Directions for Use





B. Braun Water for Injections B.P

Solvent for parenteral use

Composition

100 ml of solvent contain Water for injections

100 ml

Pharmaceutical form

Solvent for parenteral use Clear, colourless solution

Pharmacotherapeutic group

Solvents and diluting agents, incl. irrigating solutions, ATC code: V07AB

Pharmacodynamic and Pharmacokinetic properties

Since B. Braun Water for Injections is only used as a vehicle solution for administration of additives, pharmacodynamic and pharmacokinetic properties depend on the respective additives.

Indications

Preparation and dilution of parenteral preparations

Contraindications

None known if used according to the instructions given.

Special warnings and precautions for use

B. Braun Water for Injections is strongly hypotonic. It must not be injected as such.

Otherwise it will cause haemolysis and hypotonic electrolyte disorders.

<u>Please note:</u> The safety information of the additive provided by the respective manufacturer must be taken into account.

Interactions with other medicinal products and other forms of interaction

None known

Pregnancy and lactation

The use of WFI during pregnancy and lactation depends on the nature of the medicine that is added to the WFI.

Dosage

Dosage and duration of use depend on the instructions given for the medicinal product to be dissolved/diluted.

Method of administration

The method of administration depends on the instructions given for the medicinal product to be dissolved/diluted. The medicinal products should be reconstituted or diluted immediately before use. Not to be used as such for intravenous administration.

Overdose

Symptoms and treatment

Should B. Braun Water for Injections B.P have inadvertently been infused as such, any disorders of the fluid and electrolyte balance must be corrected according to the serum analytical values.

If massive haemolysis occurs, appropriate emergency treatment must be instituted immediately.

Undesirable effects

None known if used according to the instructions given for the medicinal product to be dissolved/diluted.

Note:

Patients should inform their doctor or pharmacist if they notice any side effect not mentioned in this leaflet.

Expiry date

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

Presentation

5 ml, 10 ml, 20 ml plastic ampoules. 50 ml, 100 ml, 250 ml, 500 ml, 1000 ml plastic containers. Not all presentation is available locally.

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Instructions for storage /use / handling

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

No special requirements for disposal.

The containers are for single use only. After use discard container and any remaining contents.

Only to be used if solution is clear, colourless and the container and its closure are undamaged.

Use the liquid immediately after opening of the

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Miniplasco







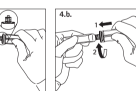


Miniplasco basic

















Product Registration Holder:

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Manufactured by:

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