SULPHUR - sulfur pellet REMEDY MAKERS

Disclaimer: This homeopathic product has not been evaluated by the Food and Drug Administration for safety or efficacy. FDA is not aware of scientific evidence to support homeopathy as effective.

SULPHUR 6X (Sulfur)

ECZEMA RASH, ACNE, OR OTHER INDICATIONS

SULFUR

WARNING:Keep this and all medications out of reach of children.

INDICATIONS: To be use according to standard homeopathic indications, for self-limiting conditions such as these indicated above or as directed by a physician.

WARNING: Use only if cap and seal are unbroken. If symptoms persist for more than 3 days or worsen, discontinue (STOP) use and consult your physician. As with any drug. If you are pregnant or nursing (breast-feeding) a baby, seek the advise of a health professional before using this product. Store tightly closed in a cool area.

Directions:(adult/children) Dissolve 3 or 4 pellets in mouth or under tongue 3 times a day or as directed by a physician. Children 2 years and older take 1/2 the adult dose.

Inactive Ingredient: Lactose and Sucrose. Free from yeast, wheat, corn and soy.

Questions or comments.(877)REM4YOU. Fax (909)594-4205 Pomona, CA. 91768. USA www.remedymakers.com

Other information: Contain approx.137 - 141 pellets.



SULPHUR

(Sulfur)

Approx. 0.1mcg per gram

ECZEMA RASH, ACNE OR OTHER INDICATIONS

2 Drams (1/4 ounce) Lot # JBB-AEB-C

Exp.10/2021

Drug Facts: Active Ingredient Listed above. To be used according to standard homeopathic indications for self-limiting conditions such as those indicated above or as directed by a physician. Warning: Use only if cap and seal are unbroken. Keep this and all medication out of reach of children. If symptoms persist for more than 3 days or worsen, discontinue (STOP) use and consult your physician. As with any drug, if you are pregnant or nursing (breast-feeding) a baby, seek the advise of a health professional before using this product. Store tightly closed in a cool area.

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HOMEOPATHIC MEDICINE

SULPHUR

sulfur pellet

Product Information

Product Type

HUMAN OTC DRUG

Item Code (Source)

NDC:10191-1578

Route of Administration

SUBLINGUAL

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
SULFUR (UNII: 70 FD1KFU70) (SULFUR - UNII:70 FD1KFU70)	SULFUR	6 [hp_X]		

Inactive Ingredients			
Ingredient Name	Strength		
SUCROSE (UNII: C151H8M554)			
LACTOSE (UNII: J2B2A4N98G)			

Packaging	ackaging					
# Item Code	Package Description	Marketing Start Date	Marketing End Date			
1 NDC:10191-1578-2	137 in 1 VIAL, GLASS					

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
unapproved homeopathic		12/08/1998			

Labeler - REMEDY MAKERS (018543582)

Revised: 11/2011 REMEDY MAKERS