



5/12615068/1020

Directions for use  
Read carefully!



# B. Braun 0.9% Sodium Chloride Injection B.P.

## Composition

100 ml of solution contain

### Active ingredients:

Sodium Chloride 0.9 g

### Excipients:

Water for injections

Theoretical osmolarity: 308 mOsm/l

Titration acidity: < 0.3 mmol/l

pH: 4.5 - 7.0

### Electrolyte concentrations:

Sodium 154 mmol/l

Chloride 154 mmol/l

## Pharmaceutical form

Solution for injection

## Pharmaco-therapeutic group

Vehicle solution

## Indications

Solvent or diluent for compatible electrolyte concentrates or drugs.

## Contraindications

None

## Precautions for use and special warnings

B. Braun 0.9% Sodium Chloride Injection B.P. should only be administered with particular caution to patients with

- hypernatraemia
- hyperchloraemia

## Interactions

When mixing with other medicaments, possible incompatibilities should be considered.

## Dosage

The quantity to be chosen depends on the desired concentration of the medicament to be dissolved.

For the use of this solution as solvent/diluent for compatible electrolyte concentrates or medicaments, the instructions for use relating to the medicament to be added should be observed.

### Method of administration

Intravenous or subcutaneous injection.

## Overdose

### Symptoms

Overdose of B. Braun 0.9% Sodium Chloride solution may result in hypernatraemia, hyperchloraemia, hyperhydration, hyperosmolarity of the serum, and metabolic acidosis.

### Emergency treatment, antidotes

Immediate stop of administration, administration of diuretics with continuous monitoring of serum electrolytes, correction of electrolyte and acid-base imbalances.

## Undesirable effects

Administration of larger amounts of the solution may lead to hypernatraemia and hyperchloraemia.

**Note:** Patients are advised to inform their doctor or pharmacist of any adverse reaction they experience in connection with the administration of this drug.

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Dokument = 148 x 210 mm

DIN A5

2 Seiten

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GIF (MPc, MPw) – folded

0.9% Sodium Chloride Injection B.P.

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Production site: Penang

Lätus



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### Usage during pregnancy

No adverse reaction has been reported. However, risk versus benefits should be assessed before using.

### Storage

The product should not be stored above the temperature stated on the label.

### Shelf life

The product must not be used beyond the expiry date stated on the label.

### Presentation

In plastic ampoules "Mini-Plasco®" and "Mini-Plasco® Connect" of 5 ml, 10 ml, 20 ml in box of 20's.

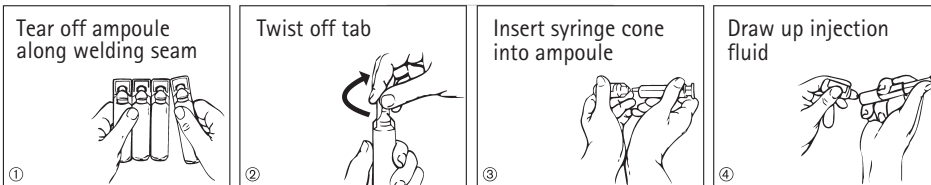
### Instruction for use/handling

The solution is supplied in single-dose containers. Discard unused contents.

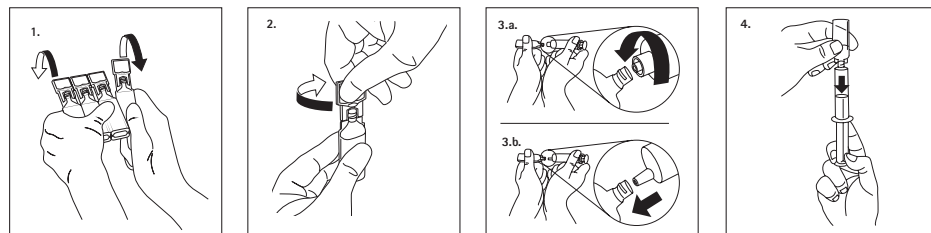
Use contents immediately after opening the container.

Only to be used if solution is clear and the container undamaged.

### Mini-Plasco® Handling



### Mini-Plasco® Connect Handling



**B | BRAUN**

Product registration holder  
and manufactured by:  
**B. Braun Medical Industries  
Sdn. Bhd.**  
11900 Bayan Lepas, Penang,  
Malaysia.

