### **Newborn Use Only**

2025

	High risk medication in A PINCH Medicines list under New South Wales Clinical Excellence Commission.  Different brands of insulin are not bioequivalent. Do not substitute between brands.[13]  Actrapid is the ANMF group's recommended short-acting insulin for IV infusion in neonates.								
	International units are hereafter referred to as "units".								
	High risk of hypoglycaemia.								
	Insulin binds to the plastic of giving	· · · · · · · · · · · · · · · · · · ·	=	pared insulin solution					
	into a receptacle prior to connectir Insulin concentrations ≤ 0.05 Unit/r			ditioning and flushing					
Indication	Treatment of persistent hyperglyca	•	erea even arter precon	didoning and nashing.					
	[For treatment of hyperkalaemia, se		nia].						
Action	Insulin is a polypeptide hormone t			late uptake, utilisation					
	and storage of glucose resulting in a								
	in the form of glycogen and facilit			-					
	lipolysis, proteolysis and gluconeog into fat.	genesis, ennances prote	in synthesis and conve	rsion of excess glucose					
Drug type	Polypeptide hormone – lowers bloc	od glucose.							
Trade name	Actrapid [Novo Nordisk]								
Presentation	100 units/mL in a 10 mL vial and 3 i	mL Penfill.							
Dose	Starting dose: 0.05 unit/kg/hour.		VY						
	Dose range: 0.01 to 0.1 unit/kg/hou								
D	Titrate in small increments to blood								
Dose adjustment	Therapeutic hypothermia: Limited e euglycemia [3].	evidence in neonates. H	igner dose may be requ	ired to maintain					
	ECMO: Data limited in preterm neo	nates to make recomme	endation.						
	Renal impairment: Limited data in r			re renal failure.					
	Hepatic impairment: Limited data i	n neonates. Close monit	oring of BGL advised du	e to lability of BGL [4].					
Maximum dose		$\rightarrow$							
Total cumulative									
dose Route	IV								
Preparation		of giving sets. Flush the	nlastic tuhing with 20 n	nt of prepared insulin					
i i c pai acion	NOTE: 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.								
-	solution into a receptacle prior to	- 7	. This is to saturate the						
-	solution into a receptacle prior to	- 7	. This is to saturate the						
-	solution into a receptacle prior to NOTE: Refer to Appendix for tables	connecting to the infant							
-	NOTE: Refer to Appendix for tables	connecting to the infant s to assist with concentr	ration selection.	e binding.					
	NOTE: Refer to Appendix for tables Weight suggestions for infusion co	s to assist with concentroncentrations below are	ration selection. e a guide only. Clinicians	e binding.  s may choose infusion					
	NOTE: Refer to Appendix for tables Weight suggestions for infusion co concentration different to the sug	s to assist with concentroncentrations below are	ration selection. e a guide only. Clinicians	e binding.  s may choose infusion					
	NOTE: Refer to Appendix for tables Weight suggestions for infusion co	s to assist with concentroncentrations below are	ration selection. e a guide only. Clinicians	e binding.  s may choose infusion					
	NOTE: Refer to Appendix for tables Weight suggestions for infusion co concentration different to the sug volumes.  Infant weight	s to assist with concentroncentrations below are	ration selection. e a guide only. Clinicians	e binding.  s may choose infusion					
