

**BEFORE THE  
BOARD OF PHARMACY  
DEPARTMENT OF CONSUMER AFFAIRS  
STATE OF CALIFORNIA**

In the Matter of the Accusation Against:

**PHARMACY RESOURCES INC.; GREGG  
N. PEDERSON, PRESIDENT AND PIC  
5290 E. Yale Circle, No. 101  
Denver, CO 80222**

**Non-Resident Pharmacy License No. NRP  
45418  
Non-Resident Sterile Compounding Permit  
No. NSC 99697**

Respondent.

Case No. 6021

OAH No. 2018020256

**STIPULATED SURRENDER OF  
LICENSE AND ORDER**

**DECISION AND ORDER**

The attached Stipulated Surrender of License and Order is hereby adopted by the Board of Pharmacy, Department of Consumer Affairs, as its Decision in this matter.

This Decision shall become effective at 5:00 p.m. on August 2, 2018.

It is so ORDERED on July 3, 2018.

BOARD OF PHARMACY  
DEPARTMENT OF CONSUMER AFFAIRS  
STATE OF CALIFORNIA



By \_\_\_\_\_

Victor Law, R.Ph.  
Board President

1 XAVIER BECERRA  
Attorney General of California  
2 DAVID E. BRICE  
Supervising Deputy Attorney General  
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7 *Attorneys for Complainant*

8 **BEFORE THE**  
9 **BOARD OF PHARMACY**  
10 **DEPARTMENT OF CONSUMER AFFAIRS**  
11 **STATE OF CALIFORNIA**

12 In the Matter of the Accusation Against:

Case No. 6021

13 **PHARMACY RESOURCES INC.; GREGG**  
14 **N. PEDERSON, PRESIDENT AND PIC**  
15 **5290 E. Yale Circle, No. 101**  
16 **Denver, CO 80222**

OAH No. 2018020256

**STIPULATED SURRENDER OF  
LICENSE AND ORDER**

17 **Non-Resident Pharmacy Permit No. NRP**  
18 **1126**  
19 **Non-Resident Sterile Compounding Permit**  
20 **No. NSC 99697**

21 Respondent.

22 IT IS HEREBY STIPULATED AND AGREED by and between the parties to the above-  
23 entitled proceedings that the following matters are true:

24 PARTIES

25 1. Virginia Herold (Complainant) is the Executive Officer of the Board of Pharmacy  
26 (Board). She brought this action solely in her official capacity and is represented in this matter by  
27 Xavier Becerra, Attorney General of the State of California, by Patricia Webber Heim, Deputy  
28 Attorney General.

1           2.    Pharmacy Resources Inc.; Gregg N. Pederson, President and Pharmacist-In-Charge  
2 (PIC) (Respondent) is represented in this proceeding by attorney Ivan Petrzelka, whose address is  
3 Ivan Petrzelka, California Pharmacy Lawyers, 49 Discovery, Suite 240, Irvine, CA 92618.

4           3.    On or about October 3, 2011, the Board issued Non-Resident Pharmacy Permit No.  
5 NRP 1126 to Respondent. The Non-Resident Pharmacy Permit was in full force and effect at all  
6 times relevant to the charges brought in Accusation No. 6021, expired on October 1, 2017, and  
7 has not been renewed.

8           4.    On or about November 3, 2011, the Board issued Non-Resident Sterile Compounding  
9 Permit No. NSC 99697 to Respondent. The Non-Resident Sterile Compounding Permit expired  
10 on October 1, 2016, and has not been renewed.

11   JURISDICTION

12           5.    Accusation No. 6021 was filed before the Board, and is currently pending against  
13 Respondent. The Accusation and all other statutorily required documents were properly served  
14 on Respondent on January 17, 2018. Respondent timely filed its Notice of Defense contesting the  
15 Accusation. A copy of Accusation No. 6021 is attached as Exhibit A and incorporated by  
16 reference.

17   ADVISEMENT AND WAIVERS

18           6.    Respondent has carefully read, fully discussed with counsel, and understands the  
19 charges and allegations in Accusation No. 6021. Respondent also has carefully read, fully  
20 discussed with counsel, and understands the effects of this Stipulated Surrender of License and  
21 Order.

