## Vecuronium

Newborn use only

Alaut	Use alout modioation. Use viel of accoing signific	ant antiant have when word in avery 1
Alert	t High-alert medication: High risk of causing significant patient harm when used in error. <sup>1</sup> This drug should be administered in the presence of personnel trained in advanced airwa	
	management. Reversal agents should be immedia	
	Suggest regular cessation of infusion for a few to	
		for continued paralysis and adequacy of sedation or
	analgesia.	
		y flushed to avoid unintended paralysis during later
	use of the line.	
	Eye lubricant should be used whilst patient is rece	iving vecuronium.
Indication	1. Skeletal muscle relaxation or paralysis in med	hanically ventilated infants.
	2. For elective endotracheal intubation.	
Action	Nondepolarizing neuromuscular blockers are competitive acetylcholine (ACh) antagonists that bi	
	directly to nicotinic receptors on the postsynaptic	membrane, thus blocking the binding of ACh so the
	motor endplate cannot depolarize. Onset of actio	n is 1-2 minutes; duration of action is 30-40 minutes.
Drug Type	Non-depolarising neuromuscular blocking agent.	
Trade Name	Vecure Powder for injection, Vecure Sun Powder	for injection
Presentation	10 mg of vecuronium bromide in glass vial (powd	
Dose	Intubation	
	IV bolus – 0.1 mg/kg	
	Muscle relaxation*#	
	Intermittent IV bolus	
	0.1 mg/kg (0.03-0.15 mg/kg) IV push eve	ry 1-2 hours as required
		ry 1 2 hours as required.
	Continuous IV infusion (with or without	loading dose)
		am/kg/hour). Start 20 minutes post bolus recovery.
		red neuromuscular blockade is achieved.
	fittate in 10% dose increments diffitues	red field official blockade is achieved.
	* Provide eye protection and instil lubricating eye	drops avany 2 hours
	<sup>#</sup> Sensation remains intact; additional sedation &	
Dece ediustreent		analgesia should be used for painful procedures.
Dose adjustment	Therapeutic hypothermia – No information. ECMO – No information.	
		University dynamics of entire way be availabled
	Renal impairment – No specific dose adjustment.	
		t. However, hepatic impairment decreases clearance
<u> </u>	resulting in prolonged duration of action.	
Route		
Maximum Dose	IV bolus: 0.2 mg/kg	
	IV infusion: 0.2 mg/kg/hour.	
Total cumulative		
dose		
Droparation		
Preparation	IV bolus:	
r repai ation	<u>IV bolus:</u> Add 5 mL water for injection to 10 mg of vecuroni	um powder for reconstitution vial to make a 2
riepaiation		um powder for reconstitution vial to make a 2
r i epai ation	Add 5 mL water for injection to 10 mg of vecuron mg/mL solution.	um powder for reconstitution vial to make a 2 of vecuronium) and add to 2 mL of sodium chloride
r i cµai ativi	Add 5 mL water for injection to 10 mg of vecuron mg/mL solution.	of vecuronium) and add to 2 mL of sodium chloride
	Add 5 mL water for injection to 10 mg of vecuron mg/mL solution. <b>Further dilute</b> : From this vial, draw up 2 mL (4 mg	of vecuronium) and add to 2 mL of sodium chloride
r i cµai ation	Add 5 mL water for injection to 10 mg of vecuroni mg/mL solution. <b>Further dilute</b> : From this vial, draw up 2 mL (4 mg 0.9% to make a final volume of 4 mL with a conce	of vecuronium) and add to 2 mL of sodium chloride
r i epai ation	Add 5 mL water for injection to 10 mg of vecuron mg/mL solution. <b>Further dilute</b> : From this vial, draw up 2 mL (4 mg	of vecuronium) and add to 2 mL of sodium chloride
	Add 5 mL water for injection to 10 mg of vecuroni mg/mL solution. <b>Further dilute</b> : From this vial, draw up 2 mL (4 mg 0.9% to make a final volume of 4 mL with a conce	of vecuronium) and add to 2 mL of sodium chloride
	Add 5 mL water for injection to 10 mg of vecuroni mg/mL solution. <b>Further dilute</b> : From this vial, draw up 2 mL (4 mg 0.9% to make a final volume of 4 mL with a conce	of vecuronium) and add to 2 mL of sodium chloride
	Add 5 mL water for injection to 10 mg of vecuroni mg/mL solution. Further dilute: From this vial, draw up 2 mL (4 mg 0.9% to make a final volume of 4 mL with a conce <u>IV infusion:</u> <u>Infant &lt;1.5 Kg</u>	of vecuronium) and add to 2 mL of sodium chloride ntration of 1 mg/mL.
	Add 5 mL water for injection to 10 mg of vecuroni mg/mL solution. <b>Further dilute</b> : From this vial, draw up 2 mL (4 mg 0.9% to make a final volume of 4 mL with a conce <u>IV infusion:</u>	of vecuronium) and add to 2 mL of sodium chloride