	NOTE: Refer to Appendix for tables Weight suggestions for infusion co concentration different to the sug volumes.  Infant weight Suggested Insulin	to assist with concentrations below are gested based on expect	ration selection.  e a guide only. Clinicians ted dose and the corres $1 \text{ to } \leq 3 \text{ kg}$	e binding.  s may choose infusion sponding 24-hour fluid  ≥3 kg					
	NOTE: Refer to Appendix for tables  Weight suggestions for infusion co concentration different to the sug volumes.  Infant weight Suggested Insulin concentration	to assist with concentrations below are gested based on expect \$\$ <1 kg \$\$ 0.05 unit/mL \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$	ration selection.  e a guide only. Clinicians ted dose and the corres  1 to ≤3 kg  0.2 unit/mL	e binding.  s may choose infusion sponding 24-hour fluid  ≥3 kg  0.8 unit/mL					
	NOTE: Refer to Appendix for tables Weight suggestions for infusion co concentration different to the sug volumes.  Infant weight Suggested Insulin	to assist with concentrations below are gested based on expect	ration selection.  e a guide only. Clinicians ted dose and the corres $1 \text{ to } \leq 3 \text{ kg}$	e binding.  s may choose infusion sponding 24-hour fluid  ≥3 kg					
	NOTE: Refer to Appendix for tables  Weight suggestions for infusion co- concentration different to the suggestions.  Infant weight Suggested Insulin concentration 0.01 unit/kg/hour is equal to	to assist with concentrations below are gested based on expect \$\$ <1 kg \$\$ 0.05 unit/mL \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$	ration selection.  e a guide only. Clinicians ted dose and the corres  1 to ≤3 kg  0.2 unit/mL	e binding.  s may choose infusion sponding 24-hour fluid  ≥3 kg  0.8 unit/mL					
	NOTE: Refer to Appendix for tables  Weight suggestions for infusion co- concentration different to the sug- volumes.  Infant weight Suggested Insulin concentration 0.01 unit/kg/hour is equal to  20 mL Syringe	to assist with concentrations below are gested based on expect \$\$ <1 kg \$\$ 0.05 unit/mL \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$	ration selection.  e a guide only. Clinicians ted dose and the corres  1 to ≤3 kg  0.2 unit/mL	e binding.  s may choose infusion sponding 24-hour fluid  ≥3 kg  0.8 unit/mL					
	Weight suggestions for infusion coconcentration different to the sugvolumes.  Infant weight Suggested Insulin concentration 0.01 unit/kg/hour is equal to  20 mL Syringe It is a 2 step dilution.	connecting to the infant to assist with concentrations below are gested based on expect  <1 kg  0.05 unit/mL  0.2 mL/kg/hour	ation selection.  a guide only. Clinicians ted dose and the corres  1 to ≤3 kg  0.2 unit/mL  0.05 mL/kg/hour	e binding.  s may choose infusion sponding 24-hour fluid  ≥3 kg  0.8 unit/mL  0.0125 mL/kg/hour					
	NOTE: Refer to Appendix for tables  Weight suggestions for infusion co- concentration different to the sug- volumes.  Infant weight Suggested Insulin concentration 0.01 unit/kg/hour is equal to  20 mL Syringe	connecting to the infant to assist with concentrations below are gested based on expect  <1 kg  0.05 unit/mL  0.2 mL/kg/hour	ation selection.  a guide only. Clinicians ted dose and the corres  1 to ≤3 kg  0.2 unit/mL  0.05 mL/kg/hour	e binding.  s may choose infusion sponding 24-hour fluid  ≥3 kg  0.8 unit/mL  0.0125 mL/kg/hour					
