Meropenem

Newborn use only

Alert	The /	Antimicrohial Stewardshin To	eam recommends t	his drug is listed u	nder the following cate	egory.			
Alcit	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 by							
	methicillin-resistant Staphylococcus aureus (MRSA), vancomycin-resistant enterococci (VRE), multi-								
	resistant Gram-negative organisms and Clostridium difficile.								
Indication		Severe infections (e.g., sepsis or meningitis) caused by Gram-negative organisms resistant to other							
		conventional antibiotics but susceptible to meropenem e.g., Extended Spectrum Beta Lactamase							
	(ESBI	(ESBL)-producing organisms.							
	Note	e: Meropenem is NOT active	against many resist	ant Gram-positive	organisms, such as M	RSA and			
	most	most Staphylococcus epidermidis. Vancomycin is first-line therapy for these. Meropenem does have							
	activ	activity against penicillin-susceptible Gram-positive organisms and most anaerobic organisms. For							
	indiv	individual advice, discuss therapy with a microbiologist or infectious diseases physician.							
Action	Merc	openem is a carbapenem. It i	inhibits cell wall syr	nthesis. (1)					
		openem is a better choice th		-					
		ins a higher concentration in		luid particularly w	ith inflamed meninges	and has a			
		er incidence of seizures than	imipenem.						
Drug type		papenem antibiotic.							
Trade name		openem APOTEX, Meropene	m DBL, Meropener	n GH, Meropenem	Juno, Meropenem Ka	bi,			
		openem Sandoz, Merrem							
Presentation		mg vial							
) mg vial	•						
Dose	Non-	-CNS and Non-Pseudomonas	_		1	٦			
	-	Gestational Age at birth	Postnatal Age	Dose	Interval	_			
		< 32 ⁺⁰ weeks	0–13 days	20 mg/kg	12 hourly	4			
		< 32 ⁺⁰ weeks	14+ days	20 mg/kg	8 hourly	4			
		≥ 32 ⁺⁰ weeks	0–13 days	20 mg/kg	8 hourly				
		≥ 32 ⁺⁰ weeks	14+ days	30 mg/kg	8 hourly				
	Man	singitis and Decudences Co	maia						
	Ivien	ningitis and Pseudomonas Se		Dose	Interval	7			
		Gestational Age at birth Any	Postnatal Age			4			
		Anv	Any	40 mg/kg	8 hourly	vio			
Daga adimeter ant		•	Assess for renal impairment prior to using higher doses as meropenem is primarily excreted via						
Dose adjustment	Asses	ess for renal impairment prior	r to using higher do	ses as meropenen	is primarily excreted				
		ess for renal impairment prior	r to using higher do	ses as meropenen	is primarily excreted				
Maximum dose	Asses	ess for renal impairment prior	r to using higher do	ses as meropenen	n is primarily excreted				
Maximum dose Total cumulative	Asses	ess for renal impairment prior	r to using higher do	ses as meropenen	n is primarily excreted				
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Maximum dose Total cumulative dose Route	Asses kidne	ess for renal impairment prioreys.	r to using higher do	ses as meropenen	n is primarily excreted				
Maximum dose Total cumulative dose Route	Asses kidne IV inf	ess for renal impairment prioreys. fusion. nts <1 kg		·					
Maximum dose Total cumulative dose Route	Asses kidne IV inf Infan	ess for renal impairment prioreys. fusion. nts <1 kg 9.6 mL of water for injection	to 500 mg vial to r	nake a 50 mg/mL s	solution OR				
Maximum dose Total cumulative dose Route	Asses kidne IV inf Infan Add 9	fusion. nts <1 kg 9.6 mL of water for injection 19.1 mL of water for injection	to 500 mg vial to r	nake a 50 mg/mL s	solution OR				
Maximum dose Total cumulative dose Route	IV inf Infan Add 9 Add 1 FURT	fusion. nts <1 kg 9.6 mL of water for injection 19.1 mL of water for injection THER DILUTE	to 500 mg vial to r n to 1g vial to make	make a 50 mg/mL s e a 50 mg/mL solu	solution OR tion.				