## Vecuronium

Newborn use only

	First Step: Add 5 mL water for injection to 10 mg v 2 mg/mL solution (Note: May need multiple vials Further dilute: Draw up 10 mL/kg of this solution chloride 0.9% to make a final volume of 50mL with microgram/kg/hour. IV bolus dose from this solution: 0.25 mL = 100 m Infant ≥1.5 Kg	(20 mg/kg) and dilute with glucose 5% or sodium in a concentration of <b>0.25 mL/hour = 100</b>
	Infusion rate	Prescribed amount
		10 mg/kg vecuronium and make up to 50 ml
	1ml/hour = 200 microgram/kg/hour10 mg/kg vecuronium and make up tFirst Step: Add 5 mL water for injection to 10 mg vecuronium powder for reconstitution	
	<ul> <li>2 mg/mL solution (Note: May need multiple vials  </li> <li>Further dilute: Draw up 5 mL/kg of this solution (2 chloride 0.9% to make a final volume of 50mL with microgram/kg/hour.</li> <li>IV bolus dose from this solution: 0.5 mL = 100 mi</li> </ul>	based on bodyweight). L0 mg/kg) and dilute with glucose 5% or sodium n a concentration of <b>0.5 mL/hour = 100</b>
Administration	IV bolus: Administer over several seconds.	
	IV infusion via syringe pump. Flush line adequately after each dose with sodium incompatibility with other medications during late	n chloride 0.9% to avoid unintended paralysis and/or ar use of the line.
Monitoring	Continuous cardio-respiratory and pulse oximetry	monitoring.
		tion, and blood pressure (invasive or non-invasive).
	Monitor electrolytes and renal function.	
<b>C</b>	Monitor injection site for signs of extravasation.	- Ath - fammedation
Contraindications	Hypersensitivity to vecuronium or any component Cross-sensitivity with other neuromuscular-blocki	
	patients with previous anaphylactic reactions.	ng agents may occur, use with extreme caution in
	Severe electrolyte abnormalities.	
Precautions	Avoid prolonged usage.	
	Factors which can increase duration of neuromus	scular blockade:
	Acidosis, hypothermia, neuromuscular disease, hepatic disease, hypokalaemia, hypermagnesaemia, renal failure, and younger age. Vecuronium is lipid soluble and is predominantly excreted via the liver so poor liver function can cause prolonged effects. <b>Factors which can decrease duration of neuromuscular blockade:</b> Alkalosis and hyperkalaemia.	
Use cautiously in neonates with hepatic or renal impairment and in neonates with f electrolyte imbalance.		mpairment and in neonates with fluid and
	Suggest regular cessation of infusion, possibly eve to assess the need for continued paralysis and ade Monitoring of fluid balance is essential due to risk	
/	Aminoglycosides & general anaesthetics can incre blockade.	ase (potentiate) duration of neuromuscular
	<b>Corticosteroids:</b> Concomitant use with corticoster development of acute quadriplegic myopathy syn provided no evidence for increased risk of neuron distress syndrome (ARDS) with the use of corticos	drome (AQMS) in adults. <sup>3</sup> However, Recent trials hyopathy in patients with sepsis or acute respiratory
		duration of neuromuscular blockade.

