

**ALLERGY RELIEF- diphenhydramine hcl tablet, film coated**  
**L.N.K. International, Inc.**

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**Allergy Relief**

***Active ingredient (in each tablet)***

Diphenhydramine HCl 25 mg

***Purpose***

Antihistamine

***Uses***

- temporarily relieves these symptoms due to hay fever or other upper respiratory allergies:
  - runny nose
  - itchy, watery eyes
  - sneezing
  - itching of the nose or throat
- temporarily relieves these symptoms due to the common cold:
  - runny nose
  - sneezing

***Warnings***

**Do not use**

- to make a child sleepy
- with any other product containing diphenhydramine, even one used on skin

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

- a breathing problem such as emphysema or chronic bronchitis
- glaucoma
- difficulty in urination due to enlargement of the prostate gland

**Ask a doctor or pharmacist before use if you are**

taking sedatives or tranquilizers.

**When using this product**

- marked drowsiness may occur
- avoid alcoholic beverages
- alcohol, sedatives, and tranquilizers may increase drowsiness
- use caution when driving a motor vehicle or operating machinery
- excitability may occur, especially in children

**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.

**Directions**

- **do not take more than directed**
- take every 4 to 6 hours, or as directed by a doctor
- do not take more than 6 times in 24 hours

adults and children 12 years and over	1 to 2 tablets
children 6 to under 12 years	1 tablet
children under 6 years	do not use

**Other information**

- **each tablet contains:** calcium 30 mg
- **TAMPER EVIDENT: DO NOT USE IF OUTER PACKAGE IS OPENED OR BLISTER IS TORN OR BROKEN**
- store at 25°C (77°F); excursions permitted between 15°-30°C (59°-86°F)
- protect from moisture
- see end flap for expiration date and lot number

**Inactive ingredients**

corn starch, D&C red #27 aluminum lake, dibasic calcium phosphate dihydrate, magnesium stearate, microcrystalline cellulose, polyethylene glycol, polyvinyl alcohol, silicon dioxide, stearic acid, talc, titanium dioxide

**Questions or comments?**

**1-800-426-9391**

**Principal Display Panel**

NDC 50844-329-07

**QUALITY  
+ PLUS**

\*Compare to active ingredient in Benadryl® Allergy ULTRATAB® Tablets

**ALLERGY  
RELIEF**

Diphenhydramine HCl 25 mg  
ANTI-HISTAMINE

Allergy Relief for:

- Sneezing
- Itchy, Watery Eyes
- Runny Nose
- Itchy Throat

**36 Minitabs**

ACTUAL SIZE

**TAMPER EVIDENT: DO NOT USE IF PACKAGE IS  
OPENED OR IF BLISTER UNIT IS TORN, BROKEN  
OR SHOWS ANY SIGNS OF TAMPERING**

\*This product is not manufactured or distributed by Johnson  
& Johnson Corporation, owner of the registered trademark  
Benadryl® Allergy ULTRATAB® Tablets.

50844            REV0721G32907

Distributed by

**LNK INTERNATIONAL, INC.**

60 Arkay Drive

Hauppauge, NY 11788

USA

**Drug Facts**

**Active ingredient**  
(in each tablet)  
Diphenhydramine HCl 25 mg. . . . . Antihistamine

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**Uses**

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KEEP OUTER PACKAGE FOR COMPLETE PRODUCT INFORMATION

**QUALITY PLUS**

# ALLERGY RELIEF

Diphenhydramine HCl 25 mg

36 Minitabs

ANTIHISTAMINE

B-1603-329-07-R  
REV0721G32907

**QUALITY PLUS**

NDC 50844-329-07

\*Compare to active ingredient in Benadryl® Allergy-ULTRATAB® Tablets

# ALLERGY RELIEF

Diphenhydramine HCl 25 mg

ANTIHISTAMINE

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- Runny Nose
- Itchy, Watery Eyes
- Itchy Throat

