

December 06, 2024

**URGENT DRUG RECALL**  
**SECOND NOTICE**

Dear Valued Customer,

This is to inform you of a product recall involving:

**Product Nebivolol Tablets 2.5 mg**

Refer Attachment 1 for NDC, dosage strength, Package Size and Lot details

**See enclosed product label for ease in identifying the product**

Aurobindo Pharma USA, Inc. has initiated a voluntary Drug Product Recall for the product Nebivolol Tablets 2.5 mg from USA market due to presence of Nitrosamine Drug Substance Related Impurity (NDSRI), N-Nitroso Nebivolol above acceptable intake (AI) limit.

Nebivolol Tablets are indicated for the treatment of hypertension, to lower blood pressure. Lowering blood pressure reduces the risk of fatal and nonfatal cardiovascular events, primarily strokes and myocardial infarctions. To date, Aurobindo has not received any reports of adverse drug events that are confirmed with this recall. Patients who are prescribed Nebivolol should continue taking their medication, as the risk of harm to the patient's health may be higher if the treatment is stopped immediately without any alternative treatment. Patients should contact their pharmacist or physician who can advise them about an alternative treatment prior to returning their medication.

As per the product information leaflet, Nebivolol Tablets 2.5 mg are light blue colored, round shaped, biconvex uncoated tablets debossed with 'L' on one side and '78' on the other side.

Aurobindo USA began shipping impacted batches to customers nationwide from June 13, 2024 to November 20, 2024.

Please complete and return the enclosed product return response form (business response form) as soon as possible. Consequently, please examine your inventory and quarantine the product batches subject to this recall. In addition, if you have further distributed these batches, please notify your customers accordingly and contact Qualanex with any questions. The product can be identified by checking the product name and batch or lot number on the product label.

As this recall should be carried out to the retail level, your assistance is appreciated and necessary to ensure patient safety.



**Recall Instructions:**

Please perform the following activities:

- Immediately examine your inventory and quarantine the specified lots of Nebivolol Tablets 2.5 mg.
- Immediately discontinue the distribution of the specific lots being recalled.
- Promptly complete the Product Return Response Form even if you have no product to return. If you have further distributed this product, notify your customers of this recall by including a copy of this notice.
- In the event you have the recall product, please return it to Qualanex, LLC., using the enclosed postage paid product return label and mail to the following.  
Qualanex, LLC.  
1410 Harris Road  
Libertyville, IL 60048

Completed product response form (Yes) may be return by the following methods:

- Email to [recall@qualanex.com](mailto:recall@qualanex.com)
- Fax to 847-737-3719
- Mail to:  
Aurobindo USA C/O Qualanex LLC  
1410 Harris Rd  
Libertyville, IL 60048

Please complete and return the enclosed Product Return Response Form as soon as possible. If you need assistance in returning your product or have questions about the recall process, contact Qualanex at 888-504-2014 during the hours of 7:00 AM to 4:00 PM CST.

Please return your product along with return goods authorization form using postage paid shipping label included in your recall return packet. Once the response form is received, a return authorization will be generated and e-mailed to you to be included with your recall returns. Appropriate reimbursement for the product returns will be issued on the receipt upon the recall product.

This recall is being made with the knowledge of the Food and Drug Administration.