22           7.    Respondent is fully aware of its legal rights in this matter, including the right to a  
23 hearing on the charges and allegations in the Accusation; the right to confront and cross-examine  
24 the witnesses against them; the right to present evidence and to testify on its own behalf; the right  
25 to the issuance of subpoenas to compel the attendance of witnesses and the production of  
26 documents; the right to reconsideration and court review of an adverse decision; and all other  
27 rights accorded by the California Administrative Procedure Act and other applicable laws.





1           4.     Respondent may not reapply for any license from the Board for three (3) years from  
2 the effective date of this decision.

3           5.     If Respondent ever applies for licensure or petitions for reinstatement in the State of  
4 California, the Board shall treat it as a new application for licensure. Respondent must comply  
5 with all the laws, regulations and procedures for licensure in effect at the time the application or  
6 petition is filed, and all of the charges and allegations contained in Accusation No. 6021 shall be  
7 deemed to be true, correct and admitted by Respondent when the Board determines whether to  
8 grant or deny the application or petition.

9           6.     Respondent shall pay the agency its costs of investigation and enforcement in the  
10 amount of \$7,500.00 prior to issuance of a new or reinstated license.

11          7.     If Respondent should ever apply or reapply for a new license or certification, or  
12 petition for reinstatement of a license, by any other health care licensing agency in the State of  
13 California, all of the charges and allegations contained in Accusation, No. 6021 shall be deemed  
14 to be true, correct, and admitted by Respondent for the purpose of any Statement of Issues or any  
15 other proceeding seeking to deny or restrict licensure.

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
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
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ACCEPTANCE

I have carefully read the above Stipulated Surrender of License and Order and have fully discussed it with my attorney, Ivan Petrzelka. I understand the stipulation and the effect it will have on my Non-Resident Pharmacy Permit and Non-Resident Sterile Compounding Permit. I enter into this Stipulated Surrender of License and Order voluntarily, knowingly, and intelligently, and agree to be bound by the Decision and Order of the Board of Pharmacy.

DATED: 5/1/2018   
GREGG N. PEDERSON, PRESIDENT AND  
PHARMACIST-IN-CHARGE, PHARMACY  
RESOURCES INC.  
*Respondent*

I have read and fully discussed with Respondent Gregg N. Pederson, President and Pharmacist-In-Charge of Pharmacy Resources Inc. the terms and conditions and other matters contained in this Stipulated Surrender of License and Order. I approve its form and content.

DATED: May 6, 2018   
IVAN PETRZELKA  
*Attorney for Respondent*

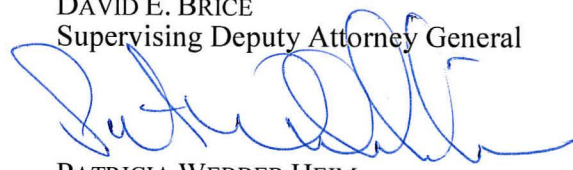
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ENDORSEMENT

The foregoing Stipulated Surrender of License and Order is hereby respectfully submitted for consideration by the Board of Pharmacy of the Department of Consumer Affairs.

Dated: 5/7/18

Respectfully submitted,  
XAVIER BECERRA  
Attorney General of California  
DAVID E. BRICE  
Supervising Deputy Attorney General



PATRICIA WEBBER HEIM  
Deputy Attorney General  
*Attorneys for Complainant*

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**Exhibit A**

**Accusation No. 6021**

1 XAVIER BECERRA  
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2 JANICE K. LACHMAN  
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3 PATRICIA WEBBER HEIM  
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7 *Attorneys for Complainant*

8 **BEFORE THE**  
**BOARD OF PHARMACY**  
9 **DEPARTMENT OF CONSUMER AFFAIRS**  
10 **STATE OF CALIFORNIA**

11 In the Matter of the Accusation Against:

Case No. 6021

12 **PHARMACY RESOURCES INCORPORATED**  
13 **GREGG N. PEDERSON, PRES./PIC**  
14 **JANET L. PEDERSON, SECTY**  
15 **5290 E. Yale Circle, No. 101**  
16 **Denver, CO 80222**

**A C C U S A T I O N**

17 **Non-Resident Pharmacy Permit No. NRP 1126**  
18 **Non-Resident Sterile Compounding Permit No.**  
19 **NSC 99697**

Respondent.

