CONDITION AND ENHANCE SYSTEM TRAVEL-SIZE NON-SURGICAL- hydroquinone, octinoxate, and zinc oxide OMP, INC.

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

Obagi[®] Condition & Enhance Clear (Hydroquinone USP, 4%) Skin Bleaching Cream

Obagi[®] Condition & Enhance Blender[®] (Hydroquinone USP, 4%) Skin Bleaching Cream

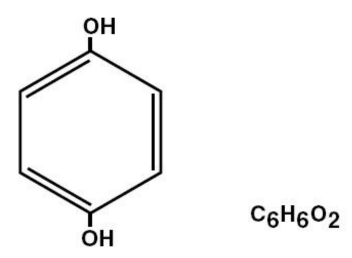
Rx Only

FOR EXTERNAL USE ONLY

DESCRIPTION

Hydroquinone is 1,4-benzenediol. Hydroquinone occurs as fine, white needles. The drug is freely soluble in water and in alcohol. Chemically, hydroquinone is designated as p-dihydroxybenzene; the empirical formula is C_6 H_6 O_2 ; molecular weight is 110.0.

Obagi[®] **Condition & Enhance Blender** contains Hydroquinone USP 40 mg/gm in a base of purified water, glycerin, cetyl alcohol, PPG-2 myristyl ether propionate, sodium lauryl sulfate, TEA-salicylate, lactic acid, phenyl trimethicone, tocopheryl acetate, sodium metabisulfite, ascorbic acid, methylparaben, saponins, disodium EDTA, BHT, and propylparaben.



Obagi[®] **Condition & Enhance Clear** contains Hydroquinone USP 40 mg/gm in a base of purified water, cetyl alcohol, glycerin, sodium lauryl sulfate, stearyl alcohol, tocopheryl acetate, ascorbic acid, sodium metabisulfite, lactic acid, saponins, disodium EDTA, methylparaben, BHT, propylparaben, and butylparaben.

CLINICAL PHARMACOLOGY

Topical application of hydroquinone produces a reversible depigmentation of the skin by inhibition of the enzymatic oxidation of tyrosine to 3, 4-dihydroxyphenylalanine (dopa) and suppression of other melanocyte metabolic processes.

Exposure to sunlight or ultraviolet light will cause repigmentation of the bleached areas, which may be prevented by the use of sunblocking agents or sunscreen agents contained in Obagi Condition & Enhance.

INDICATIONS AND USAGE

The gradual bleaching of hyperpigmented skin conditions such as chloasma, melasma, freckles, senile lentigines, and other unwanted areas of melanin hyperpigmentation.

CONTRAINDICATIONS

Prior history of sensitivity or allergic reaction to this product or any of its ingredients. The safety of topical hydroquinone use during pregnancy or in children (12 years and under) has not been established.

WARNINGS

Caution

Hydroquinone is a skin bleaching agent which may produce unwanted cosmetic effects if not used as directed. The physician should be familiar with the contents of this insert before prescribing or dispensing this medication.

Test for skin sensitivity before using by applying a small amount to an unbroken patch of skin and check in 24 hours. Minor redness is not a contraindication, but where there is itching or vesicle formation or excessive inflammatory response, further treatment is not advised. Close patient supervision is recommended.

Avoid contact with eyes. In case of accidental contact, patient should rinse eyes thoroughly with water and contact physician. A bitter taste and anesthetic effect may occur if applied to lips.

Sunscreen use is an essential aspect of hydroquinone therapy because even minimal sunlight exposure sustains melanocytic activity.

Warning

Contains sodium metabisulfite, a sulfite that may cause serious allergic type reactions (e.g., hives, itching, wheezing, anaphylaxis, severe asthma attacks) in certain susceptible persons.

PRECAUTIONS

(SEE WARNINGS)

General

Treatment should be limited to relatively small areas of the body at one time since some patients experience a transient skin reddening and a mild burning sensation which does not preclude treatment.