# Vecuronium

Newborn use only

Drug Interactions	Antimicrobials like aminoglycosides, tetracyclines, polymyxins, and clindamycin can potentiate neuromuscular blockade. <sup>3</sup>
	Inhaled anaesthetics can potentiate neuromuscular blockade. <sup>3</sup>
	Anti-epileptics can make patients resistant to vecuronium. <sup>3</sup>
	Local anaesthetics can potentiate neuromuscular blockade. <sup>3</sup>
	Aminoglycosides & general anaesthetics can increase duration of neuromuscular blockade.
	<b>Corticosteroids:</b> Concomitant use with corticosteroids has been reported to be associated with
	development of acute quadriplegic myopathy syndrome (AQMS) in adults. <sup>3</sup> However, Recent trials
	provided no evidence for increased risk of neuromyopathy in patients with sepsis or acute respiratory
	distress syndrome (ARDS) with the use of corticosteroids or neuromuscular blockers. <sup>17</sup>
	Dexamethasone and hydrocortisone may result in decreased vecuronium effectiveness, prolonged
	muscle weakness, and myopathy. <sup>3</sup>
	Adrenaline (epinephrine) can reduce (antagonise) duration of neuromuscular blockade.
Adverse	Hypoxaemia may occur because of inadequate ventilation and deterioration in pulmonary mechanics.
Reactions	Hypotension and bradycardia, particularly when used in combination with opioids.
	Prolonged paralysis after long-term use.
	Rare: Anaphylactic reaction and tachycardia.
Overdose	Supportive measure: Ventilatory support and sedation.
	Reversal of neuromuscular blockade can be achieved by neostigmine (refer to special comments).
	For information on the management of overdose, contact the Poisons Information Centre on 13 11 26
	(Australia).
Compatibility	Fluids: <sup>3</sup> glucose 5%, sodium chloride 0.9%.
	V site <sup>3</sup> glugges (aming acid colutions, advancing (aning bring), altrastadil, amikasin sulfate
	Y-site: <sup>3</sup> glucose/amino acid solutions, adrenaline (epinephrine), alprostadil, amikacin sulfate, aminophylline, amiodarone, ampicillin, atenolol, azithromycin, aztreonam, caffeine citrate, calcium
	chloride, calcium gluconate, cefazolin, ceftazidime, ceftriaxone, ciprofloxacin, clindamycin,
	dexamethasone sodium phosphate, dexmedetomidine, digoxin, diltiazem, dobutamine, dopamine,
	enalaprilat, adrenaline (epinephrine), erythromycin lactobionate, esmolol, fentanyl, fluconazole,
	fluorouracil, fosphenytoin, gentamicin, glycopyrrolate, heparin, hydralazine, hydrocortisone sodium
	succinate, insulin (regular), isoprenaline, labetalol, lidocaine, linezolid, lorazepam, magnesium sulfate,
	meropenem, metaprolol, metronidazole, midazolam, milrinone, morphine, naloxone, nicardipine,
	nitroglycerin, norepinephrine, octreotide, ondansetron, pamidronate, pentoxifylline, phenobarbital,
	phenylephrine, potassium acetate, potassium chloride, propofol at vecuronum concentrations of ≤1
	mg/mL, propranolol hydrochloride, ranitidine hydrochloride, remifentanil, sodium acetate, sodium
	bicarbonate, sodium nitroprusside, sodium phosphate, streptozocin, succinylcholine, tacrolimus,
	theophylline, ticarcillin disodium/clavulante potassium, tigecycline, tobramycin,
	sulfamethoxazole/trimethoprim, and vancomycin hydrochloride, vasopressin, verapamil,
	voriconazole, zidovudine, and zoledronic acid.
Incompatibility	Fluids: No information. No information on lipid emulsions.
	Y site: <sup>3</sup> Aciclovir, amphotericin B (all compounds), cefepime, cefotaxime, diazepam, furosemide,
	ganciclovir, ibuprofen lysine, imipenem/cilastatin sodium, methylprednisolone sodium succinate,
	micafungin sodium, pantoprazole, phenytoin, piperacillin sodium, piperacillin-tazobactam, propofol at
Chale iliter	vecuronium concentrations >1 mg/mL, sulbactam/durlobactam, and thiopental sodium.
Stability Storage	Diluted solution stable for up to 24 hours. Discard any unused solution. Store below 25°C. Protect from light.
Storage	Store below 25°C. Protect from light. Store in accordance with local policies.
Excipients	Citric acid, dibasic sodium phosphate, sodium hydroxide and/or phosphoric acid, and mannitol.
-	
Special	Muscle relaxation is reversed by neostigmine (50 microgram/kg) and atropine (20 microgram/kg).
Comments	Sugammadex is being increasingly used with extrapolated information from other populations.
	Sensation remains intact: sedation & analgesia should be used for painful procedures.
	Provide eye protection and instil lubricating eye drops every 2 hours.
	Vecuronium produces less tachycardia and hypotension when compared with pancuronium. <sup>15,16</sup>
	The neuromuscular blockade of vecuronium is of shorter duration than that of pancuronium. <sup>15,16</sup>

