

Potassium chloride Intravenous INFUSION for Adults

Who can administer

Commercially available bags containing not more than 40mmol per 500ml

- May be administered by registered competent doctor or nurse/midwife.

Infusions prepared at ward level using ampoules

- May be administered by registered competent doctor or nurse/midwife, PROVIDED the [guidelines below](#) (in Methods of Administration) have been adhered to

Higher concentrations: Administration RESTRICTED - see [Appendix 1](#)

Important information

- **Use pre-mixed infusion bags unless absolutely essential**
- Potassium chloride solutions can be **FATAL** if given inappropriately
 - Ampoules (concentrate) **MUST** be diluted before administration - see suitable dilutions below
 - Administration must be by **slow intravenous infusion through a pump**
- Any form of Potassium which contains a concentration **greater than 40mmol per litre**, is a **controlled drug** within Galway University Hospitals.
- Pain at the site of injection and phlebitis may occur during intravenous administration of solutions containing 40mmol or more potassium per litre ^(ref 1)
- For Y-site compatibility [see below](#)
- **For addition of potassium concentrate to infusion bags - see under [Methods of Administration](#) below**

Available preparations

Potassium chloride parenteral preparations

Fluid	Potassium content	Volume	Product	Comments
Sodium chloride 0.9%	20mmol	500ml	B1983	All generally available - many as stock on wards - available to order if not stock
Sodium chloride 0.9%	20mmol	1000ml	B1764	
Sodium chloride 0.9%	40mmol	1000ml	B1984	
Sodium chloride 0.9%, Glucose 5%	20mmol	500ml	B2486	Paediatric DKA policy
Sodium chloride 0.45%, Glucose 5%	20mmol	500mL	Fresenius Kabi 19-09-115	Perioperative management of some diabetics - see insulin prescription chart
Glucose 5%	40mmol	1000ml	Fresenius Kabi	All generally available - many as stock on wards - available to order if not stock
Glucose 5%	20mmol	1000ml	B1134	
Glucose 5%	20mmol	500ml	B1263	
Sodium chloride 0.9% (controlled drug) (Unlicensed)	40mmol	500ml	Fresenius Kabi 794764	Unlicensed Controlled drug in GUH
Sodium chloride 0.9% (controlled drug)	20mmol	100ml	G5028	Critical care areas only
Sodium chloride 0.9% (controlled drug)	40mmol	100ml	G5020	Critical care areas only
Potassium chloride 15% CONCENTRATE (for dilution and infusion) Controlled drug	20mmol	10ml	Â	Critical or complex care areas, ED, paediatrics, neonatal unit only.

Reconstitution

Already in solution

Infusion fluids

Ready made infusion bags as listed above

Use **Sodium Chloride 0.9% as fluid of choice** for initial replacement (unless contraindicated) as Glucose may cause a further decrease in plasma potassium levels ^(ref 1)

Methods of intravenous administration

Intravenous infusion (using an electronically controlled infusion device - i.e. pump)

NB: Pumps must never be removed while a potassium infusion is hanging - this includes when patients are being moved between units/wards

PERIPHERAL LINE	
Available as	<ul style="list-style-type: none"> • Standard pre-mixed infusion bag containing not more than 40mmol/L (preferred) • If fluid volume is an issue: use pre-mixed 40mmol in 500ml sodium chloride 0.9% and administer through a large vein • The MAXIMUM concentration that can be administered via peripheral line through a large vein is 40mmol/500mL
Administration	<ul style="list-style-type: none"> • Rate of administration should not normally exceed 10mmol per hour ^(ref 1,4) • Exceptionally, with senior decision maker, can give 20mmol per hour, but only with cardiac monitoring, and preferably through a large vein • Do not exceed 20mmol per hour • Monitor the patient for pain or phlebitis at the injection site
CENTRAL LINE	
Available as	<ul style="list-style-type: none"> • Standard pre-mixed infusion bag • Pre-mixed 20 or 40mmol in 100mL ^(unlicensed) • Pre-mixed 40mmol in 500ml sodium chloride 0.9% and administer through a large vein
Administration	<ul style="list-style-type: none"> • Rate of administration should not normally exceed 10mmol per hour, or exceptionally 20mmol/hour ^(ref 1) • In critical care areas, can give up to 40mmol per hour if absolutely necessary, but only with cardiac monitoring ^(ref 2) • Do not exceed 40mmol per hour ^(ref 2)

