ANILINUM- anilinum pellet HOMEOLAB USA INC.

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.

HOMEOPATHIC MEDICINE NDC 60512-6180-1

ACTIVE INGREDIENT HPUS

ANILINUM 4X

(Aniline)

ECZEMA OF THE KNEES

USE

For self-limiting condition listed above or as directed by a health professional.

WARNINGS

Do not use if pellet-dispenser seal is broken.

Stop use and ask a doctor if symptoms persist more than 3 days or worsen.

If pregnant or breast-feeding, ask a health professional before use.

Keep out of reach of children.

DIRECTIONS

Adults: Allow 3 or 4 pellets to dissolve in the mouth 3 times a day until symptoms are relieved or as directed by a health professional.

OTHER INFORMATION

Store at room temperature.

INACTIVE INGREDIENTS

Lactose, sucrose.

QUESTIONS?

1-800-404-4666

The letters 'HPUS' indicate that the component in this product is officially monographed in the Homeopathic Pharmacopoeia of the United States.

These claims have not been reviewed by the Food and Drug Administration. They are based on traditional homeopathic practice.

80 Pellets

Pellet dispenser

Mfd for: HOMEOLAB USA INC., 3025 De L'Assomption, Montreal, QC, H1N 2H2, CANADA

Product of Canada

LABEL

HOMEOPATHIC MEDICINE

ANILINUM

Aniline NDC 60512-6180-1

ECZEMA OF THE KNEES *

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ANILINUM

anilinum pellet

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HUMAN OTC DRUG NDC:60512-6180 Product Type Item Code (Source)

ORAL **Route of Administration**

Active Ingredient/Active Moiety					
Ingredient Name	Basis of Strength	Strength			
ANILINE (UNII: SIR7XX2F1K) (ANILINE - UNII:SIR7XX2F1K)	ANILINE	4 [hp_X]			

Inactive Ingredients					
Ingredient Name	Strength				
LACTOSE (UNII: J2B2A4N98G)					
SUCROSE (UNII: C151H8 M554)					

Packaging						
# Item Code	Package Description	Marketing Start Date	Marketing End Date			
1 NDC:60512-6180-1	80 in 1 TUBE					

Marketing Information						
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date			
unapproved homeopathic		10/11/1995				

Labeler - HOMEOLAB USA INC. (202032533)

Registrant - HOMEOLAB USA INC. (202032533)

Establishment						
Name	Address	ID/FEI	Business Operations			
HOMEOLAB USA INC.		202032533	manufacture(60512-6180)			

Revised: 10/2013 HOMEOLAB USA INC.