

Who can administer

Administration RESTRICTED- see Appendix 1Â

Important information

- Unlicensed preparation
- Serum potassium levels should be corrected before and monitored throughout treatment with enoximone. Abnormal levels can predispose patients to arrhythmias
- See under 'Dose' for adjustments required in renal impairment'

Available preparations

Enoximone 100mg per 20ml ampoule (Perfan)

Reconstitution

Already in solution

- Draw up using a 5 micron filter needle
- Dilute further prior to administration

Infusion fluids

Sodium chloride 0.9% ONLY

Methods of intravenous administration

If a central venous access device is unavailable, administer via a large peripheral vein monitoring insertion site closely using a recognised phlebitis scoring tool ^(ref 1)

Slow intravenous injection

- Add required dose to equal volume of infusion fluid (to produce a 2.5mg/mL solution)Â
- Do not use more dilute solutions as crystal formation may occur

Intermittent intravenous infusion (using an electronically controlled infusion device)

- Add required dose to equal volume of infusion fluid (to produce a 2.5mg/mL solution)
- Administer as per 'Dose' overleaf

Continuous intravenous infusion (using an electronically controlled infusion device)

- Add 100mg (20mL) to 20mL infusion fluid ^(ref 1,2) i.e. final infusion contains 100mg in 40mL (2.5mg/mL solution)
- Do not use more dilute solutions as crystal formation may occur
- Rate adjusted according to response see 'Dose'

Â

Dose in adults

Congestive Heart failure

Initial therapy

- Slow intravenous injection
 - $\circ\,$ Give 0.5 to 1 mg /kg at a rate not greater than 12.5mg per minute
 - Further doses of 0.5mg/kg may be given every 30 minutes until a satisfactory response is achieved or a total initial dose of 3mg/kg is achieved
 - OR
- Intermittent infusion
 - Give at a rate of 90micrograms/kg/minute administered over 10 to 30 minutes until the required haemodynamic response is achieved

Maintenance dose

- Slow intravenous injection
 - The initial dose (up to a max of 3mg/kg) may be repeated every three to six hours as required adjusted according to the patient response
 - This should be administered at rate not greater than 12.5mg per minute
 - Normal maximum daily dose is 24mg/kg/day (i.e. 1,680mg/day in 70kg patient)
 - OR
- Continuous infusion
 - Give 5 20 micrograms/kg/minute following loading dose, according to haemodynamic response.
 - Normal maximum daily dose is 24mg/kg/day (i.e. 1,680mg/day in 70kg patient)

Renal impairment

• In patients with renal impairment, the dose or dosage frequency may need to be reduced

Monitoring

- Enoximone has a high pH and may cause venous irritation and tissue damage in cases of extravasation
- Monitor **potassium** levels: See under Important Information above
- Platelet count to be assessed before and during therapy.
- Severe **gastrointestinal symptoms** may occur; can be managed by reducing dose/temporarily interrupting the regime.
- Liver function tests; if clinically significant increases in hepatic enzymes therapy should be discontinued.
- Contains **ethanol** (10.4%); caution for use in those with a history of alcohol excess, pregnant or breastfeeding women and high-risk groups such as liver disease or epilepsy.
- Contains **propylene glycol** -monitor for adverse events e.g. hyperosmolality, lactic acidosis, renal dysfunction, cardiotoxicity, CNS disorders, respiratory depression, dyspnoea, liver dysfunction, haemolytic and haemoglobinuria.

Further information

 Occasionally, enoximone has produced 'furring of the lines'. In practice a dedicated line is recommended for its administration ^(ref 1)

Storage

Store below $25^{\circ}C$

Do not freeze.

References

Summary of Product Characteristics Perfan July 1st 2022 1: Medusa NHS guide - downloaded June 2023 2: Standard Medication Concentrations for Continuous Infusions in Adult Critical Care. Intensive Care Society December 2020 version 4.1.Â

Therapeutic classification

Phosphodiesterase enzyme inhibitor