
 36           **offering for sale of a legend drug that is adulterated,**  
 37           **misbranded, counterfeit, suspected counterfeit, or is otherwise**  
 38           **unfit for distribution.**



- 1           **(2) The adulteration, misbranding, or counterfeiting of a**  
2           **legend drug.**
- 3           **(3) The receipt of a legend drug that is adulterated,**  
4           **misbranded, stolen, obtained by fraud or deceit, counterfeit,**  
5           **or suspected counterfeit, and the delivery or proffered**  
6           **delivery of the legend drug for pay or otherwise.**
- 7           **(4) The alteration, mutilation, destruction, obliteration, or**  
8           **removal of the whole or a part of the labeling of a legend drug**  
9           **or the commission of another act with respect to a legend drug**  
10           **that results in the legend drug being misbranded.**
- 11           **(5) Forging, counterfeiting, simulating, or falsely representing**  
12           **a legend drug using a mark, stamp, tag, label, or other**  
13           **identification device without the authorization of the**  
14           **manufacturer.**
- 15           **(6) The purchase or receipt of a legend drug from a person**  
16           **that is not licensed to distribute legend drugs to the purchaser**  
17           **or recipient.**
- 18           **(7) The sale or transfer of a legend drug to a person that is not**  
19           **authorized under the law of the jurisdiction in which the**  
20           **person receives the legend drug to purchase or receive legend**  
21           **drugs from the person selling or transferring the legend drug.**
- 22           **(8) Failure to maintain or provide records as required under**  
23           **this chapter.**
- 24           **(9) Providing the board, a representative of the board, or a**  
25           **state or federal official with false or fraudulent records or**  
26           **making false or fraudulent statements regarding a matter**  
27           **related to this chapter.**
- 28           **(10) The wholesale distribution of a legend drug that was:**
- 29               **(A) purchased by a public or private hospital or other**  
30               **health care entity;**
- 31               **(B) donated or supplied at a reduced price to a charitable**  
32               **organization; or**
- 33               **(C) stolen or obtained by fraud or deceit.**
- 34           **(11) Obtaining or attempting to obtain a legend drug by**  
35           **fraud, deceit, misrepresentation, or engaging in fraud, deceit,**  
36           **or misrepresentation in the distribution of a legend drug.**
- 37           **(12) Failure to obtain, authenticate, or provide a required**  
38           **pedigree.**

1           **(13) The receipt of a legend drug through wholesale**  
 2           **distribution without first receiving a required pedigree**  
 3           **attested to as accurate and complete by the wholesale drug**  
 4           **distributor.**

5           **(14) Distributing a legend drug that was previously dispensed**  
 6           **by a retail pharmacy or distributed by a practitioner.**

7           **(15) Failure to report an act prohibited by this section.**

8           **(b) The board may impose the following sanctions if, after a**  
 9           **hearing under IC 4-21.5-3, the board finds that a person has**  
 10           **violated subsection (a):**

11           **(1) Revoke the wholesale drug distributor's license issued**  
 12           **under this chapter if the person is a wholesale drug**  
 13           **distributor.**

14           **(2) Assess a civil penalty against the person. A civil penalty**  
 15           **assessed under this subdivision may not be more than ten**  
 16           **thousand dollars (\$10,000) per violation.**

17           SECTION 55. IC 25-26-14-26 IS AMENDED TO READ AS  
 18           FOLLOWS [EFFECTIVE JULY 1, 2005]: Sec. 26. **(a)** A person ~~that~~  
 19           **who knowingly or intentionally** engages in the wholesale distribution  
 20           of a legend drug without a license issued under this chapter commits a  
 21           Class D felony.