Pregnancy Category C

Animal reproduction studies have not been conducted with topical hydroquinone. It is also not known whether hydroquinone can cause fetal harm when used topically on a pregnant woman or affect reproductive capacity. It is not known to what degree, if any, topical hydroquinone is absorbed systemically. Topical hydroquinone should be used on pregnant women only when clearly indicated.

Nursing mothers

It is not known whether topical hydroquinone is absorbed or excreted in human milk. Caution is advised when topical hydroquinone is used by a nursing mother.

Pediatric usage

Safety and effectiveness in children below the age of 12 years have not been established.

ADVERSE REACTIONS

No systemic adverse reactions have been reported. Occasional hypersensitivity (localized contact dermatitis) may occur, in which case the medication should be discontinued and the physician notified immediately.

DOSAGE AND ADMINISTRATION

A thin application should be applied to the affected area twice daily or as directed by a physician. If no improvement is seen after three (3) months of treatment, use of this product should be discontinued. Sun exposure should be limited 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 repigmentation.

HOW SUPPLIED

Obagi Condition and Enhance Blender is available as follows:

2 oz. (57 gm) bottle NDC 62032-115-36 1 oz. (28.5 gm) bottle NDC 62032-115-10

Obagi Condition and Enhance Clear is available as follows:

2 oz. (57 gm) bottle NDC 62032-117-36

Store at 25°C (77°F); excursion permitted to 15°C-30°C (59°F-86°F).

OBAGI® MEDICAL

OMP, Inc. Long Beach, CA 90802 USA 1-800-636-7546 80707910U Rev. 6/07

PRINCIPAL DISPLAY PANEL - Kit Carton

OBAGI® **CONDITION & ENHANCE SYSTEM**For Use with Nonsurgical Procedures

Travel Size





OBAGI" CONDITION & ENHANCE SYSTEM

OBAGI

CONDITION & ENHANCE SYSTEM

For Use with Nonsurgical Procedures

Skincare and protection should be a fundamental part of good height and beautiful, vibrant skin, Your skin needs special care to stay young and health; The Obagit* Condition & Enhance System is the leading prescription skincare program that works at the cellular level for skin that looks and acts younger and healther. Whether you're planning a surpical or nonsurpical procedure, the Obegi Condition & Enhance System uses proven ingredients and penetrating technologies to help ensure a successful outcome.

Foaming Gel (Cleanser)

Foaming Gel (Cleaver)

Coapi Forning Fel is a sup-five downer that removes inputities, oil and mide-sq. This appeal formal prepares the skin for the transformation process, it is designed for use with non-surgical procedures.

DRECTIONES Apple to demo face and need with moistered fingerings in the morning and events, Prices completely with warm water.

INEXCELLIBRATE Virties water, such insured act serious acids, occanidoropyl betters, social haureth suffice also bestedensis betty use, medicago safes (dafe)a district, belongs offerede serious charments in colds (in fundicisal) extract, acid un chief de, serious quantities, proceptions phonosynthesis, entry burstens, respiration, propiparation, isobulybamben, fragmence, D&C Red No. 33, FD&C Vellow No. 5.

Toner (Skin Preparation)

Obagi Toner, formulated with alum (a natural astringent), adjusts the pH of the skin for optimal penetration of the treatment ingredients in the System. DIRECTIONS: Apply after cleaning in the morning and evening. Saturate cotton pad and wipe gently over face and neck. Avaid eyeld area. Do not nice off.

WARRINGS: Avoid contact with eyes, in case of accidental contact, patient should rinse eyes thoroughly with water and contact a physician. A bitter state and assentate is effect may occur if applied to lips. Sunscreen use is an essential aspect of hydroquimone therapy because even minimal surlight exposure sustains melamocytic activity.

melanosytic activity.

CAUTION: Contains sodium metabisuffite, a suffite that may cause serious allergic-type reactions (e.g., hives, ltching, wheezing, anaphilaxis, severe authma attacks) in certain succeptible persons.

