

**NU-DERM SYSTEM NORMAL-OILY SKIN TRANSFORMATION- hydroquinone, homosalate, octisalate, and zinc oxide  
OBAGI COSMECEUTICAL LLC**

*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 NU-DERM® SYSTEM**

**OBAGI NU-DERM®**

**Rx only**

**For external use only**

**DESCRIPTION**

Hydroquinone, USP 4% is 1, 4-benzenediol. The drug is freely soluble in water and in alcohol. Chemically, hydroquinone is designated as p-dihydroxybenzene; the empirical formula is C<sub>6</sub>H<sub>6</sub>O<sub>2</sub>; molecular weight is 110.11 g/mol. The chemical structure is in the diagram below.



**Each gram of Obagi Nu-Derm® Clear contains:**

**ACTIVE:** Hydroquinone, USP 4% (40 mg/g)

**INACTIVES:** ascorbic acid, BHT, butylparaben, cetyl alcohol, disodium EDTA, glycerin, lactic acid, methylparaben, propylparaben, saponins, sodium lauryl sulfate, sodium metabisulfite, stearyl alcohol, tocopheryl acetate, water (aqua)

**Each gram of Obagi Nu-Derm Blender® contains:**

**ACTIVE:** Hydroquinone, USP 4% (40 mg/g)

**INACTIVES:** ascorbic acid, BHT, cetyl alcohol, disodium EDTA, glycerin, lactic acid, methylparaben, phenyl trimethicone, PPG-2 myristyl ether propionate, propylparaben, saponins, sodium lauryl sulfate, sodium metabisulfite, TEA-salicylate, tocopheryl acetate, water (aqua)

## **Each gram of Obagi Nu-Derm® Sunfader® contains:**

**ACTIVES:** Hydroquinone, USP 4% (40mg/g); Octinoxate, USP 7.5%; Oxybenzone, USP 5.5%

**INACTIVES:** ascorbic acid, BHT, butylparaben, cetyl alcohol, disodium EDTA, glycerin, methylparaben, propylparaben, saponins, sodium lauryl sulfate, sodium metabisulfite, stearyl alcohol, tocopheryl acetate, water (aqua)

## **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 sunscreen agents such as those contained in Obagi Nu-Derm Sunfader® and Obagi Nu-Derm® Sun Shield Matte Broad Spectrum SPF 50.

## **INDICATIONS AND USAGE**

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

## **CONTRAINDICATIONS**

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

## **WARNINGS**

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 product.

Test for skin sensitivity before using by applying a small amount to an unbroken patch of skin and check within 24 hours. Some users of this product may experience a mild skin irritation. If skin irritation becomes severe, stop use and consult a doctor. Close patient supervision is recommended.

## **WARNINGS**

**Avoid contact with eyes, nose, mouth and lips. In case of accidental contact, patient should rinse thoroughly with water and contact a physician. Sunscreen use is an essential aspect of hydroquinone therapy because even minimal sunlight exposure sustains melanocytic activity.**

**Contains sodium metabisulfite, a sulfite that may cause allergic-type reactions including anaphylactic symptoms and life-threatening or less**

**severe asthmatic episodes in certain susceptible people. The overall prevalence of sulfite sensitivity in the general population is unknown and probably low. Sulfite sensitivity is seen more frequently in asthmatic than in nonasthmatic people.**

## **PRECAUTIONS**

**(also see WARNINGS):**

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 product should be discontinued and the physician notified immediately.

**To report SUSPECTED ADVERSE REACTIONS, contact Obagi Cosmeceuticals LLC, at 1-800-636-7546 or FDA at 1-800-FDA-1088 or [www.fda.gov/medwatch](http://www.fda.gov/medwatch).**

## **DOSAGE AND ADMINISTRATION**

A thin application should be applied once or 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 or protective clothing to cover bleached skin when using and after using this product in order to prevent darkening from reoccurring.