22           **(b) A person who engages in the wholesale distribution of a**  
 23           **legend drug and:**

24           **(1) who, with intent to defraud or deceive:**

25           **(A) fails to obtain or deliver to another person a complete**  
 26           **and accurate required pedigree concerning a legend drug**  
 27           **before:**

28           **(i) obtaining the legend drug from another person; or**

29           **(ii) transferring the legend drug to another person; or**

30           **(B) falsely swears or certifies that the person has**  
 31           **authenticated any documents related to the wholesale**  
 32           **distribution of legend drugs;**

33           **(2) who knowingly or intentionally:**

34           **(A) destroys, alters, conceals, or fails to maintain a**  
 35           **complete and accurate required pedigree concerning a**  
 36           **legend drug in the person's possession;**

37           **(B) purchases or receives legend drugs from a person not**  
 38           **authorized to distribute legend drugs in wholesale**

- 1           **distribution;**
- 2           **(C) sells, barter, brokers, or transfers a legend drug to a**
- 3           **person not authorized to purchase the legend drug in the**
- 4           **jurisdiction in which the person receives the legend drug in**
- 5           **a wholesale distribution;**
- 6           **(D) forges, counterfeits, or falsely creates a pedigree;**
- 7           **(E) falsely represents a factual matter contained in a**
- 8           **pedigree; or**
- 9           **(F) fails to record material information required to be**
- 10          **recorded in a pedigree; or**
- 11          **(3) who:**
- 12           **(A) possesses a required pedigree concerning a legend**
- 13           **drug;**
- 14           **(B) knowingly or intentionally fails to authenticate the**
- 15           **matters contained in the pedigree as required; and**
- 16           **(C) distributes or attempts to further distribute the legend**
- 17           **drug;**
- 18          **commits a Class D felony.**

19           SECTION 56. IC 25-26-14-27 IS AMENDED TO READ AS  
 20           FOLLOWS [EFFECTIVE JULY 1, 2005]: Sec. 27. A wholesale drug  
 21           distributor that fails to comply with the conditions **and requirements**  
 22           described in section 17, **17.2, 17.3, 17.8, 17.9, or 20** of this chapter  
 23           commits a Class D felony."

24           Page 18, between lines 13 and 14, begin a new paragraph and insert:  
 25           "SECTION 60. IC 34-24-1-1 IS AMENDED TO READ AS  
 26           FOLLOWS [EFFECTIVE JULY 1, 2005]: Sec. 1. (a) The following  
 27           may be seized:

- 28           (1) All vehicles (as defined by IC 35-41-1), if they are used or are
- 29           intended for use by the person or persons in possession of them to
- 30           transport or in any manner to facilitate the transportation of the
- 31           following:
- 32           (A) A controlled substance for the purpose of committing,
- 33           attempting to commit, or conspiring to commit any of the
- 34           following:
- 35           (i) Dealing in or manufacturing cocaine, a narcotic drug, or
- 36           methamphetamine (IC 35-48-4-1).
- 37           (ii) Dealing in a schedule I, II, or III controlled substance
- 38           (IC 35-48-4-2).

- 1 (iii) Dealing in a schedule IV controlled substance  
2 (IC 35-48-4-3).
- 3 (iv) Dealing in a schedule V controlled substance  
4 (IC 35-48-4-4).
- 5 (v) Dealing in a counterfeit substance (IC 35-48-4-5).
- 6 (vi) Possession of cocaine, a narcotic drug, or  
7 methamphetamine (IC 35-48-4-6).
- 8 (vii) Dealing in paraphernalia (IC 35-48-4-8.5).
- 9 (viii) Dealing in marijuana, hash oil, or hashish  
10 (IC 35-48-4-10).
- 11 (B) Any stolen (IC 35-43-4-2) or converted property  
12 (IC 35-43-4-3) if the retail or repurchase value of that property  
13 is one hundred dollars (\$100) or more.
- 14 (C) Any hazardous waste in violation of IC 13-30-6-6.
- 15 (D) A bomb (as defined in IC 35-41-1-4.3) or weapon of mass  
16 destruction (as defined in IC 35-41-1-29.4) used to commit,  
17 used in an attempt to commit, or used in a conspiracy to  
18 commit an offense under IC 35-47 as part of or in furtherance  
19 of an act of terrorism (as defined by IC 35-41-1-26.5).
- 20 (2) All money, negotiable instruments, securities, weapons,  
21 communications devices, or any property used to commit, used in  
22 an attempt to commit, or used in a conspiracy to commit an  
23 offense under IC 35-47 as part of or in furtherance of an act of  
24 terrorism or commonly used as consideration for a violation of  
25 IC 35-48-4 (other than items subject to forfeiture under  
26 IC 16-42-20-5 or IC 16-6-8.5-5.1 before its repeal):
- 27 (A) furnished or intended to be furnished by any person in  
28 exchange for an act that is in violation of a criminal statute;
- 29 (B) used to facilitate any violation of a criminal statute; or
- 30 (C) traceable as proceeds of the violation of a criminal statute.
- 31 (3) Any portion of real or personal property purchased with  
32 money that is traceable as a proceed of a violation of a criminal  
33 statute.
- 34 (4) A vehicle that is used by a person to:
- 35 (A) commit, attempt to commit, or conspire to commit;
- 36 (B) facilitate the commission of; or
- 37 (C) escape from the commission of;
- 38 murder (IC 35-42-1-1), kidnapping (IC 35-42-3-2), criminal