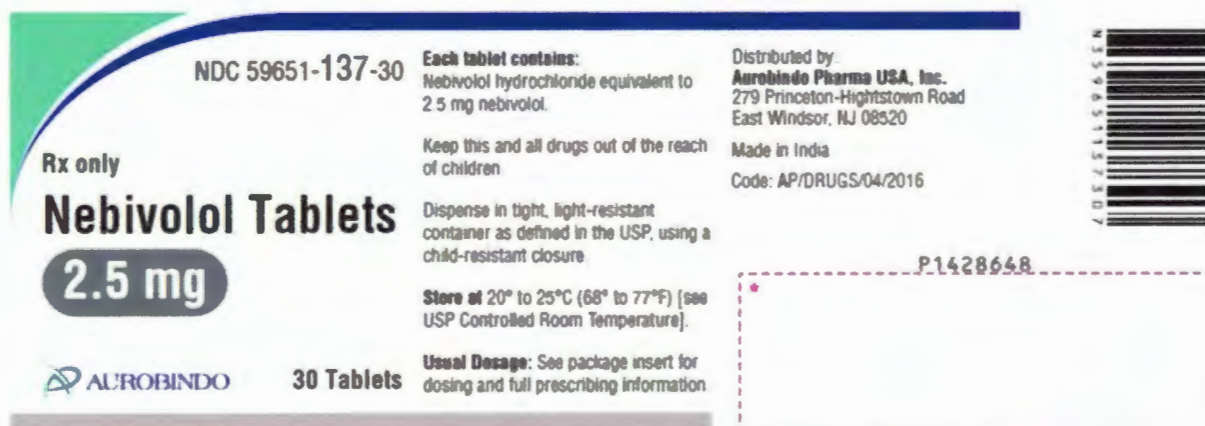
Sincerely,

**Linda Torres**

Digitally signed by Linda Torres  
DN: cn=Linda Torres, o=Aurobindo Pharma LLC,  
ou=QA Product Complaints,  
email=l.torres@aurobindousa.com, c=US  
Date: 2014.12.26 07:09:26 -0500

Ms. Linda Torres  
QA Product Complaint Manager  
Aurobindo Pharma USA, Inc  
279 Princeton-Hightstown Rd  
East Windsor, NJ 08520-1401  
Direct Email: [LTorres@aurobindousa.com](mailto:LTorres@aurobindousa.com)  
Phone-732-839-9419  
Fax-732-289-6189

The product label is as shown below:



**NDC 59651-137-30**

**Rx only**

**Nebivolol Tablets**

**2.5 mg**

**30 Tablets**

**AUROBINDO**

**Each tablet contains:**  
Nebivolol hydrochloride equivalent to  
2.5 mg nebivolol.

Keep this and all drugs out of the reach  
of children

Dispense in tight, light-resistant  
container as defined in the USP, using a  
child-resistant closure

Store at 20° to 25°C (68° to 77°F) [see  
USP Controlled Room Temperature].

**Usual Dosage:** See package insert for  
dosing and full prescribing information

Distributed by:  
**Aurobindo Pharma USA, Inc.**  
279 Princeton-Hightstown Road  
East Windsor, NJ 08520


