I-MAX LIGHTENING 5- hydroquinone cream MAXLIFE USA, 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.

ACTIVE INGREDIENTS: HYDROQUINONE USP 2%

PURPOSE:

SKIN LIGHTENING

USES:

FOR THE GRADUAL FADING OF DARK AREAS OF THE SKIN.

WARNINGS:

AVOID CONTACT WITH EYES. SOME USERS MAY EXPERIENCE MILD SKIN IRRITATION.

IF IRRITATION BECOMES SEVERE, STOP USE AND CONSULT A DOCTOR.

THIS PRODUCT IS NOT INTENDED FOR USE IN THE PREVENTION OF SUNBURN AND CONTAINS AN ALPHA HYDROXY ACID (AHA) THAT MAY INCREASE YOUR SKIN'S SENSITIVITY TO THE SUN AND PARTICULARLY THE POSSIBILITY OF SUNBURN. SUN EXPOSURE SHOULD BE LIMITED BY USING A SUNSCREEN AGENT OR PROTECTIVE CLOTHING TO COVER BLEACHED SKIN AFTER TREATMENT IS COMPLETED TO PREVENT DARKENING FROM REOCCURING.

DO NOT USE ON CHILDREN UNDER 12 YEARS OF AGE UNLESS DIRECTED BY A DOCTOR.

DIRECTIONS:

ADULTS: APPLY A SMALL AMOUNT AS A THIN LAYER ON THE AFFECTED AREA TWICE DAILY, OR USE AS DIRECTED BY A DOCTOR. IF NO IMPROVEMENT IS SEEN AFTER 3 MONTHS OF TREATMENT, USE OF THIS PRODUCT SHOULD BE DISCONTINUED. LIGHTENING EFFECT OF THIS PRODUCT MAY NOT BE NOTICEABLE WHEN USED ON VERY DARK SKIN.

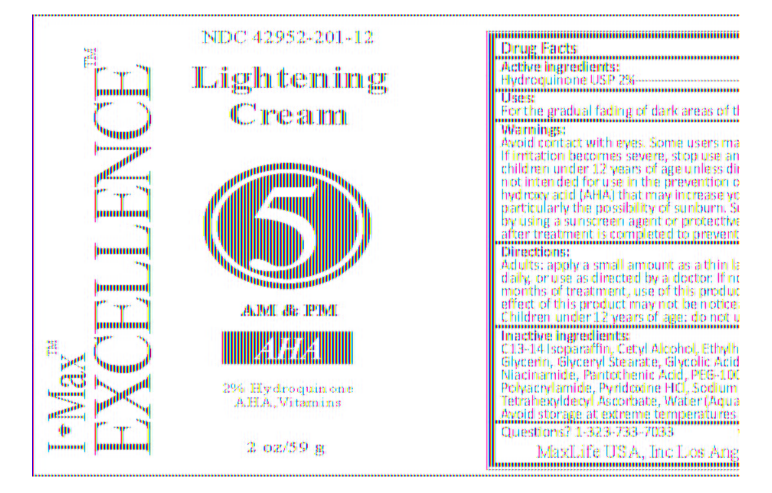
CHILDREN UNDER 12 YEARS OF AGE: DO NOT USE UNLESS DIRECTED BY A DOCTOR.

INACTIVE INGREDIENTS:

C13-14 ISOPARAFFIN, CETYL ALCOHOL, ETHYLHEXYLGLYCERIN, ETHYLHEXYL STEARATE, GLYCERIN, GLYCERYL STEARATE, GLYCOLIC ACID, ISOPROPYL MYRISTATE, LAURETH-7, NIACINAMIDE, PANTOTHENIC ACID, PEG-100 STEARATE, PHENOXYETHANOL, POLYACRYLAMIDE, PYRIDOXINE HCL, SODIUM HYDROXIDE, SODIUM METABISULFITE, TETRAHEXYLDECYL ASCORBATE, WATER (AQUA), XANTHAN GUM.

QUESTIONS? 1-323-733-7033

KEEP OUT OF REACH OF CHILDREN.



I-MAX LIGHTENING 5 hydroguinone cream **Product Information Product** Type HUMAN OTC DRUG Item Code (Source) NDC:42952-201 **Route of Administration** TOPICAL **Active Ingredient/Active Moiety Ingredient Name Basis of Strength** Strength HYDROQUINONE (UNII: XV74C1N1AE) (HYDROQUINONE - UNII:XV74C1N1AE) HYDROQUINONE 2 g in 100 g **Inactive Ingredients Ingredient Name** Strength C13-14 ISOPARAFFIN (UNII: E4F12ROE70) CETYL ALCOHOL (UNII: 936JST6JCN) ETHYLHEXYLGLYCERIN (UNII: 147D247K3P) ETHYLHEXYL STEARATE (UNII: EG3PA2K3K5) GLYCERIN (UNII: PDC6A3C0OX) GLYCERYL MONOSTEARATE (UNII: 2300U9XXE4) GLYCOLIC ACID (UNII: 0WT12SX38S) ISOPROPYL MYRISTATE (UNII: 0 RE8 K4LNJS)

LAURETH-7 (UNII: Z95S6G8201)						
NIACINAMIDE (UNII: 25X51I8RD4)						
PANTOTHENIC ACID (UNII: 19F5HK2737)						
PEG-100 STEARATE (UNII: YD01N1999R)						
P	PHENOXYETHANOL (UNII: HIE492ZZ3T)					
POLYACRYLAMIDE (10000 MW) (UNII: E2KR9C9V2I)						
PYRIDO XINE HYDRO CHLO RIDE (UNII: 68 Y4CF58 BV)						
SO DIUM HYDRO XIDE (UNII: 55X04QC32I)						
S	O DIUM METABISUL	FITE (UNII: 4VON5FNS3C)				
Т	ETRAHEXYLDECYL	ASCORBATE (UNII: 9LBV3F07AZ)				
W	WATER (UNII: 059QF0KO0R)					
X	XANTHAN GUM (UNII: TTV12P4NEE)					
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P	ackaging					
P #	ackaging Item Code	Package Description	N	Aarketing Start Date	Marketing End Date	
#	0 0	Package Description 59 g in 1 TUBE; Type 0: Not a Combination Prod		Marketing Start Date 9/18/2012	Marketing End Date	
#	Item Code	• •		0	Marketing End Date	
#	Item Code	• •		0	Marketing End Date	
#	Item Code	59 g in 1 TUBE; Type 0: Not a Combination Prod		0	Marketing End Date	
# 1	Item Code NDC:42952-201-12	59 g in 1 TUBE; Type 0: Not a Combination Prod	luct 09	0	Marketing End Date Marketing End Date	
# 1	Item Code NDC:42952-201-12	59 g in 1 TUBE; Type 0: Not a Combination Prod Ormation y Application Number or Monograph C	luct 09	9/18/2012		

Labeler - MAXLIFE USA, INC. (785111431)

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MAXLIFE USA, INC.