- 1 confinement (IC 35-42-3-3), rape (IC 35-42-4-1), child molesting  
 2 (IC 35-42-4-3), or child exploitation (IC 35-42-4-4), or an offense  
 3 under IC 35-47 as part of or in furtherance of an act of terrorism.  
 4 (5) Real property owned by a person who uses it to commit any  
 5 of the following as a Class A felony, a Class B felony, or a Class  
 6 C felony:
- 7 (A) Dealing in or manufacturing cocaine, a narcotic drug, or  
 8 methamphetamine (IC 35-48-4-1).
  - 9 (B) Dealing in a schedule I, II, or III controlled substance  
 10 (IC 35-48-4-2).
  - 11 (C) Dealing in a schedule IV controlled substance  
 12 (IC 35-48-4-3).
  - 13 (D) Dealing in marijuana, hash oil, or hashish (IC 35-48-4-10).
- 14 (6) Equipment and recordings used by a person to commit fraud  
 15 under IC 35-43-5-4(11).
- 16 (7) Recordings sold, rented, transported, or possessed by a person  
 17 in violation of IC 24-4-10.
- 18 (8) Property (as defined by IC 35-41-1-23) or an enterprise (as  
 19 defined by IC 35-45-6-1) that is the object of a corrupt business  
 20 influence violation (IC 35-45-6-2).
- 21 (9) Unlawful telecommunications devices (as defined in  
 22 IC 35-45-13-6) and plans, instructions, or publications used to  
 23 commit an offense under IC 35-45-13.
- 24 (10) Any equipment used or intended for use in preparing,  
 25 photographing, recording, videotaping, digitizing, printing,  
 26 copying, or disseminating matter in violation of IC 35-42-4-4.
- 27 (11) Destructive devices used, possessed, transported, or sold in  
 28 violation of IC 35-47.5.
- 29 (12) Cigarettes that are sold in violation of IC 24-3-5.2, cigarettes  
 30 that a person attempts to sell in violation of IC 24-3-5.2, and other  
 31 personal property owned and used by a person to facilitate a  
 32 violation of IC 24-3-5.2.
- 33 (13) Tobacco products that are sold in violation of IC 24-3-5,  
 34 tobacco products that a person attempts to sell in violation of  
 35 IC 24-3-5, and other personal property owned and used by a  
 36 person to facilitate a violation of IC 24-3-5.
- 37 **(14) If a person is convicted of an offense specified in**  
 38 **IC 25-26-14-26(b) or IC 35-43-10, the following real or**