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ACTUAL SIZE 

36 Minitabs

**QUALITY PLUS**

**ALLERGY RELIEF**

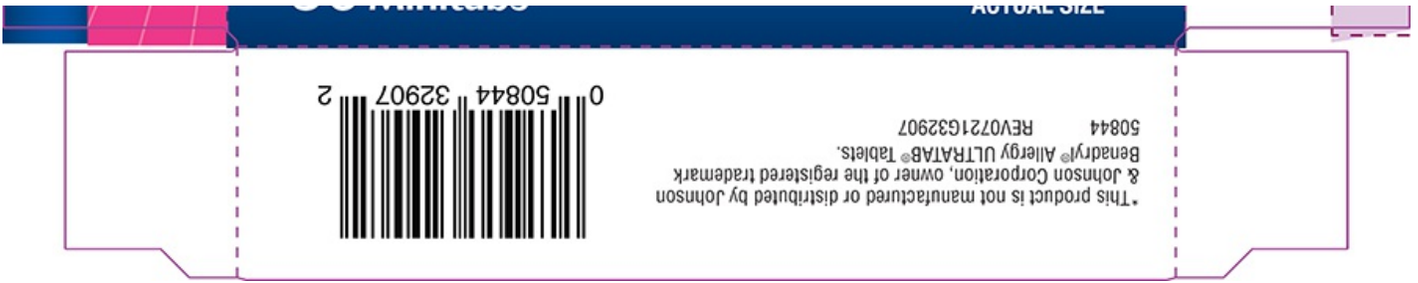
Diphenhydramine HCl 25 mg

ANTIHISTAMINE

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Distributed by  
**LNK INTERNATIONAL, INC.**  
60 Arkay Drive  
Hauppauge, NY 11788  
USA

No Print Area Lot no. & Exp. date



**Quality Plus 44-329**

**ALLERGY RELIEF**

diphenhydramine hcl tablet, film coated

**Product Information**

<b>Product Type</b>	HUMAN OTC DRUG	<b>Item Code (Source)</b>	NDC:50844-329
<b>Route of Administration</b>	ORAL		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>DIPHENHYDRAMINE HYDROCHLORIDE</b> (UNII: TC2D6JAD40) (DIPHENHYDRAMINE - UNII:8GTS82S83M)	DIPHENHYDRAMINE HYDROCHLORIDE	25 mg

**Inactive Ingredients**

Ingredient Name	Strength
<b>STARCH, CORN</b> (UNII: O8232NY3SJ)	
<b>D&amp;C RED NO. 27 ALUMINUM LAKE</b> (UNII: ZK64F7XSTX)	
<b>DIBASIC CALCIUM PHOSPHATE DIHYDRATE</b> (UNII: O7TSZ97GEP)	
<b>MAGNESIUM STEARATE</b> (UNII: 70097M6I30)	
<b>MICROCRYSTALLINE CELLULOSE</b> (UNII: OP1R32D61U)	
<b>POLYETHYLENE GLYCOL, UNSPECIFIED</b> (UNII: 3WJQ0SDW1A)	
<b>POLYVINYL ALCOHOL, UNSPECIFIED</b> (UNII: 532B59J990)	
<b>SILICON DIOXIDE</b> (UNII: ETJ7Z6XBU4)	
<b>STEARIC ACID</b> (UNII: 4ELV7Z65AP)	
<b>TALC</b> (UNII: 7SEV7J4R1U)	
<b>TITANIUM DIOXIDE</b> (UNII: 15FIX9V2JP)	

**Product Characteristics**

<b>Color</b>	pink	<b>Score</b>	no score
<b>Shape</b>	OVAL	<b>Size</b>	11mm
<b>Flavor</b>		<b>Imprint Code</b>	44;329
<b>Contains</b>			

**Packaging**

		<b>Marketing Start</b>	<b>Marketing End</b>
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#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:50844-329-02	1 in 1 CARTON	03/02/1990	
1		12 in 1 BLISTER PACK; Type 0: Not a Combination Product		
2	NDC:50844-329-08	2 in 1 CARTON	03/02/1990	
2		12 in 1 BLISTER PACK; Type 0: Not a Combination Product		
3	NDC:50844-329-07	3 in 1 CARTON	03/02/1990	
3		12 in 1 BLISTER PACK; Type 0: Not a Combination Product		
4	NDC:50844-329-22	4 in 1 CARTON	03/02/1990	
4		12 in 1 BLISTER PACK; Type 0: Not a Combination Product		
5	NDC:50844-329-12	100 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	03/02/1990	
6	NDC:50844-329-51	365 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	03/02/1990	
7	NDC:50844-329-82	1 in 1 CARTON	03/02/1990	
7		100 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product		
8	NDC:50844-329-78	600 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	03/02/1990	

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M012	03/02/1990	

**Labeler** - L.N.K. International, Inc. (038154464)

## Establishment

Name	Address	ID/FEI	Business Operations
LNK International, Inc.		038154464	pack(50844-329)

## Establishment

Name	Address	ID/FEI	Business Operations
LNK International, Inc.		832867837	manufacture(50844-329) , pack(50844-329)

## Establishment

Name	Address	ID/FEI	Business Operations
LNK International, Inc.		832867894	manufacture(50844-329)

## Establishment

Name	Address	ID/FEI	Business Operations
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LNK International, Inc.		868734088	manufacture(50844-329)
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## Establishment

Name	Address	ID/FEI	Business Operations
LNK International, Inc.		967626305	pack(50844-329)

Revised: 5/2025

L.N.K. International, Inc.