

**FEXOFENADINE HYDROCHLORIDE AND PSEUDOEPHEDRINE HYDROCHLORIDE-  
fexofenadine hydrochloride and pseudoephedrine hydrochloride tablet, film  
coated, extended release  
Ohm Laboratories Inc.**

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**Fexofenadine Hydrochloride and Pseudoephedrine Hydrochloride Extended-  
Release Tablets, 60 mg/120 mg**

***Drug Facts***

<b><i>Active ingredients (in each extended-release tablet)</i></b>	<b><i>Purpose</i></b>
Fexofenadine HCl, USP 60 mg	Antihistamine
Pseudoephedrine HCl, USP 120 mg	Nasal Decongestant

**Uses**

- temporarily relieves these symptoms due to hay fever or other upper respiratory allergies:
  - runny nose
  - sneezing
  - itchy, watery eyes
  - itching of the nose or throat
- temporarily relieves nasal congestion due to the common cold, hay fever or other upper respiratory allergies
- reduces swelling of nasal passages
- temporarily relieves sinus congestion and pressure
- temporarily restores freer breathing through the nose

**Warnings**

**Do not use**

- if you have ever had an allergic reaction to this product or any of its ingredients
- if you are now taking a prescription monoamine oxidase inhibitor (MAOI) (certain drugs for depression, psychiatric, or emotional conditions, or Parkinson's disease), or for 2 weeks after stopping the MAOI drug. If you do not know if your prescription drug contains an MAOI, ask a doctor or pharmacist before taking this product.
- if you have difficulty swallowing

**Ask a doctor before use if you have**

- heart disease
- thyroid disease
- glaucoma
- high blood pressure
- diabetes
- trouble urinating due to an enlarged prostate gland

- kidney disease. Your doctor should determine if you need a different dose.

**When using this product**

- **do not take more than directed**
- do not take at the same time as aluminum or magnesium antacids
- do not take with fruit juices (see Directions)

**Stop use and ask a doctor if**

- an allergic reaction to this product occurs. Seek medical help right away.
- symptoms do not improve within 7 days or are accompanied by a fever
- you get nervous, dizzy, or sleepless

**If pregnant or breast-feeding,**ask a health professional before use.

**Keep out of reach of children.**In case of overdose, get medical help or contact a Poison Control Center right away (1800-222-1222).

**Directions**

- do not divide, crush, chew or dissolve the tablet; swallow tablet whole

adults and children 12 years of age and over	take 1 tablet with a glass of water every 12 hours on an empty stomach; do not take more than 2 tablets in 24 hours
children under 12 years of age	do not use
adults 65 years of age and older	ask a doctor
consumers with kidney disease	ask a doctor

**Other information**

- do not use if carton is opened or if individual blister units are torn or opened
- store between 20° to 25°C (68° to 77°F)
- Meets dissolution test 6

**Inactive ingredients**

colloidal silicon dioxide, croscarmellose sodium, dibasic calcium phosphate dihydrate, ethyl cellulose, ferric oxide yellow, hypromellose, lactose monohydrate, magnesium stearate, microcrystalline cellulose, polyethylene glycol, povidone, stearic acid.

**Questions or comments?**

Call toll-free **1-800-818-4555 weekdays**

Manufactured by:

**Sun Pharmaceutical Industries Limited**

Survey No. 1012, Dadra-396 193,  
U.T. of D & NH and Daman & Diu, India.

Distributed by:

Ohm Laboratories Inc.

New Brunswick, NJ 08901

**PRINCIPAL DISPLAY PANEL - 20 Tablet Blister Pack Carton**

†Compare To  
the active ingredients of  
Allegra-D<sup>®</sup>

NDC 51660-037-21

ohm<sup>®</sup>

NON-DROWSY

Original Prescription Strength

Fexofenadine HCl 60 mg/Antihistamine  
Pseudoephedrine HCl 120 mg/Nasal Decongestant  
Extended-Release Tablets, USP

Allergy & Congestion

Indoor and Outdoor Allergies

DO NOT USE IF INDIVIDUAL BLISTER UNIT IS OPEN OR TORN

12 Hour

Relief of:

- Nasal and Sinus Congestion  
Due to Colds or Allergies
- Sneezing; Runny Nose; Itchy,  
Watery Eyes and Itchy Nose or  
Throat Due to Allergies