- 1           **personal property:**
- 2           **(A) Property used or intended to be used to commit,**
- 3           **facilitate, or promote the commission of the offense.**
- 4           **(B) Property constituting, derived from, or traceable to the**
- 5           **gross proceeds that the person obtained directly or**
- 6           **indirectly as a result of the offense.**
- 7           (b) A vehicle used by any person as a common or contract carrier in
- 8           the transaction of business as a common or contract carrier is not
- 9           subject to seizure under this section, unless it can be proven by a
- 10          preponderance of the evidence that the owner of the vehicle knowingly
- 11          permitted the vehicle to be used to engage in conduct that subjects it to
- 12          seizure under subsection (a).
- 13          (c) Equipment under subsection (a)(10) may not be seized unless it
- 14          can be proven by a preponderance of the evidence that the owner of the
- 15          equipment knowingly permitted the equipment to be used to engage in
- 16          conduct that subjects it to seizure under subsection (a)(10).
- 17          (d) Money, negotiable instruments, securities, weapons,
- 18          communications devices, or any property commonly used as
- 19          consideration for a violation of IC 35-48-4 found near or on a person
- 20          who is committing, attempting to commit, or conspiring to commit any
- 21          of the following offenses shall be admitted into evidence in an action
- 22          under this chapter as prima facie evidence that the money, negotiable
- 23          instrument, security, or other thing of value is property that has been
- 24          used or was to have been used to facilitate the violation of a criminal
- 25          statute or is the proceeds of the violation of a criminal statute:
- 26               (1) IC 35-48-4-1 (dealing in or manufacturing cocaine, a narcotic
- 27               drug, or methamphetamine).
- 28               (2) IC 35-48-4-2 (dealing in a schedule I, II, or III controlled
- 29               substance).
- 30               (3) IC 35-48-4-3 (dealing in a schedule IV controlled substance).
- 31               (4) IC 35-48-4-4 (dealing in a schedule V controlled substance) as
- 32               a Class B felony.
- 33               (5) IC 35-48-4-6 (possession of cocaine, a narcotic drug, or
- 34               methamphetamine) as a Class A felony, Class B felony, or Class
- 35               C felony.
- 36               (6) IC 35-48-4-10 (dealing in marijuana, hash oil, or hashish) as
- 37               a Class C felony.
- 38          SECTION 61. IC 35-43-10 IS ADDED TO THE INDIANA CODE

1 AS A NEW CHAPTER TO READ AS FOLLOWS [EFFECTIVE  
2 JULY 1, 2005]:

3 **Chapter 10. Legend Drug Deception**

4 **Sec. 1. The definitions in IC 25-26-14 apply throughout this**  
5 **chapter.**

6 **Sec. 2. A person who knowingly or intentionally:**

- 7 (1) possesses a contraband legend drug;  
8 (2) sells, delivers, or possesses with intent to sell or deliver a  
9 contraband legend drug;  
10 (3) forges, counterfeits, or falsely creates a label for a legend  
11 drug or falsely represents a factual matter contained on a  
12 label of a legend drug; or  
13 (4) manufactures, purchases, sells, delivers, brings into  
14 Indiana, or possesses a contraband legend drug;

15 **commits legend drug deception, a Class D felony.**

16 **Sec. 3. A person:**

- 17 (1) who knowingly or intentionally manufactures, purchases,  
18 sells, delivers, brings into Indiana, or possesses a contraband  
19 legend drug; and  
20 (2) whose act under subdivision (1) results in the death of an  
21 individual;

22 **commits legend drug deception resulting in death, a Class A**  
23 **felony."**

24 Page 18, after line 42, begin a new paragraph and insert:

25 "SECTION 63. IC 35-48-7-5 IS AMENDED TO READ AS  
26 FOLLOWS [EFFECTIVE JULY 1, 2005]: Sec. 5. As used in this  
27 chapter, "identification number" refers to **the following:**

28 (1) The unique number contained on any of the following:

29 ~~(1)~~ (A) A valid driver's license of a recipient or a recipient's  
30 representative issued under Indiana law or the law of any other  
31 state.

32 ~~(2)~~ (B) A recipient's or a recipient representative's valid  
33 military identification card.

34 ~~(3)~~ (C) A valid identification card of a recipient or a recipient's  
35 representative issued by:

36 ~~(A)~~ (i) the bureau of motor vehicles and described in  
37 IC 9-24-16-3; or

38 ~~(B)~~ (ii) any other state and that is similar to the identification

1 card issued by the bureau of motor vehicles.  
 2 ~~(A)~~ **(D)** If the recipient is an animal:  
 3 ~~(A)~~ **(i)** the valid driver's license issued under Indiana law or  
 4 the law of any other state;  
 5 ~~(B)~~ **(ii)** the valid military identification card; or  
 6 ~~(C)~~ **(iii)** the valid identification card issued by the bureau of  
 7 motor vehicles and described in IC 9-24-16-3 or a valid  
 8 identification card of similar description that is issued by any  
 9 other state;  
 10 of the animal's owner.

