GOLD COSMETICS BLEACH CREAM FORTE- hydroquinone cream cream Peer Pharm Ltd

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.

Bleach Cream Forte

Hydroquinone 8%

Face cream for gradual fading of dark spots

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Apply a very thin layer once a day, only at night, all over the face

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Sun exposure should be limit by using a sunscreen agent, a sun blocking agent, or protective clothing to cover bleached skin

when using and after using this product in order to prevent darkening from reoccurring.

Proper use of skin bleaching creams is extremely important, and directions should be followed carefully at all times.

Large skin areas, such as the entire face, should not be bleached at once due to the possibility of skin discoloration.

Product should be used on the darkened area of skin only.

For external use only.

Avoid contact with eyes. If contact occurs, rinse eyes thoroughly with water.

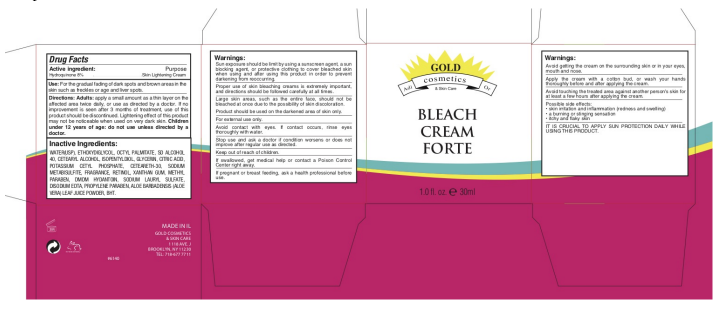
Stop use and ask a doctor if condition worsens or does not improve after regular use as directed.

Keep out of reach of children.

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

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

Keep out of reach of children



GOLD COSMETICS BLEACH CREAM FORTE

hydroquinone cream cream

Droduct	Information
Product	Imormation

Product Type HUMAN OTC DRUG Item Code (Source) NDC:69435-1901

Route of Administration TOPICAL

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
HYDRO Q UINO NE (UNII: XV74C1N1AE) (HYDRO Q UINO NE - UNII: XV74C1N1AE)	HYDROQUINONE	8 mg in 100 mL	

Inactive Ingredients		
Ingredient Name	Strength	
GLYCERIN (UNII: PDC6A3C0OX)		
JOJOBA OIL (UNII: 724GKU717M)		
DECYL OLEATE (UNII: ZGR06DO97T)		
SODIUM DITHIONATE (UNII: RPF7Z41GAW)		
XANTHAN GUM (UNII: TTV12P4NEE)		
PHENOXYETHANOL (UNII: HIE492ZZ3T)		
CITRIC ACID MONO HYDRATE (UNII: 2968 PHW8 QP)		
POTASSIUM SORBATE (UNII: 1VPU26JZZ4)		
ALOE VERA LEAF (UNII: ZY8 1Z8 3H0 X)		
BUTYLATED HYDRO XYTO LUENE (UNII: 1P9 D0 Z171K)		
WATER (UNII: 059QF0KO0R)		
CETOSTEARYL ALCOHOL (UNII: 2DMT128M1S)		
STEARYL ALCOHOL (UNII: 2KR8914H1Y)		
CETEARETH-30 (UNII: 1R9 DCZ5FOX)		
STEARETH-21 (UNII: 53J3F32P58)		
STEARETH-2 (UNII: V56DFE46J5)		
EDETATE DISODIUM ANHYDROUS (UNII: 8 NLQ36 F6 MM)		
CHLORPHENESIN (UNII: I670 DAL4SZ)		
CETYL ALCOHOL (UNII: 936JST6JCN)		
ALLANTO IN (UNII: 344S277G0Z)		
ALPHA-TOCOPHEROL ACETATE (UNII: 9E8X80D2L0)		

ı	Packaging			
ı	# Item Code Package Description		Marketing Start Date	Marketing End Date
ı	1 NDC:69435-1901-1	30 mL in 1 TUBE; Type 0: Not a Combination Product	09/05/2019	

Marketing Infor	mation		
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part358 A	09/05/2019	

Labeler - Peer Pharm Ltd (514678390)

Registrant - Peer Pharm Ltd (514678390)

Establishment			
Name	Address	ID/FEI	Business Operations
Peer Pharm Ltd		514678390	manufacture(69435-1901), label(69435-1901)

Revised: 9/2019 Peer Pharm Ltd