## **HOW SUPPLIED**

**Obagi Nu-Derm® Clear** is available as follows:

Net wt. 2 oz. (57 g) bottle  
NDC 62032-101-36

**Obagi Nu-Derm Blender®** is available as follows:

Net wt. 2 oz. (57 g) bottle  
NDC 62032-100-36

Net wt. 1 oz. (28 g) bottle  
NDC 62032-100-10

**Obagi Nu-Derm Sunfader®** is available as follows:

Net wt. 2 oz. (57 g) bottle  
NDC 62032-116-36

Store at controlled room temperature: 15° to 25°C (59° to 77°F). Keep out of direct sunlight.

 **1-800-636-7546**

**Manufactured for:**

Obagi Cosmeceuticals LLC,  
Long Beach, CA 90806

All products/brand names, whether designated by notice or not (®/TM), are trademarks of Obagi Cosmeceuticals LLC and/or its affiliates.

©2019 Obagi Cosmeceuticals LLC.  
All rights reserved. [www.obagi.com](http://www.obagi.com)

Revised 01/2019 9458404

**Obagi Nu-Derm® Clear (Hydroquinone, USP 4%)  
Skin Bleaching Cream**

**Obagi Nu-Derm Blender® (Hydroquinone, USP 4%)  
Skin Bleaching Cream**

**Obagi Nu-Derm Sunfader®  
(Hydroquinone, USP 4%; Octinoxate, USP 7.5%; Oxybenzone, USP 5.5%) Skin  
Bleaching Cream with Sunscreens**

**Sun Shield Matte Broad Spectrum SPF 50  
Net wt. 3 oz. (85 g) AM**

***Drug Facts***

<b><i>Active ingredients</i></b>	<b><i>Purpose</i></b>
Homosalate 10%	Sunscreen
Octisalate 5%	Sunscreen
Zinc Oxide 16.5%	Sunscreen

**Uses**

- helps prevent sunburn
- If used as directed with other sun protection measures (see **Directions**), decreases the risk of skin cancer and early skin aging caused by the sun

## Warnings

### For external use only

**Do not use** on damaged or broken skin

**Stop use and ask a doctor** if rash occurs

**When using this product** keep out of eyes. Rinse with water to remove.

**Keep out of reach of children.** If swallowed, get medical help or contact a Poison Control Center right away.

## Directions

- apply liberally 15 minutes before sun exposure
- use a water resistant sunscreen if swimming or sweating
- reapply at least every 2 hours
- **Sun Protection Measures.** Spending time in the sun increases your risk of skin cancer and early skin aging. To decrease this risk, regularly use a sunscreen with a Broad Spectrum SPF value of 15 or higher and other sun protection measures including:
  - limit time in the sun, especially from 10 a.m.–2 p.m.
  - wear long-sleeved shirts, pants, hats, and sunglasses
- children under 6 months: Ask a doctor

## Other information

- store at controlled room temperature: 15°C–25°C (59°F–77°F)
- protect this product from excessive heat and direct sun

## Inactive ingredients

Water (Aqua), C15-19 Alkane, Octyldodecyl Neopentanoate, Polymethylsilsesquioxane, Sorbitan Olivatate, Silica, Polyglyceryl-6 Polyricinoleate, Sodium Chloride, Xanthan Gum, Glycerin, Hydroxyacetophenone, Disodium EDTA, 1,2-Hexanediol, Caprylyl Glycol, Sodium Hydroxide, Triethoxycaprylsilane, Polyhydroxystearic Acid, Distearidimonium Hectorite, Polyglyceryl-2 Isostearate, Euphorbia Cerifera (candelilla) Wax, Beeswax, Dimethicone