11 **(2) The identification number or phrase designated by the**  
 12 **central repository.**

13 SECTION 64. IC 35-48-7-8 IS AMENDED TO READ AS  
 14 FOLLOWS [EFFECTIVE JULY 1, 2005]: Sec. 8. The advisory  
 15 committee shall provide for a controlled substance prescription  
 16 monitoring program that includes the following components:

- 17 (1) Each time a controlled substance designated by the advisory  
 18 committee under IC 35-48-2-5 through IC 35-48-2-10 is  
 19 dispensed, the dispenser shall transmit to the central repository the  
 20 following information:  
 21 (A) The recipient's name.  
 22 (B) The recipient's or the recipient representative's  
 23 identification number **or the identification number or phrase**  
 24 **designated by the central repository.**  
 25 (C) The recipient's date of birth.  
 26 (D) The national drug code number of the controlled substance  
 27 dispensed.  
 28 (E) The date the controlled substance is dispensed.  
 29 (F) The quantity of the controlled substance dispensed.  
 30 (G) The number of days of supply dispensed.  
 31 (H) The dispenser's United States Drug Enforcement Agency  
 32 registration number.  
 33 (I) The prescriber's United States Drug Enforcement Agency  
 34 registration number.  
 35 (J) An indication as to whether the prescription was  
 36 transmitted to the pharmacist orally or in writing.  
 37 (2) The information required to be transmitted under this section  
 38 must be transmitted not more than fifteen (15) days after the date



- 1 on which a controlled substance is dispensed.
- 2 (3) A dispenser shall transmit the information required under this
- 3 section by:
- 4 (A) an electronic device compatible with the receiving device
- 5 of the central repository;
- 6 (B) a computer diskette;
- 7 (C) a magnetic tape; or
- 8 (D) a pharmacy universal claim form;
- 9 that meets specifications prescribed by the advisory committee.
- 10 (4) The advisory committee may require that prescriptions for
- 11 controlled substances be written on a one (1) part form that cannot
- 12 be duplicated. However, the advisory committee may not apply
- 13 such a requirement to prescriptions filled at a pharmacy with a
- 14 Type II permit (as described in IC 25-26-13-17) and operated by
- 15 a hospital licensed under IC 16-21, or prescriptions ordered for
- 16 and dispensed to bona fide enrolled patients in facilities licensed
- 17 under IC 16-28. The committee may not require multiple copy
- 18 prescription forms and serially numbered prescription forms for
- 19 any prescriptions written. The committee may not require
- 20 different prescription forms for any individual drug or group of
- 21 drugs. Prescription forms required under this subdivision must be
- 22 jointly approved by the committee and by the Indiana board of
- 23 pharmacy established by IC 25-26-13-3.
- 24 (5) The costs of the program.

25 SECTION 65. [EFFECTIVE JULY 1, 2005] (a) **IC 25-26-14, as**

26 **amended by this act, applies:**

- 27 (1) **after June 30, 2005, for an initial license issued under**
- 28 **IC 25-26-14, as amended by this act; and**
- 29 (2) **on the first expiration date occurring after December 31,**
- 30 **2005, for renewal of a license issued under IC 25-26-14, before**
- 31 **amendment by this act.**
- 32 (b) **The Indiana board of pharmacy established by IC 25-26-13-3**
- 33 **may establish an electronic pedigree pilot program to authenticate,**
- 34 **track, and trace legend drugs. The pilot program must include**
- 35 **participation of drug manufacturers, wholesale drug distributors,**
- 36 **and pharmacies that are licensed in Indiana. The board may**
- 37 **establish the requirements and guidelines for the pilot program.**
- 38 (c) **Before June 30, 2007, the Indiana board of pharmacy**

1        **established by IC 25-26-13-3 shall conduct a study of the electronic**  
2        **pedigree pilot program. The study must include consultation with**  
3        **manufacturers, distributors, and pharmacies that participate in the**  
4        **electronic pedigree pilot program. The study may include the**  
5        **consultation with manufacturers, distributors, and pharmacies that**  
6        **do not participate in the electronic pedigree pilot program. Based**  
7        **on the results of the study, the board shall determine a date to**  
8        **implement a mandatory electronic pedigree program. However, the**  
9        **board may not implement a mandatory electronic pedigree**  
10       **program until after the board has completed the study under this**  
11       **subsection.**

12        **(d) This SECTION expires December 31, 2007."**

13        Renumber all SECTIONS consecutively.

(Reference is to SB 590 as reprinted February 11, 2005.)

**and when so amended that said bill do pass.**

---

Representative Becker