	Weight suggestions for infusion coconcentration different to the sugrestions.  Infant weight Suggested Insulin concentration 0.01 unit/kg/hour is equal to  20 mL Syringe It is a 2 step dilution. Step 1. Draw up insulin and add con	connecting to the infant to assist with concentrations below are gested based on expect  <1 kg  0.05 unit/mL  0.2 mL/kg/hour  mpatible fluid* to make  0.05 unit/mL	a guide only. Clinicians ted dose and the correst to ≤3 kg  0.2 unit/mL  0.05 mL/kg/hour  a diluted solution as pe  0.2 unit/mL	e binding.  s may choose infusion sponding 24-hour fluid  ≥3 kg  0.8 unit/mL  0.0125 mL/kg/hour  r table below:  0.8 unit/mL					
	Weight suggestions for infusion co- concentration different to the sug- volumes.  Infant weight Suggested Insulin concentration 0.01 unit/kg/hour is equal to  20 mL Syringe It is a 2 step dilution. Step 1. Draw up insulin and add con- Insulin concentration Volume of Insulin (100 units/mL)	connecting to the infant to assist with concentrations below are gested based on expect  <1 kg  0.05 unit/mL  0.2 mL/kg/hour  mpatible fluid* to make  0.05 unit/mL  0.2 mL (20 units)	a guide only. Clinicians ted dose and the correst to ≤3 kg  0.2 unit/mL  0.05 mL/kg/hour  a diluted solution as pe  0.2 unit/mL  0.2 mL (20 units)	e binding.  s may choose infusion sponding 24-hour fluid  ≥3 kg  0.8 unit/mL  0.0125 mL/kg/hour  r table below:  0.8 unit/mL  0.2 mL (20 units)					
	Weight suggestions for infusion co- concentration different to the sug- volumes.  Infant weight Suggested Insulin concentration 0.01 unit/kg/hour is equal to  20 mL Syringe It is a 2 step dilution. Step 1. Draw up insulin and add con- Insulin concentration Volume of Insulin	connecting to the infant to assist with concentrations below are gested based on expect  <1 kg  0.05 unit/mL  0.2 mL/kg/hour  mpatible fluid* to make  0.05 unit/mL  0.2 mL (20 units)  9.8 mL	a guide only. Clinicians ted dose and the correst to ≤3 kg  1 to ≤3 kg  0.2 unit/mL  0.05 mL/kg/hour  a diluted solution as pe  0.2 unit/mL  0.2 mL (20 units)  9.8 mL	e binding.  s may choose infusion sponding 24-hour fluid  ≥3 kg  0.8 unit/mL  0.0125 mL/kg/hour  r table below:  0.8 unit/mL  0.2 mL (20 units)  9.8 mL					
	Weight suggestions for infusion co- concentration different to the sug- volumes.  Infant weight Suggested Insulin concentration 0.01 unit/kg/hour is equal to  20 mL Syringe It is a 2 step dilution. Step 1. Draw up insulin and add con- Insulin concentration Volume of Insulin (100 units/mL)	connecting to the infant to assist with concentrations below are gested based on expect  <1 kg  0.05 unit/mL  0.2 mL/kg/hour  mpatible fluid* to make  0.05 unit/mL  0.2 mL (20 units)	a guide only. Clinicians ted dose and the correst to ≤3 kg  0.2 unit/mL  0.05 mL/kg/hour  a diluted solution as pe  0.2 unit/mL  0.2 mL (20 units)	e binding.  s may choose infusion sponding 24-hour fluid  ≥3 kg  0.8 unit/mL  0.0125 mL/kg/hour  r table below:  0.8 unit/mL  0.2 mL (20 units)					