PIGREGEMENT Sylvocytione USP 40 migram is base of purified water only alsohal glycom, sodium juryl suffits, steary latched, temphreyl acceler, social seconds oid, sodium results office, but one day approving discolum EDITA, metrylparaben, BHT, propylparaben and burylparaben. See enclosed Package Insert for full prescription information,

Rx ONLY, FOR EXTERNAL USE ONLY,

Exfoderm® Forte (Exidation Enhancer)

Obagi Estaderm Forte estallates dead surface skin cells and smoothes mughness, aiding in penetration of treatment ingredients in the System. **DIRECTIONS:** Apply to the face in the marring, following the application of Obegi Clear, as directed by a physician. Follow with the appropriate Obegi sur protection.

san processor.

CAUTION: FOR EXTERNAL USE ONLY, Avoid contact with the eyes,
A mild burning sensation of the skin is to be expected. If burning
is severe, discontinue use and consult a physicial,
invariant is severe, discontinue use and consult a physicial,
invariant is severe, discontinue use and consult a physicial,
invariant is severe, discontinue use and consult in the consult in the consultation of the consultation of

CAUTION: Contains sodium metabisulfite, a sulfite that may c serious allergic-type reactions (e.g. hives, tiching, wheezing, anaphykaxis, severe asthma attacks) in certain susceptible persons.

INGREDIENTS: Hydroculone USP 40 mg/ym in a brae of purified water, gloenin, only alsohid, PPG-5 mg/trajt effore propionate, sodium ibuny's adition. ITA-assipaths before add, pheny inmeditioner, toogheyst sectate, acdrum metabolishte, asonoble and, methylpaniben, saponins, disedium EDTA, IB-17 and proxylpaniben.

Rx ONLY, FOR EXTERNAL USE ONLY,

Healthy Skin Protection SPF 35

Obeg Healthy Skin Protection contains 9% microrized zinc oxide to protect the newer healther skin created by skin transformation. This high concentration of microrized zinc oxide provides protection against long UVA mys linked to deep permittive aging.

DRECTIONS: Apply liberally to all exposed areas. Apply at least 15 minutes before sur-exposure and reapply frequently after prolonged swimming, excessive perspiration, vigorous activity or toweling.

DRUG FACTS

Warnings: us for external use only.

Store at 25°C (77°F); excursion permitted to 15°-30°C (59°-86°F) Dist, by OMP, Inc., Long Beach, CA 90806 Made in U.S.A. 40707910U Obagi is a registered trademark of OMP, Inc.

Obagi products are physician dispensed and should be used under the guidance of your skincare specialist.



productigentale numero water, apre narrosportal pria puer, narranga virginiana (policia) haberel distillare condassium durin, sodium PCA, partherno DMOM hydamotin, policionale ed to allementi, antiva officiarilas fasqe), had estrati, borago officiarilas excast, calendular officiarilas flower extract, sapo diodepropynyl bulgkarichamata, fasquarose, FDAC dillar No. 1.

Clear (Skin Bleaching and Corrector Cream) NDC 62032-117-36 Obagi Clear corrects uneven skin color and brown spots, continuing the

INDICATIONS: For the gradual bleaching of hyperoigniented skin condition such as disposed, melasma, freckles, sente tendigines and other unwanted areas of melanin hypergigmentation,

areas in region hyperpoinnesses, DOSANCE AND ADMINISTRATION. A thin layer should be applied to the affected are in the recent good exercise, or as deviced by a physicism. First in proceedings a sean after these mentules of transfers, used this product should be discontinued. On exposure should be firsted by using a sursorom agent, a sun blooking agent, or protective durbing to over blooched sin chaing and after usage of this product in order to prevent ineignmentation.

