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Alert	High risk of hypo and hyperglycaemia necessitating close monitoring of blood sugar levels.
	Insulin binds to the plastic of giving sets. Flush the plastic tubing with 20 mL of prepared insulin solution
	into a receptacle prior to connecting to the infant. This is to saturate the binding.
	Insulin concentrations ≤ 0.05 units/mL are not reliably delivered even after preconditioning and flushing.
Indication	Treatment of hyperkalaemia:
	• Infants with serum potassium ( $K^+$ ) $\geq$ 7.0 mmol/L
	Infants with hyperkalaemia and abnormal ECG
	Management of severe cardiotoxicity or cardiac arrest due to hyperkalaemia
Action	Insulin and glucose activate cellular sodium-potassium ATPase resulting in a potassium shift into the
	intracellular space.
Drug type	Polypeptide hormone – lowers blood glucose and potassium levels.
Trade name	Actrapid (Novo Nordisk)
	Humulin R (Eli Lilly)
Presentation	Vial: 100 units/mL in a 10 mL vial.
	Penfill cartridge: 100 units/mL in 3mL penfill
Dose	Treatment of hyperkalaemia with insulin—glucose 25% infusion
	Starting dose: 0.1 unit/kg/hour.
	Dose range: 0.05 to 0.2 unit/kg/hour.
	Titrate infusion rate to serial serum potassium and blood glucose concentrations.
	Treatment of hyperkalaemia with insulin-only infusion
	Starting dose: 0.1 unit/kg/hour.
	Dose range: 0.05 to 0.2 unit/kg/hour.
	Titrate infusion rate to serial serum potassium and blood glucose concentrations.
	Must have adequate maintenance fluids to achieve a glucose: insulin ratio of at least 2.5g:1unit
	to prevent hypoglycaemia.
	Management of severe cardiotoxicity or cardiac arrest due to hyperkalaemia
	0.2 units/kg of insulin in glucose 50% IV over 15 to 30 minutes.
	(Use this preparation only if there is insufficient time to prepare insulin—glucose 25%
	infusion).
Dose adjustment	Therapeutic hypothermia: Limited data in neonates.
	ECMO: Limited data in neonates.
	Renal impairment: Limited data in neonates.
	Hepatic impairment: Limited data in neonates. Close monitoring of BGL advised due to lability of BGL.
Maximum dose	N/A
Total cumulative	N/A
dose	
Route	Intravenous
Preparation	INSULIN—GLUCOSE 25% INFUSION – Run via central line.
	Infusion strength Prescribed amount
	1 mL/kg/hour = 0.1 unit/kg/hour 5 units insulin and make up to 50 mL
	Draw up 0.5 mL (50 units of insulin) and add 9.5 mL sodium chloride 0.9%, glucose 5% or glucose 10% to
	make a final volume of 10 mL with a concentration of 5 units/mL.
	<b>FURTHER DILUTE</b> : Draw up 1 mL (5 units of insulin) of solution and dilute with glucose 25% [25 mL glucose
	50% plus 24 mL water for injection] to make a final volume of 50 mL with a concentration/dose rate of <b>1</b>
	mL/kg/hour = 0.1 units/kg/hour.
	INSULIN ONLY INFUSION – Can be infused peripherally.
	Must have adequate maintenance fluids to achieve a glucose: insulin ratio of at least 2.5g:1unit to prevent
	i viust nave adequate maintenance nuius to achieve a giucose: insulin ratio of at least 2.5g:10ht to prevent
	hypoglycaemia.  Infusion strength Prescribed amount

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	1 mL/kg/hour = 0.2 unit/kg/hour	10 units insulin and make up to 50 mL	
		d add 9.5 mL sodium chloride 0.9%, glucose 5% or glucose 3	 10% to
	make a final volume of 10 mL with a co		1070 10
		nits of insulin) of solution and dilute with glucose 5%, glucos	se 10% or
		olume of 50 mL with a concentration/dose rate of <b>1 mL/kg</b>	
	0.2 units/kg/hour.	,	
	Cardiac arrest due to hyperkala	<u>emia</u>	
	Infusion strength	Prescribed amount	
	1 mL/kg/hour = 0.2 units/kg/hour	10 units insulin and make up to 50 mL	
		d make up to 50mL with glucose 50% (this contains 25g of g over 15 to 30 minutes. Glucose:insulin ratio = 2.5g:1unit.	lucose).