## Questions or comments?

**1.800.636.7546**

Monday–Friday 9 a.m.– 4 p.m. Pacific Time

Distributed by Obagi Cosmeceuticals LLC, Long Beach, CA 90806

**PRINCIPAL DISPLAY PANEL - Kit Carton**

NDC# 62032-531-00

OBAGI®  
MEDICAL

OBAGI NU-DERM® SYSTEM

NORMAL OILY

Skin Transformation Kit

NORMAL OILY

OBAGI MEDICAL

NDC# 65085-001-00

OBAGI MEDICAL

FRONT PANEL OBAGI NU-DERM® SYSTEM

NORMAL OILY Skin Transformation Kit

NORMAL OILY

OBAGI MEDICAL

92032 51900 5

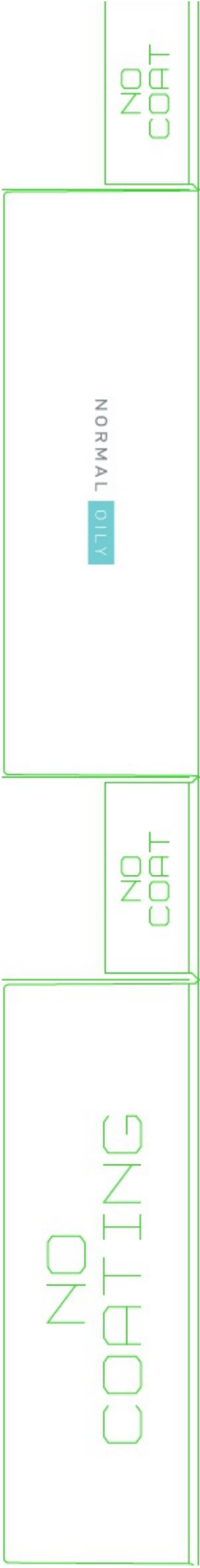
NORMAL OILY Skin Transformation Kit



The #1 prescription-strength, physician-dispensed skincare system. A total systemic approach to skin care. Addresses signs of skin aging, gently exfoliates to promote cell turnover and suppresses melanocyte activity to reduce hyperpigmentation.

OBAGI MEDICAL

Vertical text on the left side of the page, including a barcode and various product details.



# NU-DERM SYSTEM NORMAL-OILY SKIN TRANSFORMATION

hydroquinone, homosalate, octisalate, and zinc oxide kit

## Product Information

<b>Product Type</b>	HUMAN PRESCRIPTION DRUG	<b>Item Code (Source)</b>	NDC:62032-531
---------------------	-------------------------	---------------------------	---------------

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:62032-531-00	1 in 1 CARTON	12/02/2019	

## Quantity of Parts

Part #	Package Quantity	Total Product Quantity
<b>Part 1</b>	1 BOTTLE, PLASTIC	57 g
<b>Part 2</b>	1 BOTTLE, PLASTIC	198 mL
<b>Part 3</b>	1 BOTTLE, PLASTIC	198 mL
<b>Part 4</b>	1 BOTTLE, PLASTIC	57 g
<b>Part 5</b>	1 BOTTLE, PLASTIC	48 g
<b>Part 6</b>	1 BOTTLE, PLASTIC	57 g
<b>Part 7</b>	1 TUBE	85 g

## Part 1 of 7

### OBAGI NU-DERM EXFODERM FORTE

face and neck (excluding shaving preparations) lotion

## Product Information

<b>Route of Administration</b>	TOPICAL
--------------------------------	---------

## Other Ingredients

Ingredient Kind	Ingredient Name	Quantity
INGR	<b>WATER</b> (UNII: 059QF0KO0R)	
INGR	<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
INGR	<b>DIMETHICONE</b> (UNII: 92RU3N3Y1O)	
INGR	<b>METHYLPARABEN</b> (UNII: A2I8C7HI9T)	
INGR	<b>PROPYLPARABEN</b> (UNII: Z8IX2SC1OH)	
INGR	<b>POLYSORBATE 60</b> (UNII: CAL22UVI4M)	
INGR	<b>TROLAMINE</b> (UNII: 9O3K93S3TK)	
INGR	<b>MEDIUM-CHAIN TRIGLYCERIDES</b> (UNII: C9H2L21V7U)	
INGR	<b>CETOSTEARYL ALCOHOL</b> (UNII: 2DMT128M1S)	
INGR	<b>CETYL ALCOHOL</b> (UNII: 936JST6JCN)	