Step 2: Draw up diluted insulin and add compatible fluid\* as per table below:

### **Newborn Use Only**

	B	Callete Louis Louis	and the second s	de alte autobre e entre				
	Prepare two separate 20 mL syringe							
	Insulin concentration	0.05 unit/mL	0.2 unit/mL	0.8 unit/mL				
	Volume of diluted insulin from step 1	0.5 mL (1 unit)	2 mL (4 unit)	8 mL (16 unit)				
	Volume of compatible fluid*	19.5 mL	18 mL	12 mL				
	Total volume	20 mL	20 mL	20 mL				
	*Compatible fluid: glucose 5%, gluc	ose 10% or sodium chlo	ride 0.9%					
	50 mL Syringe							
	It is a 2 step dilution							
	Step 1: Draw up insulin and add co	mnatible fluid* to make	a diluted solution as no	r table below:				
	Insulin concentration	0.05 unit/mL	0.2 unit/mL	0.8 unit/mL				
	Volume of Insulin	0.05 unit/inc	0.2 unit/inc	0.0 unity iii				
	(100 units/mL)	0.2 mL (20 units)	0.2 mL (20 units)	0.5 mL (50 units)				
	Volume of compatible fluid*	9.8 mL	9.8 mL	24.5 mL				
	Total volume	10 mL solution	10 mL solution	25 mL solution				
	Total volume	(2 units/mL)	(2 units/mL)	(2 units/mL)				
	Step 2: Draw up diluted insulin and	add compatible fluid* a	s per table below.	*				
	Insulin concentration	0.05 unit/mL	0.2 unit/mL	0.8 unit/mL				
	Volume of diluted insulin from							
	step 1	1.25 mL (2.5 unit)	5 mL (10 unit)	20 mL (40 unit)				
	Volume of compatible fluid*	48.75 mL	45 mL	30 mL				
	Total volume	50 mL	50 mL	50 mL				
	Use 20 mL of this solution to flush	the plastic tubing						
	*Compatible fluid: glucose 5%, gluc	ose 10% or sodium chlo	ride 0.9%					
Administration	Intravenous: Insulin binds to the plastic of giving sets. Flush the plastic tubing with 20 mL of prepared							
	insulin solution into a receptacle p		e infant. This is to satur	ate the binding.				
	Do not filter infusion. Insulin also bi		ata a ta sulta ta forta a afra	and a City a				
	Can be infused with maintenance fl		ning insulin infusion afte	er the filter.				
Monitoring	Do not bolus other drugs through the Blood glucose level (BGL)	nis line.						
Monitoring	After Initiation of infusion:	20 minutes 2 hours has	end on the infant's risk n	rofila until stabilisad				
	On maintenance: 4–6 hour		eu on the illiant's risk p	Tome until Stabiliseu.				
	After cessation of infusion:							
	Alteration of infusion: With							
	Serum potassium concentration.							
Contraindications	Hypersensitivity to regular insulin o	r any of its components						
	During episodes of hypoglycaemia.	, ,						
Precautions	Hypoglycaemia is a common advers	se effect. Blood glucose	must be monitored clos	ely to detect				
	hypoglycaemia.							
	Do not adjust the rate of the mainte	enance solution or other	r infusions when insulin	is commenced or the				
	insulin infusion rate is altered. For e	example, if insulin is com	nmenced or the rate of t	he insulin infusion is				
	increased, do not turn down the ma		•					
	The amount of glucose being delive			sulin is commenced or				
	dose is increased, possibly causing		-					
	If ceasing insulin or changing the str		nove and replace the pr	evious line and T-piece				
	to avoid flushing through insulin re		atal ta and ta lead to a select of					
Dunca interpreta	Administer IV bolus medication via							
Drug interactions	The following may reduce insulin re							
	converting enzyme inhibitors, salicy	riales, anapolic steroids,	aipna-aurenergic block	ing agents, quinine,				
	quinidine and sulfonamides.	requirements. This sides	furocomido othornis	acid alucocorticoide				
	The following may increase insulin requirements: Thiazides, furosemide, ethacrynic acid, glucocorticoids,							

 $thy roid\ hormones,\ sympathomimetics,\ octreotide,\ growth\ hormone,\ and\ diazoxide.$ 

