

OMEPRAZOLE- omeprazole tablet, delayed release
Sun Pharmaceutical Industries, Inc.

Omeprazole Delayed-Release Tablets, 20 mg

Active ingredient(in each tablet)

Omeprazole USP 20 mg

Purpose

Acid reducer

Use

- treats frequent heartburn (occurs **2 or more** days a week)
- not intended for immediate relief of heartburn; this drug may take 1 to 4 days for full effect

Warnings

Allergy alert:

do not use if you are allergic to omeprazole

- omeprazole may cause severe skin reactions. Symptoms may include:
• skin reddening • blisters • rash

If an allergic reaction occurs, stop use and seek medical help right away.

Do not use

if you have:

- trouble or pain swallowing food, vomiting with blood, or bloody or black stools
- 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
- frequent **chest pain**

These may be signs of a serious condition. See your doctor.

Ask a doctor before use if you have:

- had heartburn over 3 months. This may be a sign of a more serious condition.
- frequent wheezing, particularly with heartburn
- unexplained weight loss
- nausea or vomiting
- stomach pain

Ask a doctor or pharmacist before use if you are

taking a prescription drug. Acid reducers may interact with certain prescription drugs.

Stop use and ask a doctor if

- your heartburn continues or worsens
- you need to take this product for more than 14 days
- you need to take more than 1 course of treatment every 4 months
- you get diarrhea
- you develop a rash or joint pain

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

- for adults 18 years of age and older
- this product is to be used once a day (every 24 hours), every day for 14 days
- it may take 1 to 4 days for full effect; some people get complete relief of symptoms within 24 hours

14-Day Course of Treatment

- swallow 1 tablet with a glass of water before eating in the morning
- take every day for 14 days
- do not take more than 1 tablet a day
- do not use for more than 14 days unless directed by your doctor
- swallow whole. Do not chew or crush tablets.

Repeated 14-Day Courses (if needed)

- you may repeat a 14-day course every 4 months
- **do not take for more than 14 days or more often than every 4 months unless directed by a doctor**
- children under 18 years of age: ask a doctor. Heartburn in children may sometimes be caused by a serious condition.

Other information

- read the directions and warnings before use
- keep the carton. It contains important information.
- store at 20 to 25° C (68 to 77° F) and protect from moisture

Inactive ingredients

anhydrous lactose, hypromellose, hypromellose acetate succinate, iron oxide red, iron oxide yellow, lactose monohydrate, methyl cellulose, monoethanolamine, propylene glycol, sodium lauryl sulfate, sodium starch glycolate, sodium stearate, sodium stearyl fumarate, talc, triethyl citrate and titanium dioxide.

The imprinting ink contains ammonium hydroxide, black iron oxide, n-butyl alcohol, propylene glycol and shellac.

Questions or Comments?

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

Tips of Managing Heartburn

- Do not lie flat or bend over after eating
- Do not wear tight-fitting clothing around the stomach
- Do not eat before bedtime
- Raise the head of your bed
- Avoid heartburn-causing foods such as rich, spicy, fatty or fried foods, chocolate, caffeine, alcohol and certain fruits and vegetables
- Eat slowly and avoid big meals
- If overweight, lose weight
- Quit smoking

Principal Display Panel - Foil

62756-377-40

Omeprazole Delayed-Release Tablet

20 mg

Acid Reducer

PUSH THROUGH

Take 1 tablet a day for 14 days

Mfg. by: **Sun Pharmaceutical Ind Ltd.**, India.

Dist. by: **Sun Pharmaceutical Ind Inc.**, NJ 08512

PGPF0520A

KEEP OUTER CARTON FOR COMPLETE WARNINGS AND PRODUCT INFORMATION, READ WARNINGS AND DIRECTIONS ON CARTON BEFORE USE.