Patient Instructions

■ Blender® (Skin Lightener and Blending Cream) NDC 62032-115-18 Obegi Render maximizes skin transformation when prescribed with tretinoin cream. Ellender evens skin color and tone, making skin lack and act younger

INDECATIONS: For the gradual bleaching of hyperpigmented skin condition such as chlosama, melasma, freddles, senile lientigines and other unwanted stope of preferring hyperpigmentation.

such as cirkaaria, melsions, feed-this, perils bringines and other convoled sense of meliar in laper granuments. Dossetta, AND ANNINISTRATION: A this flayer should be supplied to the affected are in the menting and venturing or as directed by a physician. Eno improvement is seen start three monits of treatment, use of this product should be obtained using a survivant agent a sun blooking agent or protected obthings to over blookhold with discriptions of the product obtained to the control of the product should be obtained to the product of the product of

where using uses prospect, group as a eyes, was a war war as an Stop use and ask a physician if a Rash or intation develops and persi Keep out of the mech of children, is it swelfowed, pet medical help or contact a Poison Cantrel Center inventising. octions:
Mappy prevently and everly 15 minutes before our exposure children under als months of age; Asis a physician,
Peopply as readed o covel chying, owinning or perspiring.

Directions: as
For other order at mattra

Set used type, wherehing a property

Other Information. Best or coronder over term

Set with a set of the se

Inactive Ingredients: Burgaraten, celeryl deatel, o issperaffin, defraredomine cred phosphate, deadum reletat, isotrophorism, isotropyl poliniate, bureth-7 methylamisen, phonosynthanol polyenylamish, polyenheral, polyenhera 93, polyenhera 93, polyenhera 94, polyenhera 95, polyenhera 95,



1 FL OZ (30 mL)

25£32 Protection Healthy Skin

Rx Only gunone USP, 494 NET WT, 1 OZ. (28.5 g) Blender

NET WT. 1 OZ. (28.5 g) Extodemn Forte

Kx Only 490 anoniup NEL ML 2 OZ (2) 8) Clear

2 FL. OZ. (60 mL) Toner 2 FL OZ (60 mL) Foaming Gel

This Obagia Condition & Enhance System includes:

Note: Evening routine does not include steps 4 or 6.

For Use with Monsurgical Procedures

CONDITION & ENHANCE SYSTEM

OBVCI.



MEDICAL

OBAGI"

CONDITION & ENHANCE SYSTEM

For Use with Nonsurgical Procedures

What to Expect

The desire for healthy, youthful and attractive skin is why you are undergoing facial aesthetic procedures. The Obag® Condition & Enhance System uses proven ingredients and penetrating technologies to help ensure a successful outcome, This system has been shown to improve the overall results of facial aesthetic procedures.

The length of time that it takes to condition your skin varies. You should begin to notice visible improvements to your skin within a six week period.

It is important to understand that you will experience some reactions during your skin conditioning process. These reactions are normal and part of the transformation process. You may experience one or more of

- Dryness
- Redness
- Skin texture and appearance of wrinkles may temporar
- Exfoliation
- · Sensitive skin
- Acne may temporarily worsen

In order to maximize the results of your procedure, it is very import that you follow your personal program as indicated by a skincare professional.

Your physician should discuss with you any reactions related to your facial procedure, as these vary by the type of procedure you have received.

Daily Product Routines

p.m. am Morning Evening QUANTITY QUANTITY 1 Foaming Gel Entire face 1 Foaming Gel Entire face 2 2 Toner Apply liberally with Toner Apply liberally with entire face 3 3 Clear ☐ ½ gm ☐ 1 gm Clear ☐ ½ gm □ 1/2 gm

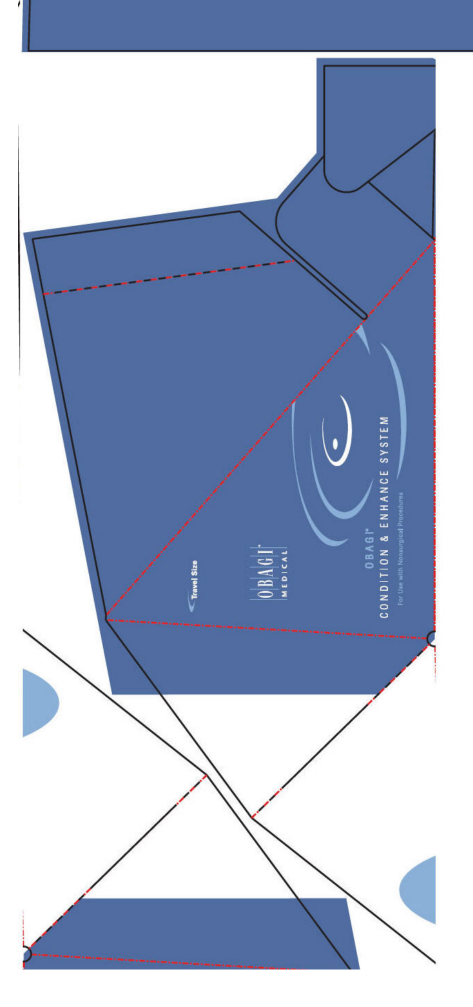
Note: Morning routine does not include step 5,