INGR	<b>LACTIC ACID, UNSPECIFIED FORM</b> (UNII: 33X04XA5AT)	
INGR	<b>EMU OIL</b> (UNII: 344821WD61)	
INGR	<b>STEARIC ACID</b> (UNII: 4ELV7Z65AP)	
INGR	<b>STEARYL ALCOHOL</b> (UNII: 2KR89I4H1Y)	
INGR	<b>Glycolic Acid</b> (UNII: 0WT12SX38S)	

### Product Characteristics

<b>color</b>	WHITE	C48325
--------------	-------	--------

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1		57 g in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
COSMETIC		01/01/1988	

## Part 2 of 7

### OBAGI NU-DERM FOAMING

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

### Product Information

<b>Route of Administration</b>	TOPICAL
--------------------------------	---------

### Other Ingredients

Ingredient Kind	Ingredient Name	Quantity
INGR	<b>WATER</b> (UNII: 059QF0K00R)	
INGR	<b>PHENOXYETHANOL</b> (UNII: HIE492ZZ3T)	
INGR	<b>METHYLPARABEN</b> (UNII: A2I8C7HI9T)	
INGR	<b>PROPYLPARABEN</b> (UNII: Z8IX2SC1OH)	
INGR	<b>BUTYLPARABEN</b> (UNII: 3QPI1U3FV8)	
INGR	<b>ETHYLPARABEN</b> (UNII: 14255EXE39)	
INGR	<b>ISOBUTYLPARABEN</b> (UNII: 0QQJ25X58G)	
INGR	<b>FD&amp;C YELLOW NO. 5</b> (UNII: I753WB2F1M)	
INGR	<b>SODIUM LAUROYL OAT AMINO ACIDS</b> (UNII: FSW2K9B9N5)	
INGR	<b>COCAMIDOPROPYL BETAINE</b> (UNII: 5OCF3O11KX)	
INGR	<b>SODIUM LAURETH-3 SULFATE</b> (UNII: BPV390UAP0)	
INGR	<b>ALOE VERA LEAF</b> (UNII: ZY81Z83H0X)	

INGR	<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
INGR	<b>MEDICAGO SATIVA WHOLE</b> (UNII: DJO934BRBD)	
INGR	<b>CHAMOMILE</b> (UNII: FGL3685T2X)	
INGR	<b>XANTHAN GUM</b> (UNII: TTV12P4NEE)	
INGR	<b>D&amp;C RED NO. 33</b> (UNII: 9DBA0SBB0L)	

### Product Characteristics

<b>color</b>	RED	C48326
--------------	-----	--------

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1		198 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
COSMETIC		01/01/1988	

## Part 3 of 7

### OBAGI NU-DERM TONER

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

### Product Information

<b>Route of Administration</b>	TOPICAL
--------------------------------	---------

### Other Ingredients

Ingredient Kind	Ingredient Name	Quantity
INGR	<b>WATER</b> (UNII: 059QF0KO0R)	
INGR	<b>GLYCERIN</b> (UNII: PDC6A3C00X)	
INGR	<b>HAMAMELIS VIRGINIANA TOP WATER</b> (UNII: NT00Y05A2V)	
INGR	<b>SODIUM PYRROLIDONE CARBOXYLATE</b> (UNII: 469OTG57A2)	
INGR	<b>DMDM HYDANTOIN</b> (UNII: BYR0546TOW)	
INGR	<b>IODOPROPYNYL BUTYLCARBAMATE</b> (UNII: 603P14DHEB)	
INGR	<b>POTASSIUM ALUM</b> (UNII: 1L24V9R23S)	
INGR	<b>PANTHENOL</b> (UNII: WW9CM0067Z)	
INGR	<b>SAGE</b> (UNII: 065C5D077J)	
INGR	<b>CALENDULA OFFICINALIS FLOWER</b> (UNII: P0M7O4Y7YD)	
INGR	<b>POLYSORBATE 80</b> (UNII: 6OZP39ZG8H)	
INGR	<b>ALLANTOIN</b> (UNII: 344S277G0Z)	

