PREMIER VALUE INFANTS GAS RELIEF- simethicone emulsion Chain Drug Consortium

Premier Value Infants' Drops Gas Relief Drug Facts

Active ingredient (in each 0.3 mL)

Simethicone 20 mg

Purpose

Antigas

Uses

relieves the discomfort of infant gas frequently caused by air swallowing or certain formulas or foods

Warnings

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
- all dosages may be repeated as needed, after meals and at bedtime or as directed by a physician. Do not exceed 12 doses per day.
- fill enclosed dropper to recommended dosage level and dispense liquid slowly into baby's mouth, toward the inner cheek
- dosage can also be mixed with 1 oz. cool water, infant formula or other suitable liquids
- clean dropper after each use replace bottle with original cap

age (yr)	weight (lb)	dose	
infants under 2	under 24	0.3 mL	
children over 2	over 24	0.6 mL	

Other information

- tamper evident: do not use if printed seal under cap is torn or missing
- store at room temperature
- do not freeze
- see bottom panel for lot and expiration date

Inactive ingredients

carboxymethylcellulose sodium, citric acid, flavors, microcrystalline cellulose, polysorbate 60, potassium sorbate, purified water, sodium benzoate, sorbitan monostearate, sorbitol, xanthan gum

Principal Display Panel

COMPARE TO ACTIVE INGREDIENT IN INFANTS' MYLICON® DROPS*

Premier Value[®]

Infants'

Gas Relief

Non-Staining formula

Simethicone 20 mg/Antigas

- Dye Free
- No Saccharin
- No Artificial Colors
- No Artificial flavors

Syringe Enclosed. This Bottle contains 100 doses (0.3 mL/dose)

1 Fl. OZ (30ml)

100 Doses

Distributed by

Pharmacy Value Alliance, LLC

407 East Lancaster Avenue,

Wayne, PA 19087

*This product is not manufactured or distributed by Infirst Healthcare Inc., the distributor of Infants' Mylicon[®] Drops.



PREMIER VALUE INFANTS GAS RELIEF								
simethicone emulsion								
Product Information								
Product Type	HUMAN OTC DRUG	Item Code	(Source)	NDC:68016-670				
Route of Administration	ORAL							
Active Ingredient/Active Moiety								
Ingred	ient Name		Basis of Streng	th Strength				
DIMETHICONE (UNII: 92RU3N3Y10) (DIMETHICONE - UNII:92RU3N3Y10)			DIMETHICONE	20 mg in 0.3 mL				
Inactive Ingredients								

	Ingredient Name						
CARBOXYMETHYLCELLULOSE SODIUM, UNSPECIFIED (UNII: K679OBS311)							
	DROUS CIT	RIC	ACID (UNII: XF417D3PSL)				
MICROCRYSTALLINE CELLULOSE (UNII: OP1R32D61U)							
POLY	SORBATE 6	0 (U	NII: CAL22UVI4M)				
рот/	ASSIUM SOR	BAT	E (UNII: 1VPU26JZZ4)				
WATER (UNII: 059QF0K00R)							
SODI	UM BENZO	ATE (UNII: OJ245FE5EU)				
SORE	ΒΙΤΑΝ ΜΟΝΟ	OSTE	ARATE (UNII: NVZ4I0H58X)				
SORE	BITOL (UNII:	506T	60A25R)				
KANT	THAN GUM (UNII:	TTV12P4NEE)				
Dro	duct Cha	ract	aristics				
					Score		
Color WHITE (white to off white, opaque)		mile (while to on while, opaque)		Size			
Shaj	•						
Flavor					Imprint Co	ae	
Cont	taine						
Cont	tains						
Pac	tains kaging tem Code		Package Description	ľ	Marketing Start Date	Marketing End Date	
Pac # It	kaging	1 in	Package Description 1 CARTON		-	-	
Pac # It 1 NC	kaging tem Code DC:68016-	30 r			Date	-	
Pac # It 1 ^{NE} 67	kaging tem Code DC:68016-	30 r	1 CARTON nL in 1 BOTTLE, DROPPER; Type 0: Not a		Date	-	
Pac # It 1 ^{NE} 67	kaging tem Code	30 r Con	1 CARTON nL in 1 BOTTLE, DROPPER; Type 0: Not a		Date	-	
Pac # It 1 ^{NE} 67 1	kaging tem Code	30 r Con	1 CARTON nL in 1 BOTTLE, DROPPER; Type 0: Not a nbination Product	0:	Date	Marketing End Date Marketing End Date	

Labeler - Chain Drug	Consortium (101668460)
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Revised: 11/2023

Chain Drug Consortium