

Mannitol Intravenous for Adults

Who can administer

May be administered by registered competent doctor or nurse/midwife

Important information

- Only the **10% solution** is now available for **ADULT** patients. (The 20% solution is stocked for use in paediatric patients)
- **Crystallisation** may occur - see under 'storage requirements' for further information
- See 'Monitoring requirements' below
- **Flush line with Glucose 5%** as there is a risk of precipitation with Sodium chloride 0.9% ^(ref 1)
- Administer via central line or large peripheral vein
- **Extravasation** causes inflammation and thrombophlebitis

Available preparations

Mannitol 10% 500mL infusion (50g in 500mL)

Reconstitution

Already in solution

Infusion fluids

Not required - product ready for infusion

Methods of intravenous administration

Slow intravenous injection (test dose for patients with marked oliguria or suspected inadequate renal function)

- **Test dose** over 3 to 5 minutes

Intermittent intravenous infusion (administer using an electronically controlled infusion device)

- See under 'dose' for details
- Administer via a giving set that incorporates a 15 micron **in-line filter** ^(ref 1) - check packaging of administration set for details)

Dose in adults

Volume in ml of Mannitol 10% solution required						
Required dose	15g	25g	50g	75g	100g	150g
10% solution	150ml	250ml	500ml	750ml	1000ml	1500ml

Reduction of intracranial pressure, cerebral volume and intraocular pressure

- Usual dose: 15 to 20mL/kg (10% infusion) administered over 30 to 60 minutes (some references suggest a range of 2.5 to 20mL/kg ^(ref 2))
- Dose may be repeated once or twice after 4 to 8 hours ^(ref 2)
- When used pre-operatively, the dose should be administered 60 to 90 minutes before surgery to obtain the maximum effect

Use in patients with oliguria or renal impairment

- A test dose (to assess renal function) of about 2mL per kg (10% infusion) should be administered over three to five minutes - this should produce a diuresis of at least 30 to 50ml/hour during the next two to three hours
- A second test dose may be given if there is inadequate response to the first test dose
- If the second test dose does not produce an adequate urine output, then need to reassess management

Promotion of elimination of renally excreted toxic substances in poisoning

- An initial loading dose of 250mL (10% infusion) may be given
- Adjust dose to maintain urinary output of at least 100mL/hour and positive fluid balance of 1 to 2 litres

Renal impairment

- Use with caution in severe renal impairment
- If the second test dose does not produce adequate urine output, reassess options

Monitoring

- Renal function, fluid balance, serum electrolytes, serum and urine osmolality
- Monitor infusion site for extravasation
- Assess cardiac function before and during treatment

Storage

- Store between 20 and 30°C ^(ref 1)
- Do not refrigerate or freeze ^(ref 1)
- Solutions may crystallise, especially if stored at low temperatures
- The administration set should contain a **15 micron in-line filter** - see under method of administration above

Crystals may be redissolved by warming before use as follows:

- Re-dissolve any crystals by warming to 37°C (Baxter products), or 60°C (Fresenius Kabi products) ^(ref 1)
- Use of dry heat (eg warming cabinet) is recommended - solutions should not be heated in water due to risk of contamination
- Gently agitate occasionally ^(ref 1)
- Microwave heating **should not be used**
- The product should be allowed to cool to 37°C before infusion

References

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1. Injectable medicines administration guide, Medusa Accessed online 31/03/2026
2. BNF- accessed online 31/03/2026

Therapeutic classification

Osmotic diuretic