### Vecuronium Newborn use only

	Prior administration of suxamethonium shortens onset and may increase depth of blockade; reduce dose and give vecuronium only after recovery from suxamethonium-induced neuromuscular blockade.
Evidence	<ul> <li>Bockade.</li> <li>Background</li> <li>Nondepolarizing neuromuscular blocking agents (NMBA) can be classified into 2 classes: steroidal (rocuronium, vecuronium, pancuronium) or benzylisoquinoline (mivacurium, atracurium, cisatracurium). Nondepolarizing neuromuscular blockers are competitive acetylcholine (ACh) antagonists that bind directly to nicotinic receptors on the postsynaptic membrane, thus blocking the binding of ACh so the motor endplate cannot depolarize. This leads to skeletal muscle paralysis.<sup>2</sup></li> <li>Paralysis occurs sequentially because of the differing sensitivity of muscles to NMBAs as well as blood flow to the area. Generally, paralysis begins with smaller, fast twitch muscles such as the eyes and larynx, then affects the limbs, neck, trunk, and upper airway, and eventually progresses to the intercostals and diaphragm until respiration terminates. Recovery from paralysis occurs in the reverse order with function of the diaphragm returning first.<sup>4</sup></li> <li>Vecuronium: vecuronium is structurally related to pancuronium. It has a greater potency, shorter duration of action, lack significant cardiovascular effects (tachycardia), and less cumulative properties. Vecuronium produces less tachycardia and hypotension when compared with pancuronium.<sup>15,16</sup> The neuromuscular blockade of vecuronium is of shorter duration than that of pancuronium.<sup>15,16</sup> Time of onset of action is 90-120 seconds after IV bolus, with a duration of effect that lasts only 30-40 minutes. Intermittent bolus dosing would need to be so frequent (i.e., every 30 to 60 minutes) that continuous IV infusion is preferred over intermittent boluses to maintain paralysis in ventilated infants.<sup>5</sup> In comparison, rocuronium is an analogue of vecuronium with a more rapid onset of action (20-100 seconds), but less potent than vecuronium, and hence larger doses (example, 600 microgram/kg of rocuronium, ompared to 100 microgram/kg of vecuronium). Tachycardia is more frequent with rocuronium, while vecuronium lack</li></ul>
	<ul> <li>microgram/kg) meaning less dose is needed in neonates compared to children.<sup>7</sup></li> <li>Efficacy</li> <li><u>Muscle relaxation</u></li> <li>Two prospective studies by Meretoja et al in 1988 and 1989 determined the dose responses with vecuronium bolus and continuous infusion in paediatric population.<sup>7,8</sup> The bolus dose required to achieve effective neuromuscular blockade in neonates was 40% less than in children. The median maintenance dose of 0.1 mg/kg is required in neonates to maintain 1 hour of neuromuscular blockade, in comparison to 0.217 mg/kg/hour in children 3-10 years old.</li> <li>Fitzpatrick et al studied vecuronium to facilitate paralysis in mechanically ventilated paediatric population (4 neonates, and 11 infants and children). A loading dose of 0.1 mg/kg was followed by an infusion of 0.1 mg/kg/hour. The titration rate was adjusted to maintain a neuromuscular block of approximately 90% as assessed by the presence of one response to a train of four (TOF) stimulation. The duration of the infusions varied from 9.5 to 179 hours. Mean recovery times after stopping the infusion were 51.7 (±17.6) and 46.8 (± 16.5) minutes for the children and neonates respectively. No adverse cardiovascular or toxic effects were noted.<sup>9</sup> Fisher et al determined the recovery period (time from injection to 90% recovery) after a bolus of vecuronium. Recovery was longest for infants (73±27 minutes), compared to children (35±6 minutes).<sup>10</sup> The longer recovery period in neonates is thought to be due higher volume of distribution.</li> <li>Hodges et al evaluated the appropriate vecuronium infusion rates in 12 neonates/infants and 18 children using train of four (TOF) monitoring. Neonates and infants required 45% less vecuronium (mean infusion rate 0.54 mg/kg/hour) than older children (0.99 mg/kg/hour) and had faster recovery (45 min vs 65 min), with no evidence of prolonged weakness.<sup>11</sup></li> </ul>
	Safety Adults with hepatic and renal failure have been shown to experience prolonged neuromuscular blockade. <sup>12,13</sup>

