QCH MAXIMUM STRENGTH ANTACID 516- aluminum hydroxide, magnesium hydroxide, simethicone liquid Chain Drug Marketing Association Inc.

QCH Maximum Strength Antacid 516

ACTIVE INGREDIENTS (in each 10 mL)

Aluminum hydroxide (equiv. to dried gel, USP) 800 mg Magnesium hydroxide 800 mg Simethicone 80 mg

PURPOSE

Antacid Antacid Antigas

USE(S)

relieves:

- acid indigestion
- heartburn
- sour stomach
- upset stomach and gas associated with these symptoms

WARNINGS

Do not take more than 60 mL in a 24-hour period, or use the maximum dosage of this product for more than 2 weeks, except under the advice and supervision of a doctor.

ASK A DOCTOR BEFORE USE IF YOU HAVE

- kidney disease
- a magnesium-restricted diet

ASK A DOCTOR OR PHARMACIST BEFORE USE IF YOU ARE

• presently taking a prescription drug. Antacids may interact with certain prescription drugs.

STOP USE AND ASK DOCTOR

if symptoms last more than two weeks

DIRECTIONS

- shake well before use
- adults and children 12 years and older: take 10 mL 20 mL two times a day, or as directed by a doctor
- children under 12 years: consult a doctor
- mL = milliliter

OTHER INFORMATION

- each 10 mL contains: magnesium 340 mg, sodium 5 mg
- store at 20°C-25°C (68°-77°F)
- do not freeze

INACTIVE INGREDIENTS

ethyl alcohol, flavor, glycerin, hydroxyethyl cellulose, methylparaben, propylene glycol, propylparaben, purified water, saccharin sodium, simethicone emulsion, sorbitol

PRINCIPAL DISPLAY PANEL

NDC 83324-122-12

QC

QUALITY CHOICE

*Compare to the Active Ingredients in Mylanta® Maximum Strength

Maximum Strength

Antacid Liquid

Antacid & Antigas

Aluminum Hydroxide 800 mg Magnesium Hydroxide 800 mg Simethicone 80 mg

Relieves:

Acid Indigestion

Heartburn

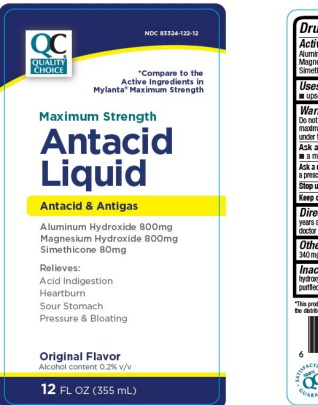
Sour Stomach

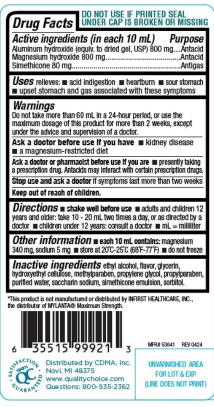
Pressure & Bloating

Original Flavor

Alcohol content 0.2% v/v

12 FL OZ (355 mL)





QCH MAXIMUM STRENGTH ANTACID 516

aluminum hydroxide, magnesium hydroxide, simethicone liquid

Product Information						
Product Type	HUMAN OTC DRUG	Item Code (Source)		NDC:83324-122		
Route of Administration	ORAL					
Active Ingredient/Active Moiety						
Ingredient Name			Basis o Strengt	- Strength		
ALUMINUM HYDROXIDE (UNII: 5QB0T2IUN0) (ALUMINUM HYDROXIDE - UNII:5QB0T2IUN0)			ALUMINUM HYDROXIDE	800 mg in 10 mL		
MAGNESIUM HYDROXIDE (UNII: NBZ 3QY004S) (MAGNESIUM CATION - UNII:T6V3LHY838)			MAGNESIUM HYDROXIDE	800 mg in 10 mL		
DIMETHICONE (UNII: 92RU3N3Y1O) (DIMETHICONE - UNII:92RU3N3Y1O)		DIMETHICONE	80 mg in 10 mL			

Ingredient Name				
ALCOHOL (UNII: 3K	9958V90M)			
GLYCERIN (UNII: PD	C6A3C0OX)			
HYDROXYETHYL C	ELLULOSE (2000 MPA.S AT 1%) (UNII: S38J6R	RZN16)		
METHYLPARABEN	UNII: A2I8C7HI9T)			
PROPYLENE GLYCO	DL (UNII: 6DC9Q167V3)			
PROPYLPARABEN (UNII: Z8IX2SC1OH)			
WATER (UNII: 059Q)	FOKOOR)			
SACCHARIN SODIU	M (UNII: SB8ZUX40TY)			
SORBITOL (UNII: 50	6T60A25R)			
Product Chara				
Color	WHITE		Score	
Shape		Size		
Flavor	MINT (Lemon mint)	Imprint Code	Imprint Code	
Contains				
Packaging				
# Item Code	Package Description	Marketing Start Date	Marketing End Date	
	355 mL in 1 BOTTLE; Type 0: Not a Combinatior Product	n 08/28/2024		
Marketing I	nformation			
Marketing I Marketing Category	nformation Application Number or Monograph Citation	n Marketing Start Date	Marketing End Date	

Labeler - Chain Drug Marketing Association Inc. (011920774)

Establishment							
Name	Address	ID/FEI	Business Operations				
Guardian Drug Company		119210276	MANUFACTURE(83324-122)				

Revised: 8/2024

Chain Drug Marketing Association Inc.