INGR	<b>ALOE VERA LEAF</b> (UNII: ZY81Z83H0X)	
INGR	<b>FD&amp;C BLUE NO. 1</b> (UNII: H3R47K3TBD)	

### Product Characteristics

<b>color</b>	BLUE	C48333
--------------	------	--------

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1		198 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
COSMETIC		01/02/1988	

## Part 4 of 7

## NU-DERM CLEAR SKIN BLEACHING AND CORRECTOR

hydroquinone cream

### Product Information

<b>Route of Administration</b>	TOPICAL
--------------------------------	---------

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>HYDROQUINONE</b> (UNII: XV74C1N1AE) (HYDROQUINONE - UNII:XV74C1N1AE)	HYDROQUINONE	40 mg in 1 g

### Inactive Ingredients

Ingredient Name	Strength
<b>EDETATE DISODIUM</b> (UNII: 7FLD91C86K)	
<b>BUTYLPARABEN</b> (UNII: 3QPI1U3FV8)	
<b>STEARYL ALCOHOL</b> (UNII: 2KR89I4H1Y)	
<b>SODIUM LAURYL SULFATE</b> (UNII: 368GB5141J)	
<b>CETYL ALCOHOL</b> (UNII: 936JST6JCN)	
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>LACTIC ACID, UNSPECIFIED FORM</b> (UNII: 33X04XA5AT)	
<b>.ALPHA.-TOCOPHEROL ACETATE</b> (UNII: 9E8X80D2L0)	
<b>ASCORBIC ACID</b> (UNII: PQ6CK8PD0R)	

**SODIUM METABISULFITE** (UNII: 4VON5FNS3C)

**WATER** (UNII: 059QF0KO0R)

**METHYLPARABEN** (UNII: A2I8C7HI9T)

**PROPYLPARABEN** (UNII: Z8IX2SC1OH)

**BUTYLATED HYDROXYTOLUENE** (UNII: 1P9D0Z171K)

### Product Characteristics

<b>Color</b>	WHITE	<b>Score</b>	
<b>Shape</b>		<b>Size</b>	
<b>Flavor</b>		<b>Imprint Code</b>	
<b>Contains</b>			

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1		57 g in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
UNAPPROVED DRUG OTHER		01/01/1988	

## Part 5 of 7

### OBAGI NU-DERM HYDRATE FACIAL MOISTURIZER

face and neck (excluding shaving preparations)

### Product Information

**Route of Administration** TOPICAL

### Other Ingredients

Ingredient Kind	Ingredient Name	Quantity
INGR	<b>WATER</b> (UNII: 059QF0KO0R)	
INGR	<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
INGR	<b>MEDIUM-CHAIN TRIGLYCERIDES</b> (UNII: C9H2L21V7U)	
INGR	<b>SODIUM HYDROXIDE</b> (UNII: 55X04QC32I)	
INGR	<b>CAESALPINIA SPINOSA RESIN</b> (UNII: WL3883U2PO)	
INGR	<b>SHEA BUTTER</b> (UNII: K49155WL9Y)	
INGR	<b>DIMETHICONE/VINYL DIMETHICONE CROSSPOLYMER (SOFT PARTICLE)</b> (UNII: 9E4CO0W6C5)	

