## TRI-SOFT SOFTENING- benzalkonium chloride solution Clinical Therapeutic Solutions

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## TRI-SOFT SOFTENING SOLUTION

**TRI-SOFT SOFTENING SOLUTION** 

## **Active Ingredient**

BENZALKONIUM CHLORIDE (0.13 %)

### Purpose

Antiseptic

#### Uses:

For antiseptic cleansing that will decrease bacteria on the skin without soap and water.

### Warnings

For external use only.

Do not use in the eyes. If eye contact occurs, rinse thoroughly with water.

Stop use and consult a physician if irritation or redness develops and persists for more than 72 hours.

Do not bandage tightly.

If pregnant or breast feeding, contact physician prior to use.

### Keep out of reach of children

if swallowed, consult physician or poison control immediately

### Directions

Mix 1 part Tri-Soft Concentrate with water q. s. to 16 parts. Saturate cotton and place over hyperkeratotic tissues for 3 minutes before reduction or debridement.

### **Other Ingredients**

Aqua (Deionized Water), Benzyl Alcohol, Phenylmercuric Nitrate, Polysorbate-20, SD-

Alcohol 40B.

Tri-Soft Softening Solution is a rapid-acting emulsifying agent combined with antiseptic properties.

It is specifically formulated for use on the skin prior to instrument reduction of hyperkeratotic lesions.

# TRI-SOFT SOFTENING SOLUTION 8oz (44577-711-08)



For External Use Only 8 FL OUNCES

PROFESSIONAL

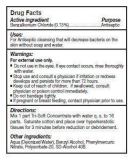
Tri-Soft Softening Solution is a rapid-acting emulsifying agent combined with antiseptic properties. It is specifically formulated for use on the skin prior to instrument reduction of hyperkeratotic lesions.

Contains: Benzalkonium Chloride in a Phenylmercuric Nitrate Solution

Directions: Mix 1 part Tri-Soft Concentrate with water q. s. to 16 parts. Saturate cotton and place over hyperkeratotic tissues for 3 minutes before reduction or debridement.

For External Use Only







TRI-SOFT SOFTENIN	G					
benzalkonium chloride solutio	on					
Product Information						
Product Type	HUMAN OTC DRUG	ltem Code (Source)		NDC:44577-711		
Route of Administration	TOPICAL					
Active Ingredient/Active	Moiety					
Ingre	dient Name		Basis of Stre	ength	Strength	
<b>BENZALKONIUM CHLORIDE</b> (UNII: F5UM2KM3W7) (BENZALKONIUM - UNII: 7N6JUD5X6Y)			BENZALKONIUM CHLORIDE		1.3 mg in 1 mL	
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Inactive Ingredients						
Ingredient Name					Strength	
WATER (UNII: 059QF0KO0R)						

	NZYL ALCOHOL			
PH	IENYLMERCURIC			
PC	DLYSORBATE 20			
AL	COHOL (UNII: 3Kg			
Pè	ackaging			
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1		236.59 mL in 1 BOTTLE; Type 0: Not a Combination Product	02/22/2017	
Μ	larketing I	nformation		
	Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
~ -	C Monograph Dru	g 505G(a)(3)	12/06/2012	
ΟI				

Labeler - Clinical Therapeutic Solutions (078402750)

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**Clinical Therapeutic Solutions**