

KESIN MAG-AL PLUS- aluminum and magnesium hydroxide w/simethicone suspension
Kesin Pharma Corporation

Kesin MAG-AL™ Plus (Aluminum & Magnesium Hydroxide w/Simethicone) Suspension

Drug Facts

Active ingredient (in each 5 mL = 1 teaspoonful).....
.....Purpose

Aluminum Hydroxide 200
mg.....Antacid

Magnesium Hydroxide 200
mg.....Antacid

Simethicone 20
mg.....Antigas

Uses

For the relief of:

- acid indigestion
- heartburn
- sour stomach
- upset stomach due to these symptoms
- pressure and bloating commonly referred to as gas

Warnings

Do not take more than 16 teaspoonfuls in a 24-hour period or use the maximum dosage for more than 2 weeks except under the advice and supervision of a physician.

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 a doctor if

- symptoms last more than 2 weeks

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

- shake well before using
- do not take more than 16 teaspoonfuls in 24 hours or use the maximum dosage for more than 2 weeks

Age	Dose
adults and children 12 years of age and older	take 2 to 4 teaspoonfuls four times a day or as directed by a physician
children under 12 years of age	consult a physician

Other information

- **each teaspoonful (5 mL) contains:**magnesium 200 mg, sodium < 1mg
- store at 20°C to 25°C (68°F to 77°F)
- Avoid freezing
- Kesin MAG-AL Plus is a milky-white mint flavored suspension supplied in the following oral dosage forms:

NDC 81033-006-30: 30 mL unit-dose cup

NDC 81033-006-01: Case containing 100 unit-dose cups of 30 mL each.

Inactive ingredients

colloidal silicon dioxide, glycerin, methylparaben, microcrystalline cellulose, mint flavor, propylene glycol, propylparaben, purified water, sorbitol, sucralose, xanthan gum

Questions or comments?

Call 1-833-537-4679

Distributed by:

Kesin Pharma

Oldsmar, FL 34677

PIS-006-V01

Rev. 10/2025



PRINCIPAL DISPLAY PANEL - 30 mL Dose Cup Tray Label

NDC: 81033-006-01

Kesin MAG-AL TM 1200 mg/1200 mg/120 mg per 30 mL

(Aluminum & Magnesium Hydroxide w/Simethicone)

Delivers: 30 mL

Case = 100 UD Cups (Do Not Break Case)

NDC: 81033-006-01

Product Name: **Kesin MAG-AL™ Plus 1200 mg/1200 mg/120 mg per 30 mL**
(Aluminum & Magnesium Hydroxide w/Simethicone)

Delivers: 30 mL Case = 100 UD Cups (Do Not Break Case)

Storage: 68°F to 77°F (20°C to 25°C)

EXP: 00-OCT-0000 LOT: K0000000 QTY: 1



(17) 001000 (10) K0000000 (30) 1



(01) 00381033006017



Packaged by and Distributed by:
Kesin Pharma Corporation - Oldsmar, FL

Internal Use Only R.01



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KESIN MAG-AL PLUS

aluminum and magnesium hydroxide w/simethicone suspension suspension

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:81033-006
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ALUMINUM HYDROXIDE (UNII: 5QB0T2IUN0) (ALUMINUM HYDROXIDE - UNII:5QB0T2IUN0)	ALUMINUM HYDROXIDE	200 mg in 5 mL
MAGNESIUM HYDROXIDE (UNII: NBZ3QY004S) (MAGNESIUM CATION - UNII:T6V3LHY838, HYDROXIDE ION - UNII:9159UV381P)	MAGNESIUM HYDROXIDE	200 mg in 5 mL
DIMETHICONE, UNSPECIFIED (UNII: 92RU3N3Y1O) (DIMETHICONE, UNSPECIFIED - UNII:92RU3N3Y1O)	DIMETHICONE, UNSPECIFIED	20 mg in 5 mL

Inactive Ingredients

Ingredient Name	Strength
SUCRALOSE (UNII: 96K6UQ3ZD4)	
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
PROPYLPARABEN (UNII: Z8IX2SC1OH)	
GLYCERIN (UNII: PDC6A3C0OX)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
SORBITOL (UNII: 506T60A25R)	
MICROCRYSTALLINE CELLULOSE (UNII: OP1R32D61U)	
METHYLPARABEN (UNII: A2I8C7HI9T)	
MINT (UNII: FV98Z8GITP)	
WATER (UNII: 059QF0KO0R)	
XANTHAN GUM (UNII: TTV12P4NEE)	

Product Characteristics

Color	white (Milky)	Score	
Shape		Size	
Flavor	CINNAMON (Mint)	Imprint Code	
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:81033-006-01	100 in 1 CASE	11/05/2025	
1	NDC:81033-006-30	30 mL in 1 CUP, UNIT-DOSE; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M001	11/05/2025	

Labeler - Kesin Pharma Corporation (117447816)

Establishment

Name	Address	ID/FEI	Business Operations
Kesin Pharma Corporation		117447816	label(81033-006) , pack(81033-006)