20 Extended-Release Tablets

NO GLUE - NO COATING

NO COATING

NVZ

**Drug Facts (continued)**

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NVZ

NO COATING

5266727



18025



Extended-Release Tablets, USP  
Pseudoephedrine HCl 120 mg/Nasal Decongestant  
Fexofenadine HCl 60 mg/Antihistamine

a SUN PHARMA company



**ohm**  
LABORATORIES INC.  
a SUN PHARMA company

**Compare To**  
the active ingredients of  
**Allegra-D®**

NDC 51660-037-21

**NON-DROWSY**  
**Fexofenadine HCl 60 mg/Antihistamine**  
**Pseudoephedrine HCl 120 mg/Nasal Decongestant**  
**Extended-Release Tablets, USP**

**Allergy & Congestion**

**12 Hour**  
*Indoor/Outdoor Relief of:*

- Nasal and Sinus Congestion Due to Colds or Allergies
- Sneezing; Runny Nose; Itchy, Watery Eyes and Itchy Nose or Throat Due to Allergies

**20 Extended-Release Tablets**

NON VARNISH  
NO INK AREA

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4 x 5 Tablets

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**Sun Pharmaceutical Industries Limited**  
Survey No. 1012, Dadra-366 193,  
U.T. of D & NH and Daman & Diu, India.  
Distributed by: Ohm Laboratories Inc.  
New Brunswick, NJ 08901

# FEXOFENADINE HYDROCHLORIDE AND PSEUDOEPHEDRINE HYDROCHLORIDE

fexofenadine hydrochloride and pseudoephedrine hydrochloride tablet, film coated, extended release

## Product Information

<b>Product Type</b>	HUMAN OTC DRUG	<b>Item Code (Source)</b>	NDC:51660-037
<b>Route of Administration</b>	ORAL		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>FEXOFENADINE HYDROCHLORIDE</b> (UNII: 2S068B75ZU) (FEXOFENADINE - UNII:E6582LOH6V)	FEXOFENADINE HYDROCHLORIDE	60 mg
<b>PSEUDOEPHEDRINE HYDROCHLORIDE</b> (UNII: 6V9V2RYJ8N) (PSEUDOEPHEDRINE - UNII:7CUC9DDI9F)	PSEUDOEPHEDRINE HYDROCHLORIDE	120 mg

## Inactive Ingredients

Ingredient Name	Strength
<b>LACTOSE MONOHYDRATE</b> (UNII: EWQ57Q8I5X)	
<b>MICROCRYSTALLINE CELLULOSE</b> (UNII: OP1R32D61U)	
<b>CROSCARMELLOSE SODIUM</b> (UNII: M28OL1HH48)	
<b>POVIDONE K30</b> (UNII: U725QWY32X)	
<b>MAGNESIUM STEARATE</b> (UNII: 70097M6I30)	
<b>DIBASIC CALCIUM PHOSPHATE DIHYDRATE</b> (UNII: O7TSZ97GEP)	
<b>SILICON DIOXIDE</b> (UNII: ETJ7Z6XBU4)	
<b>ETHYLCELLULOSE, UNSPECIFIED</b> (UNII: 7Z8S9VYZ4B)	
<b>FERRIC OXIDE YELLOW</b> (UNII: EX438O2MRT)	
<b>STEARIC ACID</b> (UNII: 4ELV7Z65AP)	
<b>HYPROMELLOSE, UNSPECIFIED</b> (UNII: 3NXW29V3WO)	
<b>POLYETHYLENE GLYCOL, UNSPECIFIED</b> (UNII: 3WJQ0SDW1A)	

## Product Characteristics

<b>Color</b>	white, yellow	<b>Score</b>	no score
<b>Shape</b>	CAPSULE (bilayer)	<b>Size</b>	17mm
<b>Flavor</b>		<b>Imprint Code</b>	724
<b>Contains</b>			

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:51660-037-21	1 in 1 CARTON	03/01/2018	
1		20 in 1 BLISTER PACK; Type 0: Not a Combination Product		
	NDC:51660			

2	NDC:51660-037-31	1 in 1 CARTON	03/01/2018	
2		30 in 1 BLISTER PACK; Type 0: Not a Combination Product		
Marketing Information				
Marketing Category	Application Number or Monograph Citation		Marketing Start Date	Marketing End Date
ANDA	ANDA090818		03/01/2018	

**Labeler** - Ohm Laboratories Inc. (184769029)

Establishment			
Name	Address	ID/FEI	Business Operations
Sun Pharmaceutical Industries Limited		650445203	analysis(51660-037) , manufacture(51660-037)

Revised: 11/2025

Ohm Laboratories Inc.