DR LIFT ANTIBACTERIAL BODY WASH- benzalkonium chloride gel Spa de Soleil

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.

Dr Lift Antibacterial Body Wash

Active Ingredient

Benzalkonium Chloride 0.13 %

Purpose

Antiseptic

Warnings

Warnings

For external use only.

When using this product: If in eyes, rinse promptly and thoroughly with water. Discontinue use if irritation or redness develop.

Stop use and ask a doctor if irritation or redness occur for more than 72 hours.

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

Keep out of reach of children.

Directions

Apply to wet skin. Scrub thoroughly. Rinse and dry. Use as needed.

Place enough product in the palm of your hands to thoroughly cover you hands. Rub hands together briskly until product is completely absorbed and hands are dry.

Inactive Ingredients:

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Sodium Chloride, Cocamidopropyl betaine, Aqua, Sodium Laureth Sulfate, Carbomer, Glycerin, *CO Humulus Lupulus (Hops) Extract, *CO Lavandula Angustifolia (Lavender) Flower/Leaf/Stem Extract, *CO Calendula Officinalis Flower Extract, *CO Chamomilla Recutita (Matricaria) Flower Extract, *CO Citrus Limon (Lemon) Peel Extract, *CO Cucumis Sativus (Cucumber) Seed Extract, *CO Camellia Sinensis Leaf Extract, *CO Pyrus Malus (Apple) Fruit Extract, *CO Spirulina Platensis Extract, Xanthan Gum.

*CO Certified Organic



| Product Information | | | | | | |
|---|----------------------|-----------------------------------|--|----------------------|----------------|--|
| Product T ype | HUMAN OTC DRUG | HUMAN OTC DRUG Item Code (Source) | | | NDC:68062-2241 | |
| Route of Administration | TOPICAL | TOPICAL | | | | |
| | | | | | | |
| Active Ingredient/Active | Mojety | | | | | |
| Ingredient Name Basis of Streng | | | | | Strength | |
| BENZALKONIUM CHLORIDE (UNII:7N6JUD5X6Y) | NUM - | BENZALKONIUM CHLORIDE | | 0.46 mg in 240 mg | | |
| | | | | | | |
| | | | | | | |
| Inactive Ingredients | | | | | | |
| Inactive Ingredients | Ingredient Name | | | | Strength | |
| | 0 | | | | Strength | |
| COCAMIDOPROPYL BETAINE | 0 | | | | Strength | |
| Inactive Ingredients COCAMIDOPROPYL BETAINE WATER (UNII: 059QF0K00R) SODIUM CHLORIDE (UNII: 451V | C (UNII: 50CF3011KX) | | | | Strength | |
| WATER (UNII: 059QF0KO0R) | C (UNII: 50CF3011KX) | | | | Strength | |
| COCAMIDOPROPYL BETAINE WATER (UNII: 059QF0KO0R) | C (UNII: 50CF3011KX) | | | | Strength | |
| COCAMIDOPROPYL BETAINE WATER (UNII: 059QF0KO0R) | C (UNII: 50CF3011KX) | | | | Strength | |

| Marketing Information | | | | | | | |
|-------------------------|--|----------------------|--------------------|--|--|--|--|
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | | | | |
| OTC monograph not final | part333E | 07/29/2020 | | | | | |
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Labeler - Spa de Soleil (874682867)

Establishment

| Name | Address | ID/FEI | Business Operations |
|---------------|---------|-----------|-------------------------|
| Spa de Soleil | | 874682867 | manufacture(68062-2241) |

Revised: 7/2020

Spa de Soleil