### **Newborn Use Only**

2025

	Beta blocking agents may mask the symptoms of hypoglycaemia and delay recovery from hypoglycaemia.
	Hypoglycaemia in the presence of concomitant use of a beta-adrenergic blocking agent may precipitate a
	hypertensive crisis.
Adverse reactions	Hypoglycaemia; hypokalaemia; and hyponatraemia.
	Urticaria and anaphylaxis (extremely rare).
	Insulin resistance may develop resulting in a larger dose requirement.
Compatibility	Fluids: sodium chloride 0.9%, glucose 5%, glucose 10%, glucose 50%.
	PN at Y site:
	Y-site:[12,13] Acetaminophen, aciclovir, alfentanil, aminophylline, amphoteiricin B lipid complex,
	anidulafungin, atenolol, atropine, azathioprine, aztreonam, caffeine citrate, calcium gluconate,
	caspofungin, cefazolin, cefepime, cefotaxime, ceftazidime, ceftriaxone, cefuroxime, chloramphenicol,
	clindamycin, cloxacillin, dexamethasone, dexmedetomidine, enalaprilat, epoetin alfa, erythromycin
	lactobionate, fentanyl, fluconazole, folic acid, foscarnet, fosfomycin, fosphenytoin, furosemide,
	ganciclovir, hydrocortisone, ibuprofen, imipenem-cilastatin, indometacin, lidocaine, linezolid, magnesium
	sulfate, Meropenem, methadone, methylprednisolone, metoclopramide, metoprolol, metronidazole,
	milrinone, naloxone, nitroglycerin, nitroprusside, octreotide, pamidronate, pancuronium, penicillin G,
	pentobarbital, pentoxifylline, phenobarbital, potassium acetate, potassium chloride, propofol, pyridoxine, remifentanil, sildenafil, sodium bicarbonate, sodium nitroprusside, streptokinase, tacrolimus,
	thiamine, ticarcillin –clavulanate, urokinase, vancomycin, vecuronium, verapamil, voriconazole.
	Variable compatibility:[12] amikacin, amiodarone, amphotericin B conventional, ampicillin, cyclosporine,
	digoxin, dobutamine, dopamine, epinephrine, furosemide, gentamicin, heparin, hydralazine, midazolam,
	morphine sulfate, multiple vitamin injection, norepinephrine (refer to Micromedex), ondansetron,
	pantoprazole, tobramycin, vasopressin.
Incompatibility	Y-site administration:[12,13] Adrenaline (epinephrine), alprostadil, amikacin, amphotericin B, ampicillin,
	calcium chloride, cefoxitin, diazepam, diazoxide, digoxin, dobutamine, dopamine, epinephrine,
	furosemide-undiluted, gentamicin, glycopyrrolate, hydralazine, ketamine, labetalol, morphine,
	phenytoin, piperacillin -tazobactam, propranolol, protamine, rocuronium, sulfamethoxazole-
	trimethoprim
Stability	Actrapid: Prepared solutions are stable at room temperature (< 25°C) for 24 hours. (extrapolated from
	Insulin Human Regular) [12]
Storage	Store human insulin between 2 and 8°C. Do not freeze.
	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	Glycerol, metacresol, zinc chloride, water for injections. Hydrochloric acid and sodium hydroxide are used
	to adjust the pH. Contains less than 1 mmol sodium (23 mg) per dose, i.e. is essentially 'sodium-free'.
Special comments	Insulin is adsorbed to the plastic of intravenous bags, syringes, and tubing which reduces the delivery of
	insulin [5-7].
	Twenty mL of insulin priming solution at concentrations of 0.1 unit/mL and 0.05 unit/mL were found to
	deliver 80% and 26.5% of the expected insulin. Insulin concentrations ≤ 0.05 unit/mL are not reliably
	delivered even after preconditioning and flushing [5, 6].
Evidence	Efficacy
	Treatment of hyperglycaemia in very low birth weight infants: Systematic review [2] of trials of insulin infusion for treatment of page 21 hyperglycaemia found that use of an insulin infusion
	insulin infusion for treatment of neonatal hyperglycaemia found that use of an insulin infusion obviates the need to decrease the concentration of glucose prescribed and optimised the
	utilisation of calories by the infant resulting in significant increases in non-protein energy intake,
	glucose intake and short-term weight gain. However, insulin infusion had no significant effect on
	death, severe intraventricular haemorrhage, retinopathy of prematurity, bacterial sepsis, fungal
	sepsis or necrotising enterocolitis; effects on other major morbidities were not assessed. These
	trials did not report an excess of hypoglycaemia, possibly due to the more liberal target BSLs:
	4.4–9.9 mmol/L [8] and 5.5–9.9 mmol/L [9]. Conclusion: Evidence
	from randomised trials in hyperglycaemic VLBW neonates is insufficient to determine the effects
	of treatment on death or major morbidities. [2] [LOE I GOR D]
	Prevention of neonatal hyperglycaemia in very low birth weight infants: Systematic review [10]
	of trials of early insulin infusion for prevention of neonatal hyperglycaemia found that use of an

