

**DR. PIERCING AFTERCARE- benzalkonium chloride swab
Swabplus, L.P.**

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. Piercing Aftercare 36CT

Use

Use: FDA Registered antiseptic for the care and cleaning of piercings.

Directions:

1. Hold swab with the colored round marker end up.
2. Bend the colored round marker tip until it "snaps".
3. Liquid flows down into the white tip.
4. Apply medicated tip to piercing. Dispose.

WARNINGS:

WARNINGS: For external use only. If redness, irritation, infection or swelling occurs, discontinue use and consult a physician. Keep out of eyes. Do not insert into the ear canal. Store at room temperature.

KEEP OUT OF REACH OF CHILDREN

Keep out of reach of children.

ACTIVE INGREDIENTS

ACTIVE INGREDIENTS: Benzalkonium Chloride 0.11%

OTHER INGREDIENTS

OTHER INGREDIENTS: Isotonic Saline Solution, Aloe Vera, Allantoin, Panthenol/Pro-Vitamin B5.

DISTRIBUTED BY

DISTRIBUTED BY: Broadway Products LLC, Sellersburg, Indiana.

LEARN MORE: www.drpiercing.com

BARCODE XM001448MO3

Dr. Piercing Afterca...roduct, 36 Swab Pierce New

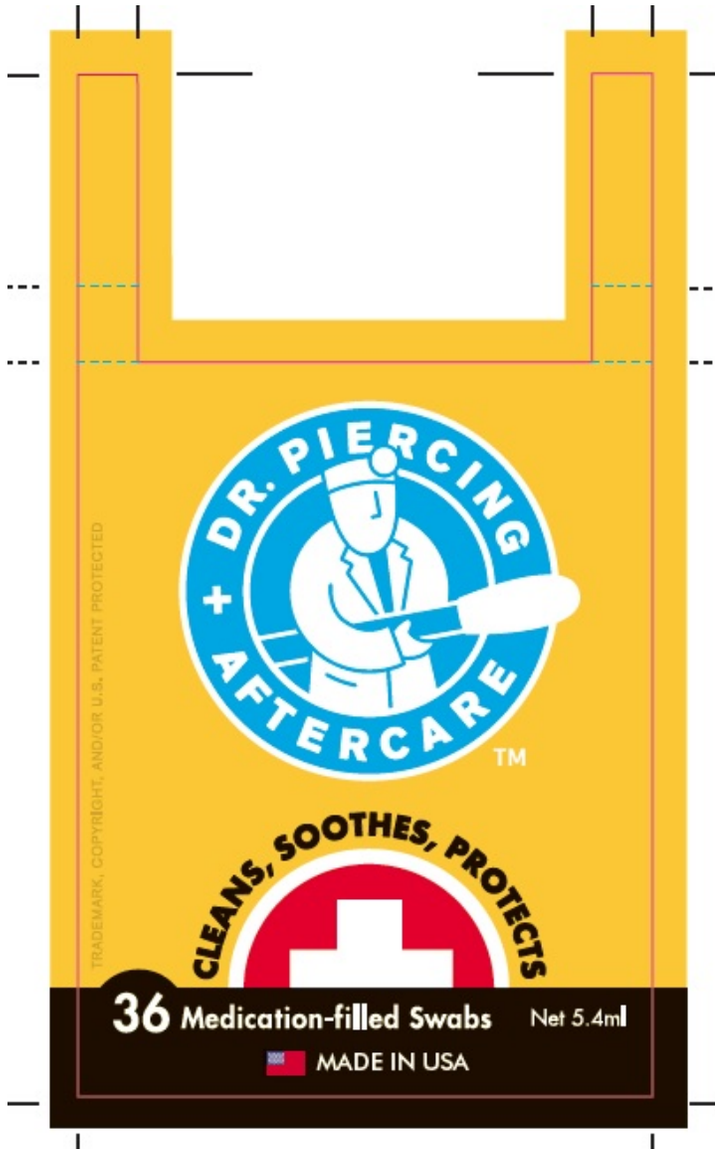
Principal Display Label

DR. PIERCING AFTERCARE™

CLEAN, SOOTHES, PROTECTS

36 Medication-filled Swabs Net 5.4mL

MADE IN USA



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X001448MO3

Dr. Piercing Afterca...roduct, 36 Swab Piece
New



DR. PIERCING AFTERCARE

benzalkonium chloride swab

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:65734-504
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7) (BENZALKONIUM - UNII: 7N6JUD5X6Y)	BENZALKONIUM CHLORIDE	1.1 mg in 1 mL

Inactive Ingredients

Ingredient Name	Strength
ALOE VERA LEAF (UNII: ZY81Z83H0X)	
METHYLCHLOROISOTHIAZOLINONE (UNII: DEL7T5QRPN)	
MAGNESIUM CHLORIDE (UNII: 02F3473H9O)	
PANTHENOL (UNII: WW9CM0O67Z)	
METHYLISOTHIAZOLINONE (UNII: 229D0E1QFA)	
MAGNESIUM NITRATE (UNII: 77CBG3UN78)	
CUPRIC NITRATE (UNII: 9TC879S2ZV)	
WATER (UNII: 059QF0KOOR)	
COCAMIDOPROPYL BETAINE (UNII: 5OCF3O11KX)	
TROLAMINE (UNII: 9O3K93S3TK)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
ALLANTOIN (UNII: 344S277G0Z)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65734-504-36	36 in 1 CASE	01/01/2021	
1		0.15 mL in 1 APPLICATOR; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333A	01/01/2021	

Labeler - Swabplus, L.P. (116984439)

Revised: 1/2023

Swabplus, L.P.