CLEAR PROOF CLARIFYING CLEANSING GEL ACNE MEDICATION- salicylic acid gel Mary Kay Inc.

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

Clear Proof Clarifying Cleansing Gel Drug Facts

Active ingredient

Salicylic Acid (2% W/W)

Purpose

Acne Medication

Uses

- for the management of acne
- helps prevent new acne pimples

Warnings

For external use only

When using this product

- skin irritation and dryness is more likely to occur if you use another topical acne medication at the same time. If irritation occurs, only use one topical acne medication at a time.
- avoid contact with the eyes

Stop use and ask a doctor if

irritation or sensitivity develops or increases

Keep out of reach of children.

If swallowed, get medical help or contact a Poison Control Center right away.

Directions

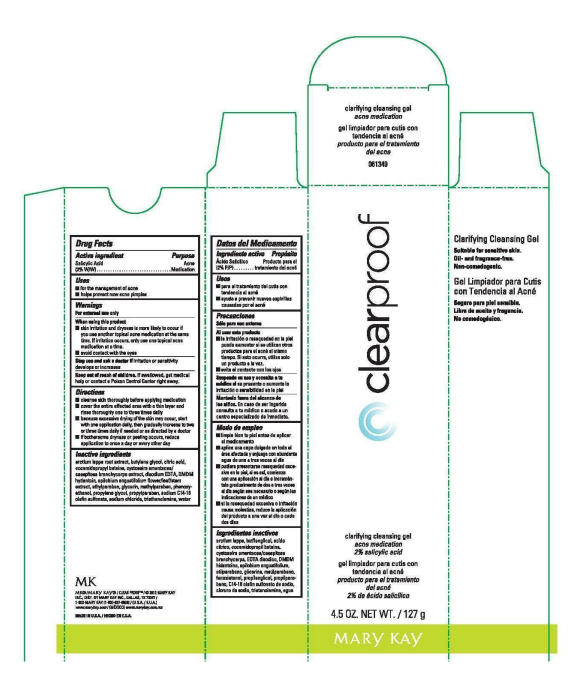
- cleanse skin thoroughly before applying medication
- cover the entire affected area with a thin layer and rinse thoroughly one to three times daily
- because excessive drying of the skin may occur, start with one application daily, then gradually increase to two or three times daily if needed or as directed by a doctor
- if bothersome dryness or peeling occurs, reduce application to once a day or every other day

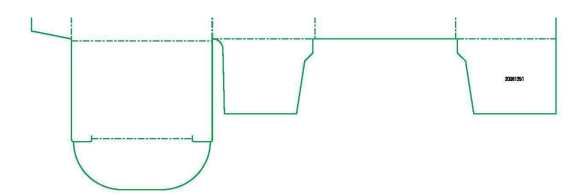
Inactive ingredients

arctium lappa root extract, butylene glycol, citric acid, cocamidopropyl betaine, cystoseira amentacea/caespitosa branchycarpa extract, disodium EDTA, DMDM hydantoin, epilobium angustifolium flower/leaf/stem extract, ethylparaben, glycerin, methylparaben, phenoxyethanol, propylene glycol, propylparaben, sodium C14-16 olefin sulfonate, sodium chloride, triethanolamine, water

Principal Display Panel - 127 g carton

clearproof clarifying cleansing gel acne medication 2% salicylic acid 4.5 OZ. NET WT. / 127 g Mary Kay





CLEAR PROOF CLARIFYING CLEANSING GEL ACNE MEDICATION

salicylic acid gel

Product Information				
coduct TypeHUMAN OTC DRUGItem Code (Source)NDC:52				
Route of Administration TOPICAL				
Active Ingredient/Active Moi	etv			
Ingredient Name Basis of Strength				Strength
SALICYLIC ACID (UNII: O414PZ4LPZ)	(SALICYLIC ACID - UNII:0414	PZ4LPZ)	SALICYLIC ACID	2 g in 100 g
Inactive Ingredients	Ingredient Name			Strength
Inactive Ingredients				
	Ingredient Name			Strength
WATER (UNII: 059QF0KO0R)	UNIL NO 1C FROM			
SODIUM C14 OLEFIN SULFONATE (COCAMIDOPROPYL BETAINE (UNII:	,			
SODIUM CHLORIDE (UNII: 451W47IQ				
TROLAMINE (UNII: 903K93S3TK)				
	86K)			
EDETATE DISODIUM (UNII: 7FLD91C86K)				
PHENOXYETHANOL (UNII: HIE492ZZ	3T)			
PHENOXYETHANOL (UNII: HIE492ZZ BUTYLENE GLYCOL (UNII: 3XUS85K	3T) (0RA)			
PHENOXYETHANOL (UNII: HIE492ZZ	3T) (0RA) 2968PHW8QP)			
PHENOXYETHANOL (UNII: HIE492ZZ BUTYLENE GLYCOL (UNII: 3XUS85K CITRIC ACID MONOHYDRATE (UNII: PROPYLENE GLYCOL (UNII: 6DC9Q	3T) KORA) 2968PHW8QP) 167V3)			
PHENOXYETHANOL (UNII: HIE492ZZ BUTYLENE GLYCOL (UNII: 3XUS855 CITRIC ACID MONOHYDRATE (UNII:	3T) KORA) 2968PHW8QP) 167V3)			
PHENOXYETHANOL (UNII: HIE492ZZ BUTYLENE GLYCOL (UNII: 3XUS855 CITRIC ACID MONOHYDRATE (UNII: PROPYLENE GLYCOL (UNII: 6DC9Q METHYLPARABEN (UNII: A218C7H19T	3T) (ORA) 2968PHW8QP) 167V3) ()			
PHENOXYETHANOL (UNII: HIE492ZZ BUTYLENE GLYCOL (UNII: 3XUS85F CITRIC ACID MONOHYDRATE (UNII: PROPYLENE GLYCOL (UNII: 6DC9Q METHYLPARABEN (UNII: A218C7H19T ETHYLPARABEN (UNII: 14255EXE39) DMDM HYDANTOIN (UNII: BYR0546T	3T) (ORA) 2968PHW8QP) 167V3) (OW)			
PHENOXYETHANOL (UNII: HIE492ZZ BUTYLENE GLYCOL (UNII: 3XUS85F CITRIC ACID MONOHYDRATE (UNII: PROPYLENE GLYCOL (UNII: 6DC9Q METHYLPARABEN (UNII: A218C7H19T ETHYLPARABEN (UNII: 14255EXE39)	3T) 3T) 3T) 3T) 3T) 3T) 3T) 3T)			

Packaging						
#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
1 N	DC:51531-1349-4	1 in 1 CARTON	08/15/2013			
1						
L		127 g in 1 TUBE; Type 0: Not a Combination Product				
1		127 g in 1 TUBE; Type 0: Not a Combination Product				
•		127 g in TTUBE; Type 0: Not a Combination Product				
1						
Ma	arketing Inf					
	arketing Inf	ormation	Marketing Start Date	Marketing End Date		

Labeler - Mary Kay Inc. (049994452)

Establishment				
Name	Address	ID/FEI	Business Operations	
Mary Kay Inc.		103978839	manufacture(51531-1349)	

Establishment

Name	Address	ID/FEI	Business Operations
Mary Kay Inc.		081158134	manufacture(51531-1349)

Establishment					
Name	Address	ID/FEI	Business Operations		
Englewood Lab Inc.		080987545	pack(51531-1349)		

Revised: 1/2019

Mary Kay Inc.