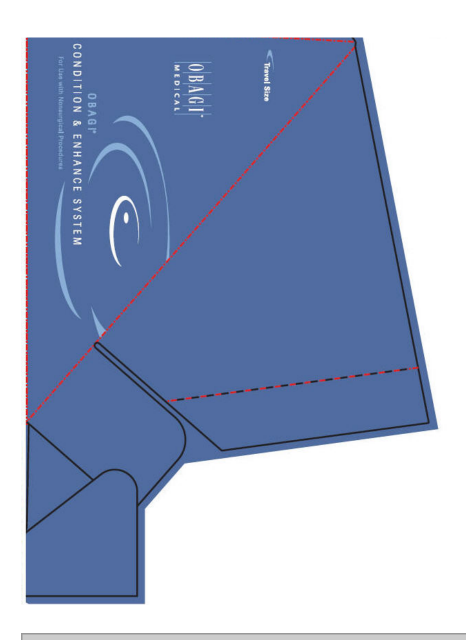
Squeeze the product onto your finger, using the bars below as reference:



Or use the distance from the tip of your pinky finger to the first joint as an approximate measurement of 1/2 gram of product.

OBAGI MEDICAL





CONDITION AND ENHANCE SYSTEM TRAVEL-SIZE NON-SURGICAL

hydroquinone, octinoxate, and zinc oxide kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:62032-510

Packaging

ı	# Item Code	Package Description	Marketing Start Date	Marketing End Date
ı	1 NDC:62032-510-60	1 in 1 CARTON		

Quantity of Parts

Quantity of Lares		
Part #	Package Quantity	Total Product Quantity
Part 1	1 BOTTLE, PLASTIC	28.5 g
Part 2	1 BOTTLE, PLASTIC	57 g
Part 3	1 BOTTLE, PLASTIC	30 mL
Part 4	1 BOTTLE, PLASTIC	60 mL

Part 5	1 BOTTLE, PLASTIC	60 mL
Part 6	1 BOTTLE, PLASTIC	28.5 g

Part 1 of 6

CONDITION AND ENHANCE BLENDER SKIN LIGHTENER AND BLENDING

hydroquinone cream

Product Information		
Item Code (Source)	NDC:62032-115	
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	40 mg in 1 g	

Inactive Ingredients		
Ingredient Name	Strength	
EDETATE DISO DIUM (UNII: 7FLD91C86K)		
PPG-2 MYRISTYL ETHER PROPIONATE (UNII: 88R97D8U8A)		
TROLAMINE SALICYLATE (UNII: H8 O40 40 BHD)		
SODIUM LAURYL SULFATE (UNII: 368GB5141J)		
CETYL ALCOHOL (UNII: 936JST6JCN)		
GLYCERIN (UNII: PDC6A3C0OX)		
LACTIC ACID (UNII: 33X04XA5AT)		
ALPHA-TOCOPHEROL ACETATE (UNII: 9E8X80D2L0)		
ASCORBIC ACID (UNII: PQ6CK8PD0R)		
SO DIUM METABISULFITE (UNII: 4VON5FNS3C)		
WATER (UNII: 059QF0KO0R)		
METHYLPARABEN (UNII: A2I8C7HI9T)		
PROPYLPARABEN (UNII: Z8IX2SC1OH)		
BUTYLATED HYDRO XYTO LUENE (UNII: 1P9 D0 Z171K)		
PHENYL TRIMETHICO NE (UNII: DR0 K5NOJ4R)		

