LOPERAMIDE HYDROCHLORIDE- loperamide hydrochloride tablet Select Brand

Drug Facts

ACTIVE INGREDIENT (IN EACH CAPLET)

Loperamide HCI USP, 2 mg

PURPOSE

Anti-diarrheal

USES

Controls symptoms of diarrhea, including Travelers' Diarrhea

WARNINGS

Allergy alert: Do not use if you have ever had a rash or other allergic reaction to loperamide HCI

Do not use

If you have bloody or black stool

Ask a doctor before use if you have

- fever
- mucus in the stool
- a history of liver disease

Ask a doctor or pharmacist before use if you are

Taking antibiotics

When using this product

• tiredness, drowsiness or dizziness may occur. Be careful when driving or operating machinery.

Stop use and ask a doctor if

- diarrhea lasts for more than 2 days
- symptoms get worse
- you get abdominal swelling or bulging. These may be signs of a serious condition.

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

- drink plenty of clear fluids to help prevent dehydration caused by diarrhea
- find right dose on chart. If possible, use weight to dose; otherwise, use age.

adults and children 12 years and over	2 caplets after the first loose stool; 1 caplet after each subsequent loose stool; but no more than 4 caplets in 24 hours
children 9-11 years (60-95 lbs)	1 caplet after the first loose stool; ¹ / ₂ caplet after each subsequent loose stool; but no more than 3 caplets in 24 hours
children 6-8 years (48-59 lbs)	1 caplet after the first loose stool; ¹ / ₂ caplet after each subsequent loose stool; but no more than 2 caplets in 24 hours
children under 6 years (up to 47 lbs)	ask a doctor

OTHER INFORMATION

- store between $20^{\circ} 25^{\circ} \text{ C} (68^{\circ} 77^{\circ} \text{ F})$
- see side panel for lot number and expiration date
- TAMPER EVIDENT: THIS PRODUCT PROTECTED WITH SEALED BLISTER UNITS. DO NOT USE IF ANY ARE TORN OR BROKEN.

INACTIVE INGREDIENTS

Anhydrous lactose, croscarmellose sodium, crospovidone, D&C yellow no.10, FD&C blue no.1, hydrogenated vegetable oil, magnesium stearate, powdered cellulose, pregelatinized starch

QUESTIONS?

Call 1800-406-7984

PRINCIPAL DISPLAY PANEL select brand® NDC 15127-338-12 Loperamide Hydrochloride Tablets USP, 2 mg ANTI-DIARRHEAL Controls The Symptoms Of Diarrhea Compare to the active ingredient of Imodium®A-D 12 Caplets* Each Caplet (*capsule-shaped tablet) contains Loperamide HCl USP, 2 mg 5048836/R1109



LOPERAMIDE	HYDROG	CHLORIDE							
loperamide hydrochlor									
Product Information	on								
Product Type		HUMAN OTC DRUG Item Code (Source)			NDC:15127-338				
Route of Administration	on	ORAL							
Active Ingredient/A	Active Moi	ety							
	Ingr	edient Name				Basi	is of St	rength	Strengtl
LOPERAMIDE HYDROC UNII:6 X9 OC3H4II)	C HLORIDE (U	NII: 77TI35393C) (LOPERAMIDE - LOPERAMIDE HYDROCHLORIDE					2 mg		
Inactive Ingredient	ts								
		Ingredient Na	ame					St	rength
ANHYDROUS LACTOS	E (UNII: 3SY5L	H9 PMK)							
CROSCARMELLOSE SO	DDIUM (UNII:	M28OL1HH48)							
CROSPOVIDONE (UNII:	68401960MK)							
D&C YELLOW NO. 10 (UNII: 35SW5U	SQ3G)							
FD&C BLUE NO. 1 (UNII: H3R47K3TBD)									
TRISTEARIN (UNII: P6O	CJ2551R)								
MAGNESIUM STEARAT	E (UNII: 7009)	7M6I30)							
POWDERED CELLULO	SE (UNII: SMD	1X3XO9M)							
STARCH, PREGELATIN	IZED CORN (UNII: O8232NY3SJ)							
Product Character	istics								
Color	green	Score 2 I			2 pieces	pieces			
Shape	CAPSU	LE	Size 91			9mm	mm		
Flavor		Imprint Code				123			
Contains			-						
Packaging									
# Item Code	Pack	age Description	Mar	rketin	g Start	Date	Ma	arketing E	nd Date
1 NDC:15127-338-66	6 in 1 BLI	STER PACK							
2 NDC:15127-338-12	12 in 1 BL	ISTER PACK							
Marketing Info	rmation								
Marketing Category		on Number or Monograph Citation		tion	ion Marketing Start Date		Marketing End Date		
ANDA	ANDA074091				02/01/1993				

Labeler - Select Brand (043562370)

Registrant - Ranbaxy Pharmaceuticals Inc. (937890044)

Establishment

Name	Address	ID/FEI	Business Operations
Ohm Laboratories Inc.		051565745	manufacture(15127-338)

Revised: 9/2012

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