### **Newborn Use Only**

2025

	insulin infusion reduced hyperglycaemia but increased death before 28 days and increased the
	risk of hypoglycaemia. The reduction in hyperglycaemia was not accompanied by significant
	effects on major morbidities; effects on neurodevelopment are awaited. The evidence does not
	support the routine use of insulin infusions to prevent hyperglycaemia in VLBW neonates. [10][LOE I GOR
	B]
	Tight glycaemic control with insulin in hyperglycaemic very low birth weight infants: RCT in infants born
	at < 30 weeks' gestation or < 1500 g with hyperglycaemia (2 consecutive BGL > 8.5 mmol/L 4 hours apart)
	randomly assigned to tight glycaemic control with insulin (target BGL 4–6 mmol/L) or restrictive
	guidelines for starting insulin (target BGL 8–10 mmol/L). Infants in the tight group had a lesser lower leg
	growth rate (P < 0.05), but greater head circumference growth (P < 0.0005) and greater weight gain (P <
	0.001) to 36 weeks' postmenstrual age than control infants. Tight group infants had lower daily BGL and
	greater incidence of hypoglycaemia (BGL < 2.6 mmol/L) (25/43 vs 12/45; P < 0.01) than controls. There
	were no significant differences in nutritional intake or in the incidences of mortality or morbidity. The
	balance of risks and benefits of insulin treatment in hyperglycaemic pre-term neonates remains
	uncertain. [1] [LOE II GOR D].
	Guidelines: ESPGHAN 2005 recommended the use of insulin should be restricted to conditions where
	reasonable changes in glucose infusion rate do not control marked hyperglycaemia. [11] Although this
	recommendation is now out of date, current evidence is consistent with this recommendation.
	Pharmacokinetics
	Following IV administration, the observed half-life of insulin ranges from 5 to 15 minutes.[12]
Practice points	
References	1. Alsweiler JM, Harding JE, Bloomfield FH. Tight glycemic control with insulin in hyperglycemic
	preterm babies: a randomized controlled trial. Pediatrics. 2012;129:639-47.
	2. Bottino M, Cowett RM, Sinclair JC. Interventions for treatment of neonatal hyperglycemia in
	very low birth weight infants. Cochrane Database Syst Rev. 2011:CD007453.
	3. Cueni-Villoz N, Devigili A, et al. Increased blood glucose variability during therapeutic hypothermia and
	outcome after cardiac arrest. Crit Care Med. 2011 Oct; 39(10):2225-31.
	4. Scheen AJ. Pharmacokinetic and toxicological considerations for the treatment of diabetes in patients
	with liver disease. Expert Opin Drug Metab Toxicol. 2014; 10:839-857.
	5. Hewson M, Nawadra V, Oliver J, Odgers C, Plummer J, Simmer K. Insulin infusions in the
	neonatal unit: delivery variation due to adsorption. J Paediatr Child Health. 2000; 36:216-20.
	6. Thompson CD, Vital-Carona J, Faustino EV. The effect of tubing dwell time on insulin
	adsorption during intravenous insulin infusions. Diabetes Technol Ther. 2012;14:912-6.
	7. Simeon PS, Geffner ME, Levin SR, et al. Continuous insulin infusions in neonates: pharmacologic
	availability of insulin in intravenous solutions. Journal of Pediatrics. 1994; 124:818-20.
	8. Collins JW, Jr., Hoppe M, Brown K, Edidin DV, Padbury J, Ogata ES. A controlled trial of insulin infusion
	and parenteral nutrition in extremely low birth weight infants with glucose intolerance. J Pediatr. 1991;
	118:921-7.
	9. Meetze W, Bowsher R, Compton J, Moorehead H. Hyperglycemia in extremely-low-birthweight
	infants. Biol Neonate. 1998;74:214-21.
	10. Sinclair JC, Bottino M, Cowett RM. Interventions for prevention of neonatal hyperglycemia in very low
	birth weight infants. Cochrane Database Syst Rev. 2011:CD007615.
	11. Koletzko B, Goulet O, Hunt J, Krohn K, Shamir. Guidelines on Paediatric Parenteral Nutrition of the
	European Society of Paediatric Gastroenterology, Hepatology and Nutrition (ESPGHAN) and the European
	Society for Clinical Nutrition and Metabolism (ESPEN), Supported by the European Society of Paediatric
	Research (ESPR). J Pediatr Gastroenterol Nutr. 2005; 41 Suppl 2:S1-87.
	12. Micromedex. Insulin Human Regular. Accessed on 26 Sep 2025.
	13. Australian Injectable Drugs Handbook, 8 <sup>th</sup> Edition. Accessed on 28 October 2020.
	https://aidh.hcn.com.au/browse/i/insulin for subcutaneous or iv use
	14. Human Insulin (rys). Product Information. Accessed on 28 October 2020.
	15. Humulin Preparations. Product Information. Accessed on 28 October 2020.
Appendix	Infusion tables to assist concentration selection