Product Characteristics			
Color WHITE		Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

Packaging

# Item Code	Package Description	Marketing Start Date	Marketing End Date
1 NDC:62032-115-10	28.5 g in 1 BOTTLE, PLASTIC		

Marketing Information				
Marketing Category Application Number or Monograph Citation		Marketing Start Date	Marketing End Date	
UNAPPROVED DRUG OTHER		0 1/0 1/19 8 8		

Part 2 of 6

CONDITION AND ENHANCE CLEAR SKIN BLEACHING AND CORRECTOR

hydroquinone cream

Product Information	
Item Code (Source)	NDC:62032-117
Route of Administration	TOPICAL

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
HYDROQUINONE (UNII: XV74C1N1AE) (HYDROQUINONE - UNII:XV74C1N1AE)	HYDROQUINONE	40 mg in 1 g

Inactive Ingredients		
Ingredient Name	Strength	
EDETATE DISO DIUM (UNII: 7FLD91C86K)		
BUTYLPARABEN (UNII: 3QPI1U3FV8)		
STEARYL ALCOHOL (UNII: 2KR8914H1Y)		
SODIUM LAURYL SULFATE (UNII: 368GB5141J)		
CETYL ALCOHOL (UNII: 936JST6JCN)		
GLYCERIN (UNII: PDC6A3C0OX)		
LACTIC ACID (UNII: 33X04XA5AT)		
ALPHA-TO COPHEROL ACETATE (UNII: 9E8 X80 D2L0)		
ASCORBIC ACID (UNII: PQ6CK8PD0R)		
SO DIUM METABISULFITE (UNII: 4VON5FNS3C)		
WATER (UNII: 059QF0KO0R)		
METHYLPARABEN (UNII: A218 C7H19 T)		
PROPYLPARABEN (UNII: Z8IX2SC1OH)		
BUTYLATED HYDRO XYTO LUENE (UNII: 1P9 D0 Z171K)		

Product Characteristics		
Color	WHITE	Score
Shape		Size
Flavor		Imprint Code

Contains

l	Packaging			
l	# Item Code	Package Description	Marketing Start Date	Marketing End Date
ı	1 NDC:62032-117-36	57 g in 1 BOTTLE, PLASTIC		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
UNAPPROVED DRUG OTHER		0 1/0 1/19 8 8	

Part 3 of 6

CONDITION AND ENHANCE HEALTHY SKIN PROTECTION SPF 35

octinoxate and zinc oxide cream

Product Information	
Item Code (Source)	NDC:62032-119
Route of Administration	TOPICAL

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
OCTINOXATE (UNII: 4Y5P7MUD51) (OCTINOXATE - UNII:4Y5P7MUD51)	OCTINOXATE	75 mg in 1 mL
ZINC OXIDE (UNII: SOI2LOH54Z) (ZINC OXIDE - UNII:SOI2LOH54Z)	ZINC OXIDE	90 mg in 1 mL

Inactive Ingredients		
Ingredient Name	Strength	
ISOPROPYL PALMITATE (UNII: 8 CRQ2TH63M)		
WATER (UNII: 059QF0KO0R)		
ETHYLHEXYL STEARATE (UNII: EG3PA2K3K5)		
CETOSTEARYL ALCOHOL (UNII: 2DMT128M1S)		
POLYSORBATE 60 (UNII: CAL22UVI4M)		
C13-14 ISOPARAFFIN (UNII: E4F12ROE70)		
METHYLPARABEN (UNII: A2I8C7HI9T)		
LAURETH-7 (UNII: Z95S6G8201)		
PROPYLPARABEN (UNII: Z8IX2SC1OH)		
EDETATE DISO DIUM (UNII: 7FLD9 1C8 6 K)		
BUTYLPARABEN (UNII: 3QPI1U3FV8)		
DIETHANO LAMINE CETYL PHO SPHATE (UNII: 4UG0316 V9S)		
PHENOXYETHANOL (UNII: HIE492ZZ3T)		
ETHYLPARABEN (UNII: 14255EXE39)		
ISOBUTYLPARABEN (UNII: 0 QQJ25X58G)		