Maximum dose Total cumulative dose Route	IV inf Infan Add 9 Add 1 FURT Draw	fusion. nts <1 kg 9.6 mL of water for injection 19.1 mL of water for injection THER DILUTE v up 2 mL (100 mg of merope	to 500 mg vial to r n to 1g vial to make enem) of the above	make a 50 mg/mL solu e a 50 mg/mL solu e solution and add	solution OR tion.				
Maximum dose Total cumulative dose Route	IV inf Infan Add 9 Add 1 FURT Draw	fusion. nts <1 kg 9.6 mL of water for injection 19.1 mL of water for injection THER DILUTE	to 500 mg vial to r n to 1g vial to make enem) of the above	make a 50 mg/mL solu e a 50 mg/mL solu e solution and add	solution OR tion.				
Maximum dose Total cumulative dose Route	IV inf Infan Add 9 Add 1 FURT Draw make	fusion. fusion. nts <1 kg 9.6 mL of water for injection 19.1 mL of water for injectio THER DILUTE v up 2 mL (100 mg of merope e a final volume of 10 mL wit	to 500 mg vial to r n to 1g vial to make enem) of the above	make a 50 mg/mL solu e a 50 mg/mL solu e solution and add	solution OR tion.				
Maximum dose Total cumulative dose Route	IV inf Infan Add 9 Add 1 FURT Draw make	fusion. nts <1 kg 9.6 mL of water for injection 19.1 mL of water for injection THER DILUTE v up 2 mL (100 mg of merope e a final volume of 10 mL wit	to 500 mg vial to r n to 1g vial to make enem) of the above th a final concentra	make a 50 mg/mL se a 50 mg/mL solution and add tion of 10 mg/mL.	solution OR tion. 8 mL sodium chloride				
Maximum dose Total cumulative dose Route	IV inf Infan Add 9 Add 1 FURT Draw make	fusion. fusion. 19.6 mL of water for injection 19.1 mL of water for injection THER DILUTE 19.0 mL of up 2 mL (100 mg of merope 19.1 mL of up 2 mL (100 mg of merope 19.1 mL of up 3 mL wite 19.6 mL of water for injection	to 500 mg vial to r n to 1g vial to make enem) of the above th a final concentra to 500 mg vial to r	make a 50 mg/mL see a 50 mg/mL solution and add tion of 10 mg/mL.	solution OR tion. 8 mL sodium chloride				
	IV inf Infan Add 9 Add 9 Infan Add 9 Add 9 Add 1	fusion. nts <1 kg 9.6 mL of water for injection 19.1 mL of water for injection THER DILUTE v up 2 mL (100 mg of merope e a final volume of 10 mL wit	to 500 mg vial to r n to 1g vial to make enem) of the above th a final concentra to 500 mg vial to r	make a 50 mg/mL see a 50 mg/mL solution and add tion of 10 mg/mL.	solution OR tion. 8 mL sodium chloride				
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Maximum dose Total cumulative dose Route	IV inf Infan Add 9 Add 9 Add 1 FURT Draw Make Infan Add 9 Add 1 FURT Draw	fusion. fusion. fusion. 19.6 mL of water for injection 19.1 mL of water for injection THER DILUTE v up 2 mL (100 mg of merope e a final volume of 10 mL wit 19.6 mL of water for injection 19.1 mL of water for injection THER DILUTE v up 4 mL (200 mg of merope	to 500 mg vial to r n to 1g vial to make enem) of the above th a final concentra to 500 mg vial to r n to 1g vial to make	make a 50 mg/mL solutes a 50 mg/mL solution and add tion of 10 mg/mL. make a 50 mg/mL solutes a 50 mg/mL solutes a 50 mg/mL solutes a solution and add	solution OR tion. 8 mL sodium chloride solution OR tion.	0.9% to			
Maximum dose Total cumulative dose Route Preparation	IV inf Infan Add 9 Add 1 FURT Draw make Infan Add 9 Add 1 FURT Draw make Add 1	fusion. fusion. fusion. nts <1 kg 9.6 mL of water for injection 19.1 mL of water for injection ther DILUTE v up 2 mL (100 mg of merope e a final volume of 10 mL wite nts≥1 kg or fluid restricted 9.6 mL of water for injection 19.1 mL of water for injection ther DILUTE v up 4 mL (200 mg of merope e a final volume of 10 mL wite	to 500 mg vial to r n to 1g vial to make enem) of the above th a final concentra to 500 mg vial to r n to 1g vial to make	make a 50 mg/mL solutes a 50 mg/mL solution and add tion of 10 mg/mL. make a 50 mg/mL solutes a 50 mg/mL solutes a 50 mg/mL solutes a solution and add	solution OR tion. 8 mL sodium chloride solution OR tion.	0.9% to			