### Vecuronium Newborn use only

Pharmacokinetics

2025

	Pharmacokinetics
	Hepatobiliary clearance is the primary route of elimination, accounting for approximately 50% of the
	dose. Vecuronium is metabolised rapidly in the liver to 3-desacetyl-vecuronium, which is 50–70% as
	potent as the parent compound. This metabolite is cleared primarily by renal elimination.
	Approximately 20–30% of vecuronium is excreted unchanged in urine. <sup>9,11,12</sup>
	Time of onset of action is 90-120 seconds after IV bolus, with a duration of effect that lasts only 30-4
	minutes. (prolonged with higher doses and in preterm infants). <sup>5,14</sup>
Practice points	Eye lubrication should be applied to all patients.
References	1. Clinical Excellence Commission. Neuromuscular blocking agents.
	https://www.cec.health.nsw.gov.au/keep-patients-safe/medication-safety/high-risk-
	medicines/neuromuscular-blocking-agents. Accessed on 7 October 2024.
	2. Cook D, Simons DJ. Neuromuscular Blockade. [Updated 2023 Nov 13]. In: StatPearls [Internet].
	Treasure Island (FL): StatPearls Publishing; 2024 Jan Available from:
	https://www.ncbi.nlm.nih.gov/books/NBK538301/.
	3. MerativeTM Micromedex <sup>®</sup> Complete IV Compatibility (electronic version). Merative, Ann Arbor
	Michigan, USA. Available at: https://www.micromedexsolutions.com/ (cited: Oct/8/2024).
	4. Warr J, Thiboutot Z, Rose L, Mehta S, Burry LD. Current Therapeutic Uses, Pharmacology, and
	Clinical Considerations of Neuromuscular Blocking Agents for Critically III Adults. Annals of
	Pharmacotherapy. 2011;45(9):1116-26.
	5. Kandasamy J, Carlo WA. Pharmacologic therapies IV. In Assisted ventilation of the neonate. 201
	Elsevier Inc.
	6. Meakin GH. Neuromuscular blocking drugs in infants and children. Continuing Education in
	Anaesthesia Critical Care & Pain. 2007;7(5):143-7.
	<ol> <li>Meretoja OA, Wirtavuori K, Neuvonen PJ. Age-dependence of the dose-response curve of</li> </ol>
	vecuronium in pediatric patients during balanced anesthesia. Anesthesia & Analgesia.
	1988;67(1):21-6.
	8. Meretoja O. Is vecuronium a long-acting neuromuscular blocking agent in neonates and infants
	British Journal of Anaesthesia. 1989;62(2):184-7.
	9. Fitzpatrick KT, Black GW, Crean PM, Mirakhur RK. Continuous vecuronium infusion for prolonge
	muscle relaxation in children. Can J Anaesth. 1991;38:169-74.
	10. Fisher DM, Miller RD. Neuromuscular effects of vecuronium (ORG NC45) in infants and children
	during N2O, halothane anesthesia. Anesthesiology. 1983;58(6):519-23.