Table 1: Infusion rates when using insulin concentration 0.05 unit/mL (Suggested weight <1kg)

Rate (mL/hr)	0.1	0.2	0.3	0.4	0.5	0.6	0.7	0.8	0.9	1
Weight (kg)		Approximate unit/kg/hour								
0.5	0.01	0.02	0.03	0.04	0.05	0.06	0.07	0.08	0.09	0.1
1	0.01	0.01	0.02	0.02	0.03	0.03	0.04	0.04	0.05	0.05
1.5	< 0.00	0.01	0.01	0.01	0.02	0.02	0.02	0.03	0.03	0.03
2	< 0.00	0.01	0.01	0.01	0.01	0.02	0.02	0.02	0.02	0.03
2.5	< 0.00	< 0.00	0.01	0.01	0.01	0.01	0.01	0.02	0.02	0.02
3	< 0.00	< 0.00	0.01	0.01	0.01	0.01	0.01	0.01	0.02	0.02
3.5	< 0.00	< 0.00	<0.00	0.01	0.01	0.01	0.01	0.01	0.01	0.01
4	< 0.00	< 0.00	<0.00	0.01	0.01	0.01	0.01	0.01	0.01	0.01
4.5	< 0.00	< 0.00	<0.00	< 0.00	0.01	0.01	0.01	0.01	0.01	0.01
5	<0.00	<0.00	<0.00	<0.00	0.01	0.01	0.01	0.01	0.01	0.01

