RANITIDINE - ACID REDUCER- ranitidine hydrochloride tablet, film coated Chain Drug Consortium, LLC.

Drug Facts

ACTIVE INGREDIENT (IN EACH TABLET)

Ranitidine 75 mg (as ranitidine hydrochloride USP, 84 mg)

PURPOSE

Acid reducer

USES

- relieves heartburn associated with acid indigestion and sour stomach
- prevents heartburn associated with acid indigestion and sour stomach brought on by eating or drinking certain foods and beverages

WARNINGS

Allergy alert: Do not use if you are allergic to ranitidine or other acid reducers

Do not use

- if you have trouble or pain swallowing food, vomiting with blood, or bloody or black stools. These may be signs of a serious condition. See your doctor.
- with other acid reducers

Ask a doctor before use if you have

- frequent **chest pain**
- frequent wheezing, particularly with heartburn
- unexplained weight loss
- nausea or vomiting
- stomach pain
- had heartburn over 3 months. This may be a sign of a more serious condition.
- heartburn with lightheadedness, sweating or dizziness
- chest pain or shoulder pain with shortness of breath; sweating; pain spreading to arms, neck or shoulders; or lightheadedness

Stop use and ask a doctor if

- your heartburn continues or worsens
- you need to take this product for more than 14 days

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 (1-800-222-1222).

DIRECTIONS

- adults and children 12 years and over:
 - to **relieve** symptoms, swallow 1 tablet with a glass of water
 - to **prevent** symptoms, swallow 1 tablet with a glass of water **30 to 60 minutes before** eating food or drinking beverages that cause heartburn
 - can be used up to twice daily (do not take more than 2 tablets in 24 hours)
- children under 12 years: ask a doctor

OTHER INFORMATION

- TAMPER EVIDENT: DO NOT USE IF THE CARTON OR PRINTED FOIL UNDER CAP IS OPEN OR TORN.
- store at 20° 25° C (68° 77° F)
- avoid excessive heat or humidity
- this product is sodium and sugar free

INACTIVE INGREDIENTS

colloidal silicon dioxide, croscarmellose sodium, hypromellose, iron oxide red, magnesium stearate, microcrystalline cellulose, polyethylene glycol 400, talc, titanium dioxide

QUESTIONS?

call **1-800-406-7984**

Read the directions, consumer information leaflet and warnings before use.

Keep the carton. It contains important information.

DISTRIBUTED BY

CHAIN DRUG CONSORTIUM

3301 NW BOCA RATON BLVD

SUITE 101, BOCA RATON, FL 33431

PRINCIPAL DISPLAY PANEL

Premier Value[®] NDC 68016-352-60 Ranitidine Tablets, USP 75 mg Acid Reducer Regular Strength 60 Tablets Prevents & Relieves Heartburn Associated with Acid Indigestion & Sour Stomach

 $^{\dagger}\text{Compare}$ to the active ingredient of Zantac 75^{\circledast}

 $^\dagger This$ product is not manufactured or distributed by Boehringer Ingelheim Pharmaceuticals Inc., owner of the registered trademark Zantac $75^{\&}$.



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Drug Facts (continued)	i

RANITIDINE - ACID REDUCER									
ra	nitidine hydrochlori	de tablet, film	n coated						
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P	roduct Informati	ion							
Р	roduct T ype		HUMAN OTC DR	UG	Ite m C	ode (Source)		NDC:68016-	352
R	oute of Administrat	ion	ORAL						
A	ctive Ingredient/	Active Moi/	ety						
		I	ngredient Nam	e			Basis o	f Strength	Strength
R	ANITIDINE HYDROC	HLORIDE (UN	II: BK76465IHM) (R	RANITIDINE - U	NII:884	KT10 YB7)	RANITID	INE	75 mg
_									
I	nactive Ingredier	nts							
			Ingredien	t Name				Sti	rength
SI	LICON DIO XIDE (UN	NII: ETJ7Z6XBU	J4)						
C	ROSCARMELLOSES	SODIUM (UNII:	M28OL1HH48)						
H	YPROMELLOSES (U	NII: 3NXW29V3	3WO)						
F	ERRIC OXIDE RED (U	JNII: 1K09F3G6	75)						
Μ	AGNESIUM STEARA	TE (UNII: 7009	7M6I30)						
C	ELLULOSE, MICROO	CRYSTALLINE	E (UNII: OP1R32D61	lU)					
-	DLYETHYLENE GLY		II: B697894SGQ)						
	ALC (UNII: 7SEV7J4R								
T	TANIUM DIO XIDE (U	JNII: 15FIX9V2J	JP)						
-									
	roduct Characte								
		pink		Score		Marketing Start Date Marketing Marketing Marketing			
		ROUN	D	Size					
Flavor				Imprint Cod	le		DR;606		
C	ontains								
Packaging									
			Package Description Marketing 9			76 1			
#	Item Code				art Date	Marketing	, End Date		
-	NDC:68016-352-60 NDC:68016-352-30	60 in 1 BOTTLE; Type 0: Not a Combination Product 30 in 1 BOTTLE; Type 0: Not a Combination Product							
2	1100.00010-352-30		in, Type 0. Not a Co		uuct				

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
ANDA	ANDA201745	07/10/2012			

Labeler - Chain Drug Consortium, LLC. (101668460)

Registrant - Ohm Laboratories Inc. (184769029)

Establishment					
Name	Address	ID/FEI	Business Operations		
Shasun Pharmaceuticals Limited		915786829	MANUFACTURE(68016-352)		

Revised: 10/2015

Chain Drug Consortium, LLC.