	11. Hodges U. Vecuronium infusion requirements in paediatric patients in intensive care units: the
	use of acceleromyography. British journal of anaesthesia. 1996;76(1):23-8.
	12. Bencini A, Scaf A, Sohn Y, Kersten-Kleef U, Agoston S. Hepatobiliary disposition of vecuronium
	bromide in man. British Journal of Anaesthesia. 1986;58(9):988-95.
	13. Lynam DP, Cronnelly R, Castagnoli KP, Canfell PC, Caldwell J, Arden J, et al. The
	pharmacodynamics and pharmacokinetics of vecuronium in patients anesthetized with
	isoflurane with normal renal function or with renal failure. Anesthesiology. 1988;69(2):227-31.
	14. Johnson PN, Miller J, Gormley AK. Continuous-infusion neuromuscular blocking agents in
	critically ill neonates and children. Pharmacotherapy: The Journal of Human Pharmacology and
	Drug Therapy. 2011;31(6):609-20.
	15. Basta SJ, Savarese JJ, Ali HH et al. Vecuronium does not alter serum histamine within the clinica
	dose range. Anesthesiology 1983;59:A273.
	16. Son SL, Waud BE, Waud DR. A comparison of the neuromuscular blocking and vagolytic effects
	of ORG NC45 and pancuronium. Anesthesiology 1981;55:12–18.
	<ol> <li>Annane D. What is the evidence for harm of neuromuscular blockade and corticosteroid use in</li> </ol>
	the intensive care unit? Seminars in respiratory and critical care medicine 2016;37:51-56.

#### Vecuronium Newborn use only

VERSION/NUMBER	DATE
Original 1.0	10/04/2017
Version 2.0	27/06/2019
Version 3.0	17/10/2024
Version 3.0 (minor errata)	1/11/2024
Current 4.0	10/07/2025
REVIEW	10/07/2030

#### Authors Contribution of the current version

Current version authors	Srinivas Bolisetty, Rebecca O'Grady, Nilkant Phad	
Evidence Review	Srinivas Bolisetty, Rebecca O'Grady, Nilkant Phad	
Expert review		
Nursing Review	Benjamin Emerson-Parker	
Pharmacy Review	Rebecca O'Grady	
ANMF Group	Bhavesh Mehta, Rebecca Barzegar, Mohammed Irfan Azeem, Thao Tran, Cindy Chen,	
contributors	Susannah Brew, Michelle Jenkins, Celia Cunha Brites, Bryony Malloy, Renae Gengaroli,	
	Samantha Hassall, Amber Seigel, Emma Watson, Tiffany Kwan, Charles Tian	
Final editing	Srinivas Bolisetty	
Electronic version	Thao Tran, Cindy Chen, Ian Callander	
Facilitator	Srinivas Bolisetty	

#### Citation for the current version

Bolisetty S, O'Grady R, Phad N, Emerson-Parker B, Mehta B, Barzegar R, Azeem MI, Tran T, Chen C, Brew S, Jenkins M, Brites CC, Malloy B, Gengaroli R, Hassall S, Watson E, Kwan T, Seigel A, Callander I. Vecuronium. Consensus formulary by the Australasian Neonatal Medicines Formulary group. Version 4 dated 10 July 2025. www.anmfonline.org