INGR	<b>CYCLOMETHICONE 5</b> (UNII: 0THT5PCI0R)	
INGR	<b>CETYL ALCOHOL</b> (UNII: 936JST6JCN)	
INGR	<b>SACCHARIDE ISOMERATE</b> (UNII: W8K377W98I)	
INGR	<b>DIMETHICONE</b> (UNII: 92RU3N3Y1O)	
INGR	<b>TOCOPHEROL</b> (UNII: R0ZB2556P8)	
INGR	<b>LAURETH-12</b> (UNII: OAH19558U1)	
INGR	<b>PHENOXYETHANOL</b> (UNII: HIE492ZZ3T)	
INGR	<b>ETHYLHEXYLGLYCERIN</b> (UNII: 147D247K3P)	
INGR	<b>STEARIC ACID</b> (UNII: 4ELV7Z65AP)	
INGR	<b>AVOCADO OIL</b> (UNII: 6VNO72PFC1)	
INGR	<b>SOY STEROL</b> (UNII: PL360EPO9J)	
INGR	<b>CAPRYLYL GLYCOL</b> (UNII: 00YIU5438U)	
INGR	<b>LEVOMENOL</b> (UNII: 24WE03BX2T)	
INGR	<b>HEXYLENE GLYCOL</b> (UNII: KEH0A3F75J)	
INGR	<b>TETRAHYDRODIFERULOYLMETHANE</b> (UNII: 00U0645U03)	
INGR	<b>PANTHENOL</b> (UNII: W99CM0067Z)	
INGR	<b>MANGIFERA INDICA SEED BUTTER</b> (UNII: 4OXD9M35X2)	
INGR	<b>SODIUM STEAROYL GLUTAMATE</b> (UNII: 65A9F4P024)	
INGR	<b>CARBOMER HOMOPOLYMER, UNSPECIFIED TYPE</b> (UNII: 0A5MM307FC)	
INGR	<b>ALLANTOIN</b> (UNII: 344S277G0Z)	
INGR	<b>GLYCERYL MONOSTEARATE</b> (UNII: 230OU9XXE4)	

### Product Characteristics

<b>color</b>	WHITE	C48325
--------------	-------	--------

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1		48 g in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
COSMETIC		11/07/2012	

### Part 6 of 7

## NU-DERM BLENDER SKIN LIGHTENER AND BLENDING

hydroquinone cream

### Product Information

<b>Route of Administration</b>	TOPICAL
--------------------------------	---------

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>HYDROQUINONE</b> (UNII: XV74C1N1AE) (HYDROQUINONE - UNII:XV74C1N1AE)	HYDROQUINONE	40 mg in 1 g

## Inactive Ingredients

Ingredient Name	Strength
<b>EDETATE DISODIUM</b> (UNII: 7FLD91C86K)	
<b>PPG-2 MYRISTYL ETHER PROPIONATE</b> (UNII: 88R97D8U8A)	
<b>TROLAMINE SALICYLATE</b> (UNII: H8O4040BHD)	
<b>SODIUM LAURYL SULFATE</b> (UNII: 368GB5141J)	
<b>CETYL ALCOHOL</b> (UNII: 936JST6JCN)	
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>LACTIC ACID, UNSPECIFIED FORM</b> (UNII: 33X04XA5AT)	
<b>.ALPHA.-TOCOPHEROL ACETATE</b> (UNII: 9E8X80D2L0)	
<b>ASCORBIC ACID</b> (UNII: PQ6CK8PD0R)	
<b>SODIUM METABISULFITE</b> (UNII: 4VON5FNS3C)	
<b>WATER</b> (UNII: 059QF0KO0R)	
<b>METHYLPARABEN</b> (UNII: A2I8C7HI9T)	
<b>PROPYLPARABEN</b> (UNII: Z8IX2SC1OH)	
<b>BUTYLATED HYDROXYTOLUENE</b> (UNII: 1P9D0Z171K)	
<b>PHENYL TRIMETHICONE</b> (UNII: DR0K5NOJ4R)	