SO DIUM HYDRO XIDE (UNII: 55X04QC32I)	
TRIETHO XYCAPRYLYLSILANE (UNII: LDC331P08E)	
CITRIC ACID MO NO HYDRATE (UNII: 2968 PHW8 QP)	

Product Characteristics		
Color	WHITE	Score
Shape		Size
Flavor		Imprint Code
Contains		

Packaging			
# Item Code	Package Description	Marketing Start Date	Marketing End Date
1 NDC:62032-119-10	30 mL in 1 BOTTLE, PLASTIC		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC MONOGRAPH FINAL	part352	0 1/0 1/20 0 2	

Part 4 of 6

CONDITION AND ENHANCE FOAMING

cleansing (cold creams, cleansing lotions, liquids, and pads) gel

Product Information Route of Administration TOPICAL

Other Ingredients		
Ingredient Kind	Ingredient Name	Quantity
INGR	WATER (UNII: 059QF0KO0R)	
INGR	GLYCERIN (UNII: PDC6A3C0OX)	
INGR	PHENOXYETHANOL (UNII: HIE492ZZ3T)	
INGR	METHYLPARABEN (UNII: A2I8 C7HI9 T)	
INGR	PROPYLPARABEN (UNII: Z8IX2SC1OH)	
INGR	BUTYLPARABEN (UNII: 3QPI1U3FV8)	
INGR	ETHYLPARABEN (UNII: 14255EXE39)	
INGR	ISOBUTYLPARABEN (UNII: 0 QQJ25X58G)	
INGR	SODIUM LAURO YL OAT AMINO ACIDS (UNII: FSW2K9B9N5)	
INGR	CO CAMIDO PRO PYL BETAINE (UNII: 50 CF3 O 11 KX)	
INGR	SODIUM LAURETH SULFATE (UNII: BPV390UAP0)	
INGR	ALOE VERA LEAF (UNII: ZY8 1Z8 3H0 X)	
INGR	SODIUM CHLORIDE (UNII: 451W47IQ8X)	

INGR	ALFALFA (UNII: DJO934BRBD)
INGR	CHAMO MILE (UNII: FGL3685T2X)
INGR	XANTHAN GUM (UNII: TTV12P4NEE)
INGR	D&C RED NO. 33 (UNII: 9DBA0SBB0L)
INGR	FD&C YELLOW NO. 5 (UNII: I753WB2F1M)

Product Characteristics					
Color	RED	Score			
Shape		Size			
Flavor		Imprint Code			
Contains					

ı	Packaging						
ı	#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
ı	1		60 mL in 1 BOTTLE, PLASTIC				

l	Marketing Information					
l	Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
l	COSMETIC		0 1/0 1/19 8 8			

Part 5 of 6

CONDITION AND ENHANCE TONER

face and neck (excluding shaving preparations) liquid

Product Information

Route of Administration TOPICAL

Other Ingredients				
Ingredient Kind	Ingredient Name	Quantity		
INGR	WATER (UNII: 059QF0KO0R)			
INGR	GLYCERIN (UNII: PDC6A3C0OX)			
INGR	SODIUM PYRROLIDONE CARBOXYLATE (UNII: 469 OTG57A2)			
INGR	DMDM HYDANTO IN (UNII: BYR0546 TOW)			
INGR	IODOPROPYNYL BUTYLCARBAMATE (UNII: 603P14DHEB)			
INGR	POTASSIUM ALUM (UNII: 1L24V9R23S)			
INGR	PANTHENOL (UNII: WV9CM0O67Z)			
INGR	SAGE (UNII: 065C5D077J)			
INGR	CALENDULA OFFICINALIS FLOWER (UNII: POM7O4Y7YD)			
INGR	POLYSORBATE 80 (UNII: 6 OZP39 ZG8 H)			
INGR	ALLANTO IN (UNII: 344S277G0Z)			