Administration	Intravenous:		
		ravenous bags, syringes and tubing which reduces the delive	
		astic, flush 20 mL of prepared insulin solution through pla	
		atient. Insulin concentrations ≤ 0.05 units/mL are not reliak	bly
	delivered even after preconditioning a	nd flushing [2].	
	Infuse with maintenance fluids.		
	Do not include in maintenance fluid re	•	
Manitarina	Insulin binds to the filter. <b>Do not filter</b>		
Monitoring		ed to detect either hypo/hyperglycaemia.	
		inutes for the first hour, every 30 minutes for the second h frequency of monitoring during weaning.	our and
		0–60 minutes of commencing glucose/insulin infusion. Seru	Im
		t to monitor response to treatment and avoid hypokalaemia	
Contraindications	Hypersensitivity to human insulin or ar		u.
contrainalcations	During episodes of hypoglycaemia.		
Precautions		sensitivity, hypoglycaemia, hyperglycaemia, and hypokalaer	mia.
	Use with caution in cardiac disease, he		
Drug interactions		irements: Octreotide, beta-adrenergic blocking agents, angi	iotensin
-	converting enzyme inhibitors, salicylates, anabolic steroids, alpha-adrenergic blocking agents, quinine,		
	quinidine, and sulfonamides.		
	The following may increase insulin req	uirements: Thiazides, furosemide, ethacrynic acid, glucocor	ticoids,
	thyroid hormones, sympathomimetics	-	
	Sympathomimetics have a potassium l		
Adverse	-	vith a high rate of hyperglycaemia and hypoglycaemia durin	g
reactions	infusion and hypoglycaemia during weaning (insulin has a longer half-life than glucose).		
	Hypokalaemia if infusion continued.		
<u> </u>	Hypertonic solution – potential for ext	ravasation.	
Overdose			
Compatibility	Fluids: Glucose 5%, glucose 10%, gluco		
		arone (variable), anidulafungin, ascorbic acid, asparaginase,	
		nzylpenicillin, bivalirudin, bleomycin, bumetanide, buprenc	-
	· · · · =	iconate, caspofungin, cefamandole, cefazolin, cefepime, cef	
		azobactam, ceftizoxime, ceftriaxone, cefuroxime, chloramp	
		alamin, cyclophosphamide, dexamethasone, dexmedetomic	
		at, epirubicin, epoetin alfa, erythromycin lactobionate, esm	
		azole, folic acid (as sodium salt), foscarnet, fosfomycin, fosp	-
		nisetron, heparin sodium, hydrocortisone, hydromorphone, indomethacin, isovuconazonium sulfate, lidocaine, linezolic	
	indproten tysine, intipenent-chastaliti,	indometriacin, isovaconazoniam sunate, naocame, imezono	',

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	<ul> <li>lorazepam, mannitol, magnesium sulfate, meropenem, methadone, methylprednisolone, metoclopramide, metoprolol, metronidazole, mixifloxacin hydrochloride, milrinone, naloxone, nitroglycerin, nitroprusside sodium, octreotide, oxacillin, palonosetron, pamidronate, pancuronium, pantoprazole (variable), paracetamol, pentoxyphylline, phenobarbital, phytomenadione, piperacillin, potassium acetate, potassium chloride; procainamide hydrochloride, promethazine hydrochloride, propofol, pyridoxine, remifentanil, sodium bicarbonate, sodium nitroprusside, streptokinase, sufentanil, tacrolimus, terbutaline, theophylline, thiamine hydrochloride, ticarcillin disodium, ticarcillin disodium-clavulanate potassium, tigecycline, urokinase, vancomycin, vecuronium, verapamil, voriconazole and zoledronic acid</li> <li>In syringe: Insulin NPH.</li> </ul>
Incompatibility	Y-site: Alprostadil, cefoperazone, cefoxitin, chlorpromazine, dantrolene sodium, diazepam, diazoxide,
	dobutamine, famotidine (variable), gentamicin (variable), glycopyrrolate, hydralazine (variable), isoprenaline, ketamine, labetalol, metaraminol (variable), micafungin, noradrenaline (norepinephrine)(variable), ondansetron (variable), phentolamine, phenylephrine, phenytoin, piperacillin- tazobactam, polymyxin, propranolol, protamine, rocuronium, sulfamethoxazole-trimethoprim, tobramycin, vasopressin (variable)
Stability	Prepared solutions are stable at room temperature (< 25°C) for 24 hours.
	A 20 mL insulin priming solution at a concentration of 0.1 units per mL was found to deliver 80% of the expected insulin (1). A 20 mL insulin priming solution with additional preconditioning for 1 hour at a concentration of 0.05 units per mL was found to deliver 26.5% of the expected insulin (2).