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Swallow whole. Do not chew or crush tablets.
THIS SINGLE 14-TABLETS BLISTER PACKAGE
CONTAINS ONE 14-DAY COURSE OF TREATMENT.
DO NOT TAKE FOR MORE THAN 14 DAYS OR
MORE OFTEN THAN EVERY 4 MONTHS UNLESS
DIRECTED BY A DOCTOR.
LOT
EXP


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Principal Display Panel - Blister Carton

NDC 62756-377-96

Compare to Prilosec OTC ® *

Treats Frequent Heartburn! (24 HR)

Compare to Prilosec OTC ® *

Omeprazole Delayed-release Tablets

20 mg

Acid Reducer

SWALLOW - DO NOT CHEW
28 (2 x 14) Unit-Dose Tablets
TWO 14-DAY COURSES OF TREATMENT
May take 1 to 4 days for full effect



Principal Display Panel - Label

NDC 62756-377-21
Treats **Frequent Heartburn!** (24 HR)
Omeprazole Delayed-release Tablets
20 mg
Acid Reducer
14 Tablets
ONE 14-DAY COURSE OF TREATMENT
May take 1 to 4 days for full effect



Principal Display Panel - Bottle Carton

NDC 62756-377-11
Compare to Prilosec OTC ® *
Treats **Frequent Heartburn!** (24 HR)
Omeprazole Delayed-release Tablets
20 mg

Acid Reducer
SWALLOW - DO NOT CHEW
28 Tablets
TWO 14-DAY COURSES OF TREATMENT
May take 1 to 4 days for full effect



OMEPRAZOLE

omeprazole tablet, delayed release

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:62756-377
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
OMEPRAZOLE (UNII: KG60484QX9) (OMEPRAZOLE - UNII:KG60484QX9)	OMEPRAZOLE	20 mg

Inactive Ingredients

Ingredient Name	Strength
ANHYDROUS LACTOSE (UNII: 3S5Y5LH9PMK)	
HYPROMELLOSE, UNSPECIFIED (UNII: 3NXW29V3WO)	
HYPROMELLOSE ACETATE SUCCINATE 12070923 (3 MM2/S) (UNII: 36BGF0E889)	
FERRIC OXIDE RED (UNII: 1K09F3G675)	
FERRIC OXIDE YELLOW (UNII: EX438O2MRT)	
LACTOSE MONOHYDRATE (UNII: EWQ57Q8I5X)	
MONOETHANOLAMINE (UNII: 5KV86114PT)	
METHYLCELLULOSE (1500 MPA.S) (UNII: PONTE48364)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
SODIUM STARCH GLYCOLATE TYPE A POTATO (UNII: 5856J3G2A2)	
SODIUM STEARATE (UNII: QU7E2XA9TG)	
SODIUM STEARYL FUMARATE (UNII: 7CV7WJK4UI)	
SODIUM LAURYL SULFATE (UNII: 368GB5141J)	
TRIETHYL CITRATE (UNII: 8Z96QXD6UM)	

TALC (UNII: 7SEV7J4R1U)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	
AMMONIA (UNII: 5138Q19F1X)	
FERROSFERRIC OXIDE (UNII: XM0M87F357)	
BUTYL ALCOHOL (UNII: 8PJ61P6TS3)	
SHELLAC (UNII: 46N107B71O)	

Product Characteristics

Color	BROWN (brownish pink)	Score	no score
Shape	OVAL (biconvex)	Size	12mm
Flavor		Imprint Code	20
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:62756-377-21	1 in 1 CARTON	01/01/2019	
1		14 in 1 BOTTLE; Type 0: Not a Combination Product		
2	NDC:62756-377-11	2 in 1 CARTON	01/01/2019	
2		14 in 1 BOTTLE; Type 0: Not a Combination Product		
3	NDC:62756-377-12	3 in 1 CARTON	01/01/2019	
3		14 in 1 BOTTLE; Type 0: Not a Combination Product		
4	NDC:62756-377-70	1 in 1 CARTON	01/01/2019	
4		14 in 1 BLISTER PACK; Type 0: Not a Combination Product		
5	NDC:62756-377-96	2 in 1 CARTON	01/01/2019	
5		14 in 1 BLISTER PACK; Type 0: Not a Combination Product		
6	NDC:62756-377-79	3 in 1 CARTON	01/01/2019	
6		14 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	ANDA207891	01/01/2019	

Labeler - Sun Pharmaceutical Industries, Inc. (146974886)

Establishment

Name	Address	ID/FEI	Business Operations
Sun Pharmaceutical Industries Limited		650445203	ANALYSIS(62756-377) , MANUFACTURE(62756-377)

Revised: 4/2025

Sun Pharmaceutical Industries, Inc.