INGR	ALOE VERA LEAF (UNII: ZY81Z83H0X)	
INGR	FD&C BLUE NO. 1 (UNII: H3R47K3TBD)	
INGR	HAMAMELIS VIRGINIANA TOP WATER (UNII: NT00Y05A2V)	

Product Characteristics					
Color	BLUE	Score			
Shape		Size			
Flavor		Imprint Code			
Contains					

F	Packaging						
#	Item Code	Package Description	Marketing Start Date	Marketing End Date			
1		60 mL in 1 BOTTLE, PLASTIC					

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
COSMETIC		0 1/0 1/19 8 8			

Part 6 of 6

CONDITION AND ENHANCE EXFODERM FORTE

face and neck (excluding shaving preparations) lotion

Product Information

Route of Administration TOPICAL

Other Ingredients					
Ingredient Name	Quantity				
WATER (UNII: 059QF0KO0R)					
GLYCERIN (UNII: PDC6 A3C0 O X)					
METHYLPARABEN (UNII: A2I8 C7HI9 T)					
PROPYLPARABEN (UNII: Z8IX2SC1OH)					
POLYSORBATE 60 (UNII: CAL22UVI4M)					
CETO STEARYL ALCOHOL (UNII: 2DMT128M1S)					
GLYCOLIC ACID (UNII: 0 WT12SX38S)					
TROLAMINE (UNII: 9O3K93S3TK)					
MEDIUM-CHAIN TRIGLYCERIDES (UNII: C9H2L21V7U)					
LACTIC ACID (UNII: 33X04XA5AT)					
CETYL ALCOHOL (UNII: 936JST6JCN)					
EMU O IL (UNII: 344821WD61)					
STEARIC ACID (UNII: 4ELV7Z65AP)					
	WATER (UNII: 059QF0KO0R) GLYCERIN (UNII: PDC6A3C0OX) METHYLPARABEN (UNII: A2I8C7HI9T) PROPYLPARABEN (UNII: Z8IX2SC10H) POLYSORBATE 60 (UNII: CAL22UVI4M) CETOSTEARYL ALCOHOL (UNII: 2DMT128M1S) GLYCOLIC ACID (UNII: 0 WT12SX38S) TROLAMINE (UNII: 903K93S3TK) MEDIUM-CHAIN TRIGLYCERIDES (UNII: C9H2L21V7U) LACTIC ACID (UNII: 33X04XA5AT) CETYL ALCOHOL (UNII: 936JST6JCN) EMU OIL (UNII: 344821WD61)				

	INGR	STEARYL ALCOHOL (UNII: 2KR89I4H1Y)	
	INGR	DIMETHICO NE (UNII: 92RU3N3Y1O)	
1			ı

Product Characteristics				
Color	WHITE	Score		
Shape		Size		
Flavor		Imprint Code		
Contains				

P	Packaging					
#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
1		28.5 g in 1 BOTTLE, PLASTIC				

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
COSMETIC		0 1/0 1/19 8 8		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
UNAPPROVED DRUG OTHER		08/20/2007		

Labeler - OMP, INC. (790553353)

Establishment			
Name	Address	ID/FEI	Business Operations
PURETEK CORPORATION		785961046	MANUFACTURE(62032-510), LABEL(62032-510), PACK(62032-510)

Establishment					
Name	Address	ID/FEI	Business Operations		
Ei INC.		105803274	MANUFACTURE(62032-510), LABEL(62032-510), PACK(62032-510), ANALYSIS(62032-510)		

Establishment					
Name	Address	ID/FEI	Business Operations		
Swiss-American Products		611921669	MANUFACTURE(62032-510)		

Establishment				
Name	Address	ID/FEI	Business Operations	
Bay Cities Container Corporation		066229618	RELABEL(62032-510), REPACK(62032-510)	

Revised: 5/2012 OMP, INC.