Storage	Store human insulin preparations between 2 and 8°C.
otor upc	Do not freeze. Human insulin preparations which have been frozen must not be used.
	Protect from excessive heat and light. Should appear clear and colourless.
	While it is suggested that insulin vials can be kept for 28 days after the first use, ANMF consensus
	recommendation is to avoid this practice because of the risk of microbial contamination and increased
	susceptibility of neonates to sepsis.
Excipients	Actrapid: glycerol, metacresol, zinc chloride, water for injection, hydrochloric acid, sodium hydroxide
Creation	Humulin R: glycerol, hydrochloric acid, metacresol, sodium hydroxide, water for injection
Special	Recommend administer insulin/glucose in same line as intravenous fluids. Recommend intravenous fluids and/or an additional glucose 25% syringe placed proximally for rapid
comments	treatment of hypoglycaemia if needed.
	Do not include insulin glucose in the total daily fluid intake.
	Frequent blood glucose and potassium measurements, especially after commencement and during
	weaning of infusion are needed for titration and safety
Evidence	Efficacy
	<ul> <li>Treatment of hyperkalaemia: A systematic review (3) of interventions for neonatal hyperkalaemia found 2 studies (4, 5) comparing insulin/glucose infusion versus rectal cation-resin. Meta-analysis of 2 studies (52 infants) found no difference in cardiac arrhythmias (RR 0.29; 95% CI 0.05, 1.65); or all-cause mortality [RR 0.18; 0.03, 1.15]. Malone 1991, using an insulin infusion 0.05 to 0.2 units/kg/hour in albumin 5%, reported reduced treatment failure (rise in K+ concentration &gt; 0.5 mmol/L or K+ &gt; 7 mmol/L) of borderline statistical significance (RR 0.07; 0.00 to 1.01; RD -1.00; -1.28 to -0.72) compared to resin (5). Hu 1999, using a glucose/insulin infusion with glucose 10−15 g:insulin 1 unit, reported a reduction in duration of hyperkalaemia (MD -12.20 hours; -20.95, -3.45); no difference in peak serum K+ (MD -0.10 mmol/L; -0.57, 0.37); a reduction in IVH (RR 0.3; 0.10, 0.93) and IVH grades ≥ 2 (RR 0.3; 0.10, 0.93) compared to resin; and no infant with hypoglycaemia in either group (4). No study compared insulin-glucose with a beta-agonist. Conclusion: The combination of insulin and glucose is preferred over treatment with rectal cation-resin for hyperkalaemia in preterm infants (3). (LOE I GOR C)</li> <li>Glucose:insulin ratio: It is recommended to neutralise insulin in the glucose-insulin infusion for</li> </ul>
	hyperkalaemia by using safe glucose:insulin ratio to prevent hypoglycemia. Several ratios ranging from 2.5:1 to 10:1 have been reported in literature (6,7). To balance the risk of hyper or hypoglycemia, a

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	historical control study compared infusions with lower glucose: insulin ratio 3.3g:1 unit (glucose 20%) versus a higher glucose:insulin ratio 5 g:1 unit (glucose 30%) for treatment of hyperkalaemia in neonates. This study reported reduced rates of moderate hyperglycaemia [77% to 21.7% (p = 0.001)] with a single infant in the lower arm having hypoglycaemia (8). (LOE III-3, GOR C).
	<ul> <li>Management of severe cardiotoxicity or cardiac arrest due to hyperkalaemia: The Pediatric Advanced Life Support guidelines (9), Advanced Cardiac Life Support guidelines (10) and a simulation trial of medication preparation and delivery (11) support the following sequence of medications to treat hyperkalaemia during paediatric cardiac: First, calcium; second, sodium bicarbonate; and third, insulin with glucose. Recommended dose [adult guideline]: Glucose plus insulin: mix 25 g (50 mL of glucose 50%) glucose and 10 units regular insulin and give IV over 15 to 30 minutes. Glucose:insulin ratio = 2.5 g:1 unit.</li> <li>Pharmacokinetics</li> <li>Following IV administration, the observed half-life of insulin ranges from 5 to 15 minutes (12).</li> </ul>
Practice points	
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	<ol> <li>American Heart Association. Web-based Integrated Guidelines for Cardiopulmonary Resuscitation and Emergency Cardiovascular Care – Part 12: Pediatric Advanced Life Support. ECCguidelines.heart.org.</li> <li>American Heart Association. Web-based Integrated Guidelines for Cardiopulmonary Resuscitation and</li> </ol>
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