

**99 PLUS HAND SANITIZER- benzealkonium chloride liquid  
Momar Incorporated**

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**99 Plus Instant Hand Sanitizer Drug Facts and Label**

**Drug Facts Box OTC-Active Ingredient Section**

benzalkonium chloride USP 0.13%

**Drug Facts Box OTC-Indications & Usage Section**

For hand-washing to decrease bacteria on the skin, only when water is not available

**Drug Facts Box OTC-Warnings Section**

For external use only

**Drug Facts Box OTC-Purpose Section**

Antiseptic

**Drug Facts Box-OTC When Using Section**

do not get into eyes

if contact occurs, rinse eyes thoroughly with water

**Drug Facts Box-OTC Stop Use Section**

irritation and redness develop

**Drug Facts Box-OTC Keep Out Of Reach Of Children Section**

If swallowed, get medical help or contact a Poison Control Center right away

**Drug Facts Box-OTC Dosage & Administration Section**

press pump twice to deliver two squirts (about a quarter size) of foaming product onto the palm of your hand

rub hands together until dry

wash hands with soap and water at earliest opportunity

**Drug Facts Box-OTC Inactive Ingredient Section**

water, glycerine, dimethicone, DMDM hydantoin, iodopropynyl butylcarbamate, methylchloroisothiazolinone, methylisothiazolinone, fragrance

## 99 Plus Instant Hand Sanitizer



Drug Facts	
<b>Active Ingredient</b>	benzalkonium chloride 0.13%
<b>Use</b>	for hand-washing to decrease the water is not available
<b>Warnings</b>	For external use only When using this product Do not get into eyes If contact occurs, rinse eyes thoroughly Stop use and ask a doctor if irritation and redness develop Keep out of reach of children. If contact a Poison Control Center right
<b>Directions</b>	press pump twice (a quarter size) of foaming product or rub hands together until dry water at earliest opportunity
<b>Inactive Ingredients</b>	water, DMDM hydantoin, iodopropynyl butylthiazolinone, methylisothiazolinone, fragrance

## 99 Plus Instant Hand Sanitizer

### 99 PLUS HAND SANITIZER

benzealkonium chloride liquid

#### Product Information

<b>Product Type</b>	HUMAN OTC DRUG	<b>Item Code (Source)</b>	NDC:63533-457
<b>Route of Administration</b>	TOPICAL		

#### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>BENZALKONIUM CHLORIDE</b> (UNII: F5UM2KM3W7) (BENZALKONIUM - UNII: 7N6JUD5X6Y)	BENZALKONIUM CHLORIDE	1.3 mg in 1 mL

#### Inactive Ingredients

Ingredient Name	Strength
<b>WATER</b> (UNII: 059QF0KO0R)	
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>DIMETHICONE</b> (UNII: 92RU3N3Y1O)	
<b>DMDM HYDANTOIN</b> (UNII: BYR0546TOW)	
<b>IODOPROPYNYL BUTYLCARBAMATE</b> (UNII: 603P14DHEB)	
<b>METHYLCHLOROISOTHIAZOLINONE</b> (UNII: DEL7T5QRPN)	

**METHYLISOTHIAZOLINONE** (UNII: 229D0E1QFA)

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:63533-457-10	1000 mL in 1 CARTRIDGE; Type 0: Not a Combination Product	06/28/2019	
2	NDC:63533-457-18	50 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	06/28/2019	
3	NDC:63533-457-14	3785 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	06/28/2019	
4	NDC:63533-457-12	1000 mL in 1 BAG; Type 0: Not a Combination Product	06/28/2019	

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M003	06/28/2019	

**Labeler** - Momar Incorporated (003266616)

**Registrant** - ABC Compounding Co., Inc. (003284353)

### Establishment

Name	Address	ID/FEI	Business Operations
ABC Compounding Co., Inc.		003284353	manufacture(63533-457)

Revised: 1/2026

Momar Incorporated