## Product Characteristics

<b>Color</b>	WHITE	<b>Score</b>	
<b>Shape</b>		<b>Size</b>	
<b>Flavor</b>		<b>Imprint Code</b>	
<b>Contains</b>			

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1		57 g in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
UNAPPROVED DRUG OTHER		01/01/1988	

# NU-DERM SUN SHIELD MATTE BROAD SPECTRUM SPF 50 SUNSCREEN

homosalate, octisalate, and zinc oxide lotion

## Product Information

**Route of Administration** TOPICAL

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>HOMOSALATE</b> (UNII: V06SV4M95S) (HOMOSALATE - UNII:V06SV4M95S)	HOMOSALATE	100 mg in 1 g
<b>OCTISALATE</b> (UNII: 4X49Y0596W) (OCTISALATE - UNII:4X49Y0596W)	OCTISALATE	50 mg in 1 g
<b>ZINC OXIDE</b> (UNII: SOI2LOH54Z) (ZINC OXIDE - UNII:SOI2LOH54Z)	ZINC OXIDE	165 mg in 1 g

## Inactive Ingredients

Ingredient Name	Strength
<b>WATER</b> (UNII: 059QF0KO0R)	
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>Hydroxyacetophenone</b> (UNII: G1L3HT4CMH)	
<b>XANTHAN GUM</b> (UNII: TTV12P4NEE)	
<b>Sodium Chloride</b> (UNII: 451W47IQ8X)	
<b>Sodium Hydroxide</b> (UNII: 55X04QC32I)	
<b>Octyldodecyl Neopentanoate</b> (UNII: X8725R883T)	
<b>Triethoxycaprylylsilane</b> (UNII: LDC331P08E)	
<b>C15-19 Alkane</b> (UNII: CI87N1IM01)	
<b>Disteardimonium Hectorite</b> (UNII: X687XDK09L)	
<b>POLYGLYCERYL-2 ISOSTEARATE</b> (UNII: 7B8OE71MQC)	
<b>Sorbitan Olivat</b> (UNII: MDL271E3GR)	
<b>DIMETHICONE</b> (UNII: 92RU3N3Y1O)	
<b>1,2-HEXANEDIOL</b> (UNII: TR046Y3K1G)	
<b>CAPRYLYL GLYCOL</b> (UNII: 00YIU5438U)	
<b>EDETATE DISODIUM</b> (UNII: 7FLD91C86K)	
<b>CANDELILLA WAX</b> (UNII: WL0328HX19)	
<b>Yellow Wax</b> (UNII: 2ZA36H0S2V)	
<b>POLYMETHYLSILSESQUIOXANE (4.5 MICRONS)</b> (UNII: 59Z907ZB69)	
<b>Silicon Dioxide</b> (UNII: ETJ7Z6XBU4)	

## Product Characteristics

<b>Color</b>	WHITE	<b>Score</b>	
<b>Shape</b>		<b>Size</b>	
<b>Flavor</b>		<b>Imprint Code</b>	
<b>Contains</b>			

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1		85 g in 1 TUBE; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC MONOGRAPH FINAL	part352	12/02/2019	

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
UNAPPROVED DRUG OTHER		12/02/2019	

**Labeler** - OBAGI COSMECEUTICAL LLC (790553353)

## Establishment

Name	Address	ID/FEI	Business Operations
PURETEK CORPORATION		785961046	MANUFACTURE(62032-531) , LABEL(62032-531) , PACK(62032-531)

## Establishment

Name	Address	ID/FEI	Business Operations
G.S. COSMECEUTICAL USA, INC.		017014734	MANUFACTURE(62032-531)

## Establishment

Name	Address	ID/FEI	Business Operations
Swiss-American CDMO, LLC		080170933	MANUFACTURE(62032-531)

Revised: 1/2023

OBAGI COSMECEUTICAL LLC