20 Complainant alleges:

21 **PARTIES**

22 1. Virginia Herold (Complainant) brings this Accusation solely in her official capacity  
23 as the Executive Officer of the Board of Pharmacy, Department of Consumer Affairs.

24 2. On or about October 3, 2011, the Board of Pharmacy issued Non-Resident Pharmacy  
25 Permit Number NRP 1126 to Pharmacy Resources Incorporated (Respondent), with Gregg N.  
26 Pederson (Pederson) as president and pharmacist-in-charge and Janet L. Pederson as secretary.  
27 The Non-Resident Pharmacy Permit was in full force and effect at all times relevant to the  
28 charges brought in the Accusation and will expire on October 1, 2017, unless renewed.

///

1 3. On or about November 3, 2011, the Board of Pharmacy issued Non-Resident Sterile  
2 Compounding Permit Number NSC 99697 to Respondent. The Non-Resident Sterile  
3 Compounding permit was in full force and effect at all times relevant to the charges brought in  
4 the Accusation; however, it expired on October 1, 2016, and has not been renewed.

5 **JURISDICTION**

6 4. This Accusation is brought before the Board of Pharmacy (Board) under the authority  
7 of the following laws. All section references are to the Business and Professions Code (Code)  
8 unless otherwise indicated.

9 5. Code section 4300 states, in pertinent part:

10 (a) Every license issued may be suspended or revoked.

11 (b) The board shall discipline the holder of any license issued by the  
12 board, whose default has been entered or whose case has been heard by the board and  
found guilty, by any of the following methods:

13 (1) Suspending judgment.

14 (2) Placing him or her upon probation.

15 (3) Suspending his or her right to practice for a period not exceeding one  
16 year.

17 (4) Revoking his or her license.

18 (5) Taking any other action in relation to disciplining him or her as the  
board in its discretion may deem proper . . .

19 6. Code section 4300.1 states:

20 The expiration, cancellation, forfeiture, or suspension of a board-issued  
21 license by operation of law or by order or decision of the board or a court of law, the  
22 placement of a license on a retired status, or the voluntary surrender of a license by a  
23 licensee shall not deprive the board of jurisdiction to commence or proceed with any  
investigation of, or action or disciplinary proceeding against, the licensee or to render  
a decision suspending or revoking the license.

24 **STATUTORY AND REGULATORY PROVISIONS**

25 7. Code section 4301 states, in pertinent part:

26 The board shall take action against any holder of a license who is guilty  
27 of unprofessional conduct . . . Unprofessional conduct shall include, but is not limited  
to, any of the following:

28 . . . .

1 (o) Violating or attempting to violate, directly or indirectly, or assisting in  
2 or abetting the violation of or conspiring to violate any provision or term of this  
3 chapter or of the applicable federal and state laws and regulations governing  
4 pharmacy, including regulations established by the board or by any other state or  
5 federal regulatory agency . . .

6 8. Code section 4076 states, in pertinent part:

7 (a) A pharmacist shall not dispense any prescription except in a container  
8 that meets the requirements of state and federal law and is correctly labeled with all  
9 of the following: . . .

10 . . . . .

11 (2) The directions for the use of the drug.

12 . . . . .

13 (8) The quantity of the drug or drugs dispensed . . .

14 9. Code section 4127.2, subdivision (a), states that “[a] nonresident pharmacy shall not  
15 compound injectable sterile drug products for shipment into the State of California without a  
16 license issued by the board pursuant to this section. The license shall be renewed annually and  
17 shall not be transferable”.

18 10. California Code of Regulations, title 16, section 1707.5 states, in pertinent part:

19 (a) Labels on drug containers dispensed to patients in California shall  
20 conform to the following format:

21 (1) Each of the following items, and only these four items, shall be  
22 clustered into one area of the label that comprises at least 50 percent of the label.  
23 Each item shall be printed in at least a 12-point sans serif typeface, and listed in the  
24 following order:

25 (A) Name of the patient.

26 (B) Name of the drug and strength of the drug. For the purposes of this  
27 section, “name of the drug” means either the manufacturer’s trade name of the drug,  
28 or the generic name and the name of the manufacturer.

