# Meropenem

## **Newborn use only**

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Alert		The Antimicrobial Stewardship Team recommends this drug is listed under the following category:						
		Restricted.						
	Widespread use of carbapenems has been linked with increasing prevalence of infections caused							
		methicillin-resistant Staphylococcus aureus (MRSA), vancomycin-resistant enterococci (VRE), multi-						
	resistant Gram-negative organisms and Clostridium difficile.  Severe infections (e.g., sepsis or meningitis) caused by Gram-negative organisms resistant to other							
Indication								
		ventional antibiotics but susce	eptible to meropen	iem e.g., Extended	Spectrum Beta Lacta	mase		
		(ESBL)-producing organisms.  Note: Meropenem is NOT active against many resistant Gram-positive organisms, such as MRSA and						
				•	_			
	most Staphylococcus epidermidis. Vancomycin is first-line therapy for these. Meropenem of activity against penicillin-susceptible Gram-positive organisms and most anaerobic organis							
	individual advice, discuss therapy with a microbiologist or infectious diseases physician.							
Action	Meropenem is a carbapenem. It inhibits cell wall synthesis. (1)							
	met openem is a carbapenem it initials cen wan synthesis. (1)							
	Mer	Meropenem is a better choice than imipenem for central nervous system infections. Meropenem						
		ins a higher concentration in	-	-	•			
		lower incidence of seizures than imipenem.						
Drug type	Carl	papenem antibiotic.						
Trade name	Mer	ropenem APOTEX, Meropener	m DBL, Meropenen	n GH, Meropenem	Juno, Meropenem Ka	ıbi,		
	Meropenem Sandoz, Merrem							
Presentation	500	) mg vial						
	100	0 mg vial						
Dose	Non	n-CNS and Non-Pseudomonas	Sepsis					
		Gestational Age at birth	Postnatal Age	Dose	Interval			
		< 32 <sup>+0</sup> weeks	0–13 days	20 mg/kg	12 hourly			
		< 32 <sup>+0</sup> weeks	14+ days	20 mg/kg	8 hourly			
		≥ 32 <sup>+0</sup> weeks	0–13 days	20 mg/kg	8 hourly			
		≥ 32 <sup>+0</sup> weeks	14+ days	30 mg/kg	8 hourly			
	Mei	ningitis and Pseudomonas Se	psis			_		
		Gestational Age at birth	Postnatal Age	Dose	Interval			
		Any	Any	40 mg/kg	8 hourly			
Dose adjustment	Asse	Assess for renal impairment prior to using higher doses as meropenem is primarily excreted via						
	kidn	neys.						
Maximum dose								
Total cumulative								
dose								
Route	IV ir	nfusion.						
Preparation		Infants ≤1kg						
		9.6 mL of water for injection	_	_				
	Add 19.1 mL of water for injection to 1g vial to make a 50 mg/mL solution.  FURTHER DILUTE  Draw up 2 mL (100 mg of meropenem) of the above solution and add 8 mL sodium chloride 0.9% to make a final volume of 10 mL with a final concentration of 10 mg/mL.							
	Infa	Infants >1kg or fluid restricted.						
		9.6 mL of water for injection	to 500 mg vial to n	nake a 50 mg/ml (	solution			
		THER DILUTE	to 500 mg viai to m	nake a 50 mg/me.	oracion.			
	Draw up 4 mL (200 mg of meropenem) of the above solution and add 6 mL sodium chloride 0.9% to							
		make a final volume of 10 mL with a concentration of 20 mg/mL.						
Administration	IV infusion over 4 hours. (5)							
		• •	tes if longer infusio	n time is not feasi	ble.			
Monitoring	May be given over 15 to 30 minutes if longer infusion time is not feasible.  Renal function.							
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	_	r function.						
	Live							