Made in India

Code: AP/DRUGS/04/2016

**P1428648**

**\*Over Printing Zone**

**Coding Area**  
(45 x 15 mm)  
Dotted lines not to be printed

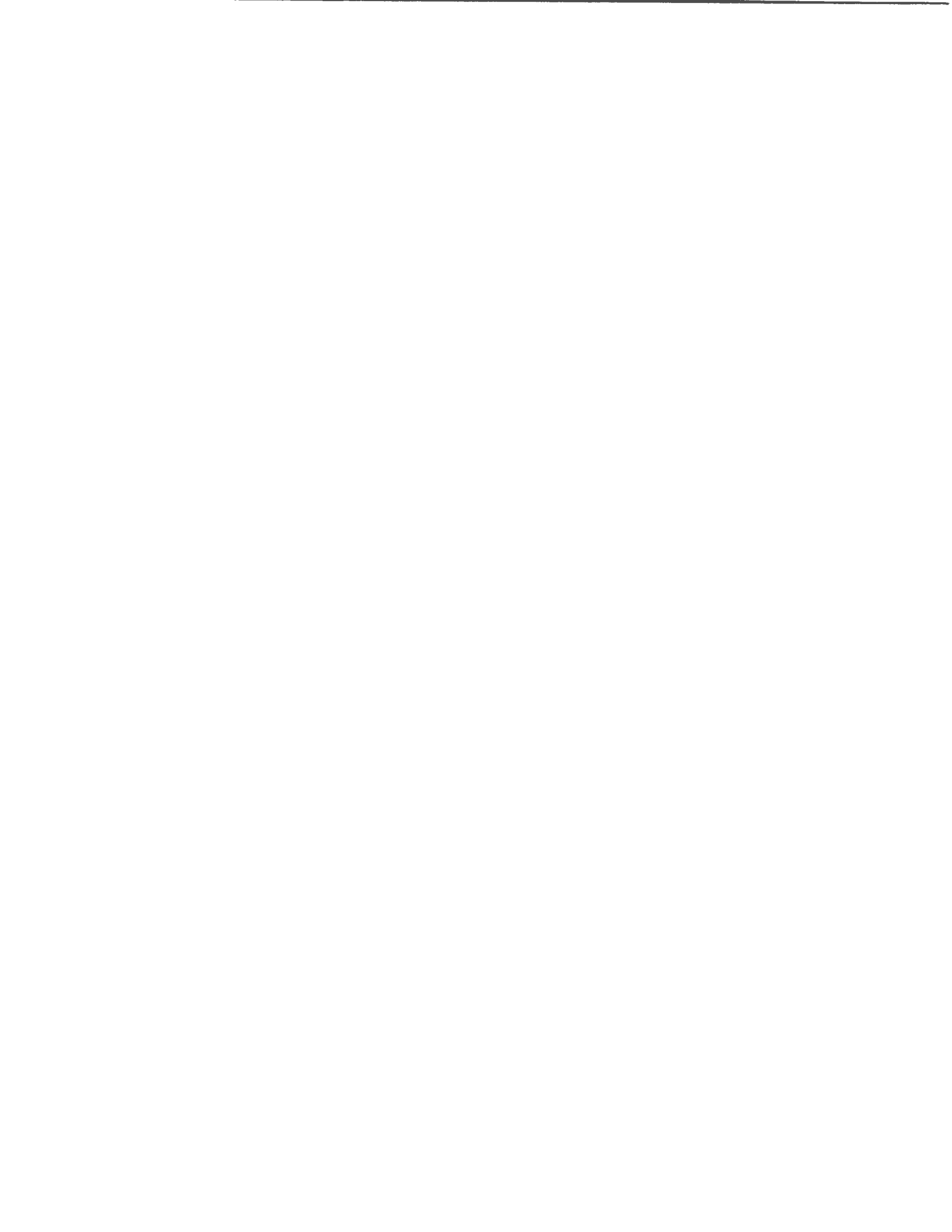




**Attachment - 1**

**Batches manufactured at APL Healthcare Limited Unit-IV**

| <b>NDC Number</b> | <b>Dosage Strength</b> | <b>Package Size</b> | <b>Lot Number</b> | <b>Expiration Date</b> |
|-------------------|------------------------|---------------------|-------------------|------------------------|
| 59651-137-30      | 2.5 mg                 | 30's HDPE Container | NB0224001A        | 04/2027                |
| 59651-137-30      | 2.5 mg                 | 30's HDPE Container | NB0224001B        | 04/2027                |





**BUSINESS RESPONSE FORM**  
**SECOND NOTICE**

**Nebivolol Tablets 2.5 mg**  
**Product Recall – December 6, 2024**  
**Recall # 446**

**Please fill out this form completely.** By doing so, this will acknowledge that you have read and understand the Product Recall instructions and have taken the appropriate action.

Customer Name \_\_\_\_\_ DEA # \_\_\_\_\_

*\*DEA # is required, if it is not provided, the processing of your form will be delayed.*

Address \_\_\_\_\_

City \_\_\_\_\_ State \_\_\_\_\_ Zip \_\_\_\_\_

Contact Name (please print) \_\_\_\_\_ Telephone # \_\_\_\_\_

Contact E-mail : \_\_\_\_\_ Fax # \_\_\_\_\_

Contact Signature \_\_\_\_\_ Date \_\_\_\_\_

**I have checked my stock and:**

\_\_\_\_\_ Do not have any stock of **recalled items**.

**OR**

I have quarantined and listed in the box below the qty. of withdrawn units, I will be returning to QUALANEX, as soon as possible. Upon receipt of this Response Form, QUALANEX will issue return authorization label(s) Please indicate the # of needed box labels \_\_\_\_\_.

| Item Description         | NDC Number   | Lot Number | Sealed Bottle Qty. To Be Returned | Open Bottle Qty. To Be Returned |
|--------------------------|--------------|------------|-----------------------------------|---------------------------------|
| Nebivolol Tablets 2.5 mg | 59651-137-30 | NB0224001A |                                   |                                 |
|                          | 59651-137-30 | NB0224001B |                                   |                                 |



**Recall # 446**

**If you did not purchase the product directly from the Manufacturer, please complete the below section.**

Purchased From: Wholesaler Name \_\_\_\_\_

City \_\_\_\_\_ State \_\_\_\_\_

Wholesaler DEA# \_\_\_\_\_

If you have any questions regarding this form or product return, please contact QUALANEX at 888-504-2014. Office hours 7am to 4pm CST Monday through Friday.

**Please fax this form to: 847-737-3719 Or E-mail [recall@qualanex.com](mailto:recall@qualanex.com)**