(C) The directions for the use of the drug.

(D) The condition or purpose for which the drug was prescribed if the  
condition or purpose is indicated on the prescription . . .

11. California Code of Regulations, title 16, section 1735.2, subdivision (h), states that  
“[a]ll chemicals, bulk drug substances, drug products, and other components used for drug

///

1 compounding shall be stored and used according to compendia and other applicable requirements  
2 to maintain their integrity, potency, quality, and labeled strength.”

3 12. California Code of Regulations, title 16, section 1735.4 states, in pertinent part:

4 (a) Each compounded drug preparation shall be affixed with a container  
5 label prior to dispensing that contains at least:

6 . . . .

7 (3) Instructions for storage, handling, and administration. For admixed IV  
8 solutions, the rate of infusion shall be included . . .

9 13. California Code of Regulations, title 16, section 1751.3 states, in pertinent part:

10 (a) Any pharmacy engaged in compounding sterile drug preparations  
11 shall maintain written policies and procedures for compounding. Any material failure  
12 to follow the pharmacy’s written policies and procedures shall constitute a basis for  
13 disciplinary action. In addition to the elements required by section 1735.5, there shall  
14 be written policies and procedures regarding the following:

15 . . . .

16 (12) Labeling of the sterile compounded drug preparations based on the  
17 intended route of administration and recommended rate of administration . . .

18 14. California Code of Regulations, title 16, section 1751.7 states, in pertinent part:

19 . . . .

20 (e)(1) Batch-produced sterile drug preparations compounded from one or  
21 more non-sterile ingredients . . . shall be subject to documented end product testing  
22 for sterility and pyrogens and shall be quarantined until the end product testing  
23 confirms sterility and acceptable levels of pyrogens. Sterility testing shall be USP  
24 chapter 71 compliant and pyrogens testing shall confirm acceptable levels of  
25 pyrogens per USP chapter 85 limits, before dispensing. This requirement of end  
26 product testing confirming sterility and acceptable levels of pyrogens prior to  
27 dispensing shall apply regardless of any sterility or pyrogen testing that may have  
28 been conducted on any ingredient or combination of ingredients that were previously  
non-sterile. Exempt from pyrogen testing are topical ophthalmic and inhalation  
preparations . . .

**COST RECOVERY**

15 15. Code section 125.3 provides, in pertinent part, that a Board may request the  
16 administrative law judge to direct a licentiate found to have committed a violation or violations of  
17 the licensing act to pay a sum not to exceed the reasonable costs of the investigation and  
18 enforcement of the case.

19 ///

1 **DRUG CLASSIFICATIONS**

2 16. Glycopyrrolate 0.2 mg/ml is a dangerous drug pursuant to Code section 4022 and is  
3 indicated for the reduction of secretions in horses. "Robinul 0.2 mg/ml" is a brand name for  
4 Glycopyrrolate.

5 17. Cimetidine 150 mg/ml is a dangerous drug pursuant to Code section 4022 and is  
6 indicated for the treatment of gastric ulcers and melanomas and stomach protection in horses.  
7 "Tagamet 300 mg/2ml" is a brand name for Cimetidine.

8 18. Cacodylate/copper 6 mg/ml is a dangerous drug pursuant to Code section 4022 and is  
9 indicated for the treatment of anemia in horses.

10 19. Methocarbamol 100 mg/ml is a dangerous drug pursuant to Code section 4022 and is  
11 indicated for the treatment of muscle spasms in horses.

12 **FACTUAL ALLEGATIONS**

13 20. On or about September 9, 2016, Board Inspector D. P. conducted a non-resident  
14 sterile compounding permit renewal inspection at Respondent's pharmacy located in Denver,  
15 Colorado, and was assisted by pharmacist-in-charge Pederson. D. P. asked Pederson what risk  
16 levels of sterile compounds were dispensed to California patients. Pederson stated that the  
17 pharmacy provided mainly non-sterile to sterile (high-risk) veterinary products.