Maximum dose Total cumulative dose Route	IV inf Infan Add 9 Add 1 FURT Draw make Infan Add 9 Add 1 FURT Draw make IV inf	fusion. fusion. fusion. 19.6 mL of water for injection 19.1 mL of water for injection THER DILUTE v up 2 mL (100 mg of merope e a final volume of 10 mL wit 19.6 mL of water for injection 19.1 mL of water for injection THER DILUTE v up 4 mL (200 mg of merope	to 500 mg vial to r n to 1g vial to make enem) of the above th a final concentra to 500 mg vial to r n to 1g vial to make enem) of the above th a concentration of	make a 50 mg/mL see a 50 mg/mL solution and add tion of 10 mg/mL. make a 50 mg/mL solution and add a 50 mg/mL solution and add of 20 mg/mL.	solution OR tion. 8 mL sodium chloride solution OR tion. 6 mL sodium chloride	0.9% to			

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	Liver function.		
	Electrolytes		
Contraindications	Hypersensitivity to penicillins, cephalosporins and carbapenems.		
Precautions	Colitis-due to risk of pseudomembranous colitis.		
	Renal impairment.		
Drug interactions	Sodium valproate- meropenem may result in clinically significant reduction in concentration of sodium		
	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,		
la compatibility	gentamicin, heparin sodium, metronidazole. Fluids: Mannitol 10%		
Incompatibility	Fidias: Mannitor 10%		
	Y-site: Dolasetron, ketamine, zidovudine.		
Stability	Use immediately after preparation.		
oud.oy	Diluted solutions are potentially unstable, particularly glucose containing solutions and should be		
	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 Enterobacteriaceae bacteraemia. A systematic review of carbapenems for the		
	treatment of patients with extended-spectrum β-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). ²		
	0.23 1.13) of empirical (tilt 0.31, 33% ci 0.00 1.23) freatment (LOE 1, GON C).		
	A retrospective case series of 100 neonates infected by extended-spectrum beta-lactamase-producing		
	Klebsiella 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). ³		
	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 gram-		
	negative 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]. ⁵		
	Pharmacokinetics:		
	Meropenem is primarily excreted via the kidneys.		
	Meropenem clearance is influenced by serum creatinine and postmenstrual age in neonates. ²		
	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. ⁶ However, a longer infusion may have greater efficacy. ⁵		
	There is a knowledge gap in pharmacokinetic (PK) studies of neonates with renal impairment. ^{2,3}		
	However, dose adjustment for renal failure may not be appropriate in cases where severe sepsis is		
	probably responsible for acute renal failure [expert opinion].		
	Pose:		
	Dose: 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		
	and the terms and the market 2 32 weeks of and then the days, and so mg/ng every o hours in		

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	infants ≥ 32 weeks GA and PNA ≥ 14 days to achieve therapeutic concentrations in infants with suspected intra-abdominal infections. ⁴		
Practice points			
References	1. Pacifici GM, Allegaert K. Clinical pharmacology of carbapenems in neonates. J Chemother 2014;26(2):67–73.		
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