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Contraindications	Hypersensitivity to penicillins, cephalosporins and carbapenems.			
Precautions	Colitis–due to risk of pseudomembranous colitis.			
	Renal impairment.			
Drug interactions				
valproate, which may cause seizures.				
Adverse reactions	Phlebitis, diarrhoea (up to 6% in children), anaemia and eosinophilia.			
Compatibility	Fluids: sodium chloride 0.9% (preferred for stability), glucose 5%, glucose 10%,			
	Y-site: Amino acid solutions, anidulafungin, caspofungin, linezolid, atropine, dexamethasone sodium, gentamicin, heparin sodium, metronidazole.			
Incompatibility	Fluids: Mannitol 10%			
incompatibility	Fidius. Maillitoi 10%			
	Y-site: Dolasetron, ketamine, zidovudine.			
Stability	Use immediately after preparation.			
Diluted solutions are potentially unstable, particularly glucose containing solutions and s				
	discarded if not used immediately.			
Storage	Vial: Store at room temperature.			
Excipients	Sodium carbonate			
Special comments	Meropenem 1 g vial contains 3.92 mmol of sodium.			
Evidence	Efficacy:			
	Carbapenems may be considered the treatment of choice for empirical treatment of patients with ESBL-producing <i>Enterobacteriaceae</i> bacteraemia. A systematic review of carbapenems for the treatment of patients with extended-spectrum $\beta$ -lactamase (ESBL)-positive <i>Enterobacteriaceae</i> bacteraemia involving 1584 patients, mostly adults showed lower mortality than non-Beta-lactam/Beta-Lactam Inhibitor combination antibiotics for definitive [risk ratio (RR) 0.65, 95% CI 0.47–0.91] and empirical (RR 0.50, 95% CI 0.33–0.77) treatment. No statistically significant differences in mortality were found between carbapenems and BL/BLIs administered as definitive (RR 0.52, 95% 0.23–1.13) or empirical (RR 0.91, 95% CI 0.66–1.25) treatment (LOE 1, GOR C).			
	A retrospective case series of 100 neonates infected by extended-spectrum beta-lactamase-producing <i>Klebsiella</i> species showed higher mortality in those neonates not started on empirical meropenem or Piperacillin + tazobactam and amikacin (OR – 17.01, 95% CI 2.41–120.23) (LOE IV, GOR C). <sup>3</sup>			
	A RCT reported a prolonged infusion (4 hours) of meropenem (20 mg/kg/dose every 8 hours and 40 mg/kg/dose every 8 hours in meningitis and Pseudomonas infection) in 102 neonates with gramnegative late onset infection is associated with higher rate of clinical improvement, microbiologic eradication, less neonatal mortality (14% versus 31%; p=0.03), shorter duration of respiratory support and less acute kidney injury compared with the conventional strategy (30 minute infusion) [LOE II GOR B]. <sup>5</sup>			
	Pharmacokinetics: Meropenem is primarily excreted via the kidneys. Meropenem clearance is influenced by serum creatinine and postmenstrual age in neonates. <sup>2</sup> A comparative pharmacokinetic study of short (30 minute) versus long (4 hour) infusion in neonates showed short infusion resulted in a higher mean drug concentration in serum (C(max)) than a prolonged infusion. <sup>6</sup> However, a longer infusion may have greater efficacy. <sup>5</sup> There is a knowledge gap in pharmacokinetic (PK) studies of neonates with renal impairment. <sup>2,3</sup> However, dose adjustment for renal failure may not be appropriate in cases where severe sepsis is probably responsible for acute renal failure [expert opinion].			
Practice points	<b>Dose:</b> Multicentre, prospective PK study conducted in USA suggested a dosing strategy of 20 mg/kg every 12 hours in infants < 32 weeks GA and PNA < 14 days; 20 mg/kg every 8 hours in infants < 32 weeks GA and PNA $\geq$ 14 days and in infants $\geq$ 32 weeks GA and PNA < 14 days; and 30 mg/kg every 8 hours in infants $\geq$ 32 weeks GA and PNA $\geq$ 14 days to achieve therapeutic concentrations in infants with suspected intra-abdominal infections. <sup>4</sup>			
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### References Pacifici GM, Allegaert K. Clinical pharmacology of carbapenems in neonates. J Chemother 2014;26(2):67-73. Vardakas KZ, Tansarli GS, Rafailidis PI, Falagas ME. Carbapenems versus alternative antibiotics for the treatment of bacteraemia due to Enterobacteriaceae producing extended-spectrum betalactamases: a systematic review and meta-analysis. J Antimicrob Chemother 2012;67(12):2793-Velaphi S, Wadula J, Nakwa F. Mortality rate in neonates infected with extended-spectrum blactamase-producing Klebsiella species and selective empirical use of meropenem. Ann Trop Paediatr 2009;29:101-10. Smith PB, Cohen-Wolkowiez M, Castro LM, Poindexter B, Bidegain M, Weitkamp JH, et al, Meropenem Study Team. Population pharmacokinetics of meropenem in plasma and cerebrospinal fluid of infants with suspected or complicated intra-abdominal infections. Pediatr Infect Dis J 2011;30(10):844-9. Shabaan AE, Nour I, Elsayed Eldegla H, Nasef N, Shouman B, Abdel-Hady H. Conventional Versus Prolonged Infusion of Meropenem in Neonates With Gram-negative Late-onset Sepsis: A Randomized Controlled Trial. Pediatric Infectious Disease Journal. 2017;36:358-63. Padari H, Metsvaht T, Korgvee LT, Germovsek E, Ilmoja ML, Kipper K, Herodes K, Standing JF, Oselin K, Lutsar I. Short versus long infusion of meropenem in very-low-birth-weight neonates. Antimicrob Agents Chemother 2012;56(9):4760-4.

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