18 21. D. P. requested various documents from Pederson, including compounding policies  
19 and procedures, completed recent patient-specific compounding records with associated  
20 prescription labels, quality assurance policies and procedures with documentation of end product  
21 testing, non-sterile to sterile compounded product testing documentation, and a list of all  
22 compounded preparations provided to California patients during the past year. Pederson told  
23 D. P. that his staff kept a log of sterile preparations compounded by the pharmacy and that the  
24 California patients could be highlighted on the log for the inspector's review. D. P. was given a  
25 log listing veterinary sterile compounded preparations that were supplied to patients with the  
26 California patients highlighted.

27 22. D. P. was also given compounding log worksheets for approximately eight sterile  
28 compounded preparations and the related prescriptions, prescription labels, and "soy-logs" (logs

1 documenting in-house sterility testing of drug products). D. P. found that the prescription labels,  
2 including the label for Glycopyrrolate 0.2 mg/ml, were not in compliance with California law.  
3 D. P. also found that four sterile compounded injectable preparations, glycopyrrolate, cimetidine,  
4 cacodylate/copper, and methocarbamol, were dispensed with beyond use dates (BUD's) greater  
5 than 180 days. D. P. requested the validation paperwork for the extended BUD's. D. P. was  
6 given a certificate of analysis and a soy log for only one of the four injectable preparations,  
7 glycopyrrolate. D. P. was advised that the extended BUD's for the other three drug products were  
8 validated some time ago and that the paperwork might not be easy to locate.

9 23. On and between September 12, 2016 and September 30, 2016, D. P. requested certain  
10 documents and information from Pederson, including a list of sterile drug products the pharmacy  
11 provided to California patients during the last year; a current copy of the pharmacy's sterile  
12 compounding policy and procedures; dispensing records of sterile preparations the pharmacy  
13 provided to California patients for the last sixth months; the master formulas, sterility and  
14 pyrogens testing data and documentation, and stability studies used to extend the BUD's for  
15 glycopyrrolate, cimetidine, cacodylate/copper, and methocarbamol; and the compounding logs,  
16 master formulas, prescriptions, labels, and sterility and pyrogens testing data and documentation  
17 for certain other compounded products the pharmacy sent to California patients.

18 24. On and between September 23, 2016 and September 30, 2016, Pederson sent D. P. an  
19 Excel spreadsheet listing the sterile compounded prescriptions the pharmacy supplied to  
20 California patients, written documentation and justification for the extended BUD's for the four  
21 drug preparations, and the pharmacy's compounding policies and procedures.

22 25. On or about October 5, 2016, D. P. sent Pederson an email stating that she still had  
23 not received the sterility and pyrogens data and documentation for glycopyrrolate, cimetidine,  
24 cacodylate/copper, and methocarbamol or the compounding logs, master formulas, prescriptions,  
25 labels, and sterility and pyrogens testing data and documentation for the other compounded  
26 products.

27 26. On or about October 7, 2016, Pederson provided D. P. with compounding worksheets  
28 and prescriptions labels for estrone AQ suspension 5 mg/ml injection, acetyl D glucosamine 50

1 mg/ml injection, and stanozolol suspension 50 mg/ml injection, and a revised Excel spreadsheet.  
 2 D. P. found that of the 164 prescriptions listed on the spreadsheet, approximately 72 prescriptions  
 3 (44%) of sterile compounded products were dispensed prior to the required 14-day quarantine  
 4 period. D. P. also found that the prescription labels for Estrone, Acetyl D Glucosamine, and  
 5 Stanozolol were not in compliance with California law and that the three drug products were  
 6 dispensed with BUD's greater than 180 days.

7 **FIRST CAUSE FOR DISCIPLINE**

8 **(Failure to Document Appropriate Beyond Use Date for Compounded Products)**

9 27. Respondent's Non-Resident Pharmacy Permit and Non-Resident Sterile  
 10 Compounding Permit are subject to disciplinary action pursuant to Code section 4301,  
 11 subdivision (o), for unprofessional conduct, in that Respondent violated California Code of  
 12 Regulations, title 16, section 1735.2, subdivision (h), as follows: Respondent failed to provide  
 13 the Board with any stability studies to support exceeding the beyond use date of 180 days for the  
 14 following sterile injectable compounded preparations:

15 <b>Compounded Preparation</b>	16 <b>Assigned Lot No.</b>	17 <b>Date Preparation Compounded</b>	18 <b>"Validated" BUD Listed on Product Worksheet</b>	19 <b>Assigned BUD</b>	20 <b>Actual BUD Days Assigned &amp; Documented</b>	21 <b>Listed Label Expiration Date</b>
22 Glycopyrrolate .2 mg/ml inj.	050816	08/05/16	270 days	05/05/17	274 days	APR '17
23 Cimetidine 150 mg/ml inj.	070916	09/07/16	360 days	09/07/17	366 days	AUG '17
24 Caco-6-Copper 6 mg/ml inj.	040816	08/03/16	240 days	04/03/17	244 days	MAR '17
25 Methocarbamol 100 mg/ml inj.	270716	07/27/16	360 days	04/19/17 (component expiration date)	236 days	APR '17
26 Estrone AQ susp 5 mg/ml inj.	060716	07/06/16	360 days	04/02/17 (component expiration date)	271 days	MAR '17
27 Glycopyrrolate .2 mg/ml inj.	050816	08/05/16	270 days	05/05/17	274 days	APR '17



Compounded Preparation	Assigned Lot No.	Date Preparation Compounded	"Validated" BUD Listed on Product Worksheet	Assigned BUD	Actual BUD Days Assigned & Documented	Listed Label Expiration Date
Stanozolol susp. 50 mg/ml inj.	210416	04/21/16	360 days	01/09/17 (component expiration date)	274 days	DEC '16

**SECOND CAUSE FOR DISCIPLINE**

**(Failure to Quarantine Batch-Produced Sterile Injectable Drug Products)**

28. Respondent's Non-Resident Pharmacy Permit and Non-Resident Sterile Compounding Permit are subject to disciplinary action pursuant to Code section 4301, subdivision (o), for unprofessional conduct, in that Respondent violated California Code of Regulations, title 16, section 1751.7, subdivision (e)(1), as follows:

a. Respondent dispensed the following sterile injectable compounded preparations, compounded from one or more non-sterile ingredients and in quantities greater than one dose, prior to documented end-product testing confirming sterility and acceptable levels of pyrogens, specifically, endotoxin testing:

Compounded Preparation	Lot #	Quantity Compounded	Date Compounded	Date Dispensed	Endotoxin Indication on Compounding Worksheet
Glycopyrrolate .2 mg/ml inj.	050816	262 ml into 12 20 ml vials	08/05/16	08/05/16	"N/A"
Cimetidine 150 mg/ml inj.	070916	260 ml into 8 30 ml vials	09/06/16	09/06/16	No endotoxin requirement listed

b. Respondent dispensed the following sterile injectable compounded preparations, compounded from one or more non-sterile ingredients and in quantities greater than one dose, prior to the 14-day quarantine period and completion of the end product testing confirming sterility and acceptable levels of pyrogens:

Rx Number	Date Written	Date Compounded	Date Dispensed	Patient Name	Compounded Preparation	Vial Quantity/Quantity Dispensed
66854	08/29/16	08/30/16	08/30/16	Armored Car	Estrone 5 mg/ml	100 ml/13 vials

Rx Number	Date Written	Date Compounded	Date Dispensed	Patient Name	Compounded Preparation	Vial Quantity/ Quantity Dispensed
66853	08/29/16	08/30/16	08/30/16	Award It	Estrone 5 mg/ml	100 ml/13 vials
66760	8/23/16	8/12/16	8/23/16	Madame Strips	Estrone 5 mg/ml	100 ml/13 vials
66618	08/12/16	8/12/16	8/12/16	Weird Haircut Steh	Tranexamic Acid 10%	100 ml/6 vials
65623	06/07/16	8/09/16	8/09/16	Zen	Ketoprofen 10%/L-Arginine 7%	100 ml/12 vials
66520	08/05/16	08/05/16	08/05/16	Raphael	Glycopyrrolate 0.mg/ml	20 ml/12 vials
66409	08/01/16	08/01/16	08/01/16	Travel Fighter	L-Arginine 10%	100 ml/13 vials
66481	08/04/16	08/04/16	8/04/16	Sunfeet	Acety D Glucosamine 100 mg/ml	50 ml/8 vials
66384	07/29/16	07/29/16	07/29/16	Velocity	Aminocaproic Acid 25%	100 ml/6 vials
66385	07/29/16	07/29/16	07/29/16	Westbrook	Estrone 5 mg/ml	100 ml/13 vials
66386	07/29/16	07/29/16	07/29/16	Magical Tech	Estrone 5 mg/ml	100 ml/13 vials
61677	08/03/15	07/29/16	07/29/16	Fancy Pants	Acety D Glucosamine 100 mg/ml	50 ml/8 vials
66355	07/28/16	07/28/16	07/28/16	Edgy Girl	Estrone 5 mg/ml	100 ml/13 vials
66019	07/05/16	07/06/16	07/06/16	Check's in the Mail	Estrone 5 mg/ml	100 ml/13 vials
66020	07/05/16	07/06/16	07/06/16	Pandora	Estrone 5 mg/ml	100 ml/13 vials
66021	07/05/16	07/06/16	07/06/16	Alaskan Fun	Estrone 5 mg/ml	100 ml/13 vials