**Table 2**: Infusion rates when using insulin concentration **0.2 unit/mL** (Suggested weight 1 to  $\leq$ 3 kg)

Rate	0.1	0.2	0.3	0.4	0.5	0.6	0.7	0.8	0.9	1	
(mL/hr)											
Weight		Approximate unit /kg/hour									
(kg)			1	1	1	1	1		1		
0.5	0.04	0.08	0.12	0.16	0.2	0.24	0.28	0.32	0.36	0.4	
1	0.02	0.04	0.06	0.08	0.1	0.12	0.14	0.16	0.18	0.2	
1.5	0.01	0.03	0.04	0.05	0.07	0.08	0.09	0.11	0.12	0.13	
2	0.01	0.02	0.03	0.04	0.05	0.06	0.07	0.08	0.09	0.1	
2.5	0.01	0.02	0.02	0.03	0.04	0.05	0.06	0.06	0.07	0.08	
3	0.01	0.01	0.02	0.03	0.03	0.04	0.05	0.05	0.06	0.07	
3.5	0.01	0.01	0.02	0.02	0.03	0.03	0.04	0.05	0.05	0.06	
4	0.01	0.01	0.02	0.02	0.03	0.03	0.04	0.04	0.05	0.05	
4.5	< 0.01	0.01	0.01	0.02	0.02	0.03	0.03	0.04	0.04	0.04	
5	< 0.01	0.01	0.01	0.02	0.02	0.02	0.03	0.03	0.04	0.04	

**Table 3**: Infusion rates when using insulin concentration **0.8 unit/mL** (Suggested weight >3kg)

Rate (mL/hr)	0.1	0.2	0.3	0.4	0.5	0.6	0.7	0.8	0.9	1
Weight (kg)	Approximate unit /kg/hour									
0.5	0.16	0.32	0.48	0.64	0.8	0.96	1.12	1.28	1.44	1.6
1	0.08	0.16	0.24	0.32	0.4	0.48	0.56	0.64	0.72	0.8
1.5	0.05	0.11	0.16	0.21	0.27	0.32	0.37	0.43	0.48	0.53
2	0.04	0.08	0.12	0.16	0.2	0.24	0.28	0.32	0.36	0.4
2.5	0.03	0.06	0.1	0.13	0.16	0.19	0.22	0.26	0.29	0.32
3	0.03	0.05	0.08	0.11	0.13	0.16	0.19	0.21	0.24	0.27
3.5	0.02	0.05	0.07	0.09	0.11	0.14	0.16	0.18	0.21	0.23

### **Newborn Use Only**

2025

4	0.02	0.04	0.06	0.08	0.1	0.12	0.14	0.16	0.18	0.2
4.5	0.02	0.04	0.05	0.07	0.09	0.11	0.12	0.14	0.16	0.18
5	0.02	0.03	0.05	0.06	0.08	0.1	0.11	0.13	0.14	0.16

Rate (mL/hr) =  $\frac{\text{Dose (unit/kg/hour) x Weight (kg)}}{\text{Concentration (unit/mL)}}$ 

Dose (unit/kg/hour) =  $\frac{\text{Rate (mL/hr) x Concentration (unit/mL)}}{\text{Rate (mL/hr) x Concentration (unit/mL)}}$ 

Weight (kg)

VERSION/NUMBER	DATE	
Original	3/05/2017	
Revised 1.1	19/05/2017	
Revised 1.2	20/06/2017	
Revised 2.0	18/03/2021	
Revised 3.0	24/06/2022	
Version 4.0	26/09/2025	
REVIEW	26/09/2030	

#### **Authors Contribution**

Original author/s	Srinivas Bolisetty
Evidence Review	David Osborn
Expert review	Charles Verge, Shihab Hameed, Uma Visser
Nursing Review	Ruth Jackson, Celia Cunha Brites, Charles Tian, Tiffany Kwan
Pharmacy Review	Rebecca O'Grady, Kerrie Knox, Thao Tran, Michelle Jenkins, Susanah Brew
ANMF Group contributors	Nilkant Phad, Bhavesh Mehta, Rebecca Barzegar, Mohammad Irfan Azeem, Rebecca
	O'Grady, Cindy Chen, Thao Tran, Celia Cunha Brites, Kerrie Knox, Susannah Brew, Bryony
	Malloy, Renae Gengaroli, Samantha Hassall, Jutta van den Boom, Amber Seigel, Tiffany
	Kwan, Charles Tian, Trong Tran, Emma Watson
Final editing	Srinivas Bolisetty
Electronic version	Cindy Chen, Thao Tran, Ian Callander
Facilitator	Srinivas Bolisetty