Addition of potassium concentrate to infusion bags

- GUH hospital policy requires that pre-mixed bags be used in preference
- If essential to prepare at ward level
 - Preparation must be done **jointly by a doctor and a nurse** in the clinic room
 - The second practitioner should always check for the correct product, dosage, dilution, mixing and labelling during the preparation of and again prior to intravenous administration
 - Both the Controlled Drug register, and the Additive label **must be signed** by the SAME doctor and nurse
 - UNUSED ampoules must immediately be returned to the CD press and signed back into the CD register by the SAME doctor and nurse
 - The addition of potassium concentrate to an existing lower concentrate potassium infusion bag may be considered (with over-labelling to reflect the TOTAL amount of mmol of potassium in the bag)
 - After adding potassium concentrate to an infusion bag, squeeze and invert bag a **MINIMUM** of ten times to avoid inadvertent administration of a potentially **fatal toxic bolus** ^(ref 1)
 - **Cannot exceed 40mmol per 500mL concentration for peripheral use**

Dose in adults

- Always prescribe in **mmol** and specify the **final volume** of infusion to avoid confusion ^(ref 1)
- Oral potassium supplements can be prescribed in conjunction with intravenous potassium ^(ref 1)
- Patients **at risk of hypokalaemia** should receive oral supplementation or maintenance potassium infusions as a means of restricting the necessity for 'rescue' high strength infusions.
- Doses can be highly variable (monitoring is essential)

The following options for administering potassium IV are listed in order of preference (four

options)

Option 1 (preferred)

- Using the premixed infusions available - up to 40mmol per litre
- Give at a usual rate of 10mmol per hour (up to 20mmol per hour **with cardiac monitoring, do not exceed 20mmol/hour**) ^(ref 1,4)

Option 2: (if fluid volume is an issue)

- Use pre-mixed 40mmol in 500ml sodium chloride 0.9%
- Administer via a **large peripheral vein**
- Give at a rate of 10mmol per hour (up to 20 mmol per hour **with cardiac monitoring**) ^(ref 1,4)

Option 3: (Critical care areas ONLY)

- If fluid volume is an issue, using the pre-mixed bag of 20mmol or 40mmol per 100mL ^(unlicensed)
- Administer via **central line only**
- Give at a rate of 10mmol per hour (up to 20 mmol per hour **with cardiac monitoring**) ^(ref 1,4)
- Expert sources advise higher concentrations of potassium chloride may be given in very severe depletion, but require specialist advice ^(ref 3)

Option 4 (Where no premixed bag is suitable)

- The addition of potassium concentrate to an existing lower concentrate potassium infusion bag may be considered (with over-labelling to reflect the TOTAL amount of mmol of potassium in the bag)
- Take careful note of maximum allowable concentrations for peripheral or central line use (see under Methods of administration)
- **Thorough mixing** of the bag after adding the potassium concentrate **is essential** (squeeze and invert bag at least ten times) ^(ref 1)
- See [guidelines above](#) under table for central/peripheral lines for guidance on how to add to bags

Monitoring

Continuous Cardiac Monitoring requirements ^(ref 1)

- Advised if the rate of infusion is greater than 10mmol potassium/hour, and **must be used** if the rate of infusion is 20mmol potassium/hour or greater
- Required if the potassium concentration being administered exceeds 80mmol per litre
- Required if the patient's serum potassium is less than or equal to **2.5mmol/L**
- Peaking of the T wave or other ECG changes associated with hyperkalemia indicate that the rate of potassium infusion is excessive and should be reduced

Site of infusion

- Monitor patient for **pain or phlebitis** which may occur at the site of infusion during peripheral administration of solutions containing potassium
- If pain occurs, either the infusion rate, or preferably, the concentration should be reduced ^(ref 2)

Storage

- **Controlled drug press** for any parenteral potassium with a concentration which exceeds 40mmol/litre of potassium
- Store below 25°C

References

1. [Best practice guidelines for the safe use of intravenous potassium in Irish Hospitals](#), October 2020 Irish Medication Safety Network
2. Uptodate- accessed online April 2026
- 3: BNF accessed online via MedicinesComplete April 2026
- 4: Injectable medicines guide. Medusa, downloaded April 2026
- 5: [GUH policy- potassium concentrate, supply and storage in GUH hospitals](#)CLN-PHAR/UCH-023

Therapeutic classification

Electrolyte