**THIRD CAUSE FOR DISCIPLINE**

**(Failure to Label Patient-Specific Sterile Compounded Prescriptions with the Required Elements)**

29. Respondent's Non-Resident Pharmacy Permit and Non-Resident Sterile Compounding Permit are subject to disciplinary action pursuant to Code section 4301, subdivision (o), for unprofessional conduct, in that Respondent violated Code section 4076, subdivisions (a)(2) and (8), and California Code of Regulations, title 16, sections 1707.5,

1 subdivision (a)(1), 1735.4, subdivision (a)(3), and 1751.3, subdivision (a)(12), as follows:  
2 Respondent failed to label California patient-specific sterile compounded prescriptions,  
3 specifically, prescriptions for Glycopyrrolate 0.2 mg/ml injection, Estrone AQ suspension 5  
4 mg/ml injection, Acetyl D Glucosamine 50 mg/ml injection, and Stanozolol suspension 50 mg/ml  
5 injection, with the directions for use of the preparation, the total quantity of the drug or drugs  
6 dispensed, and/or the intended rate of administration. Further, Respondent failed to use the  
7 California patient-centered format on the labels.

8 **MATTERS IN AGGRAVATION**

9 30. To determine the degree of discipline to be assessed against Respondent, if any,  
10 Complainant alleges as follows:

11 a. On or about January 20, 2016, the Board issued Citation and Fine No. CI 2015 66540  
12 against Respondent's Non-Resident Pharmacy Permit for violating California Code of  
13 Regulations, title 16, section 1735.2, subdivision (j) (failure to complete compounding self-  
14 assessment). The Board ordered Respondent to pay a fine of \$500 by February 19, 2016.  
15 Respondent paid the citation in full.

16 b. On or about January 20, 2016, the Board issued Citation and Fine No. CI 2015 68710  
17 against Respondent's Non-Resident Sterile Compounding Permit for violating California Code of  
18 Regulations, title 16, section 1735.2, subdivision (j) (failure to complete compounding self-  
19 assessment).

20 **PRAYER**

21 **WHEREFORE**, Complainant requests that a hearing be held on the matters herein alleged,  
22 and that following the hearing, the Board of Pharmacy issue a decision:

23 1. Revoking or suspending Non-Resident Pharmacy Permit Number NRP 1126, issued  
24 to Pharmacy Resources Incorporated;

25 2. Revoking or suspending Non-Resident Sterile Compounding Permit Number NSC  
26 99697, issued to Pharmacy Resources Incorporated;

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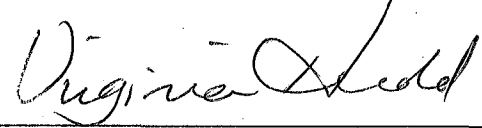
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3. Ordering Pharmacy Resources Incorporated to pay the Board of Pharmacy the reasonable costs of the investigation and enforcement of this case, pursuant to Business and Professions Code section 125.3; and,

4. Taking such other and further action as deemed necessary and proper.

DATED: 1/3/18



VIRGINIA HEROLD  
Executive Officer  
Board of Pharmacy  
Department of Consumer Affairs  
State of California  
*Complainant*

SA2016104808