# **Newborn use only**

Indication Action	The administration of antibiotics within 1 hour of the identification of sepsis is recommended.(1) The Antimicrobial Stewardship Team has listed this drug under the following categories: Unrestricted – duration up to 48 hours and restricted for duration > 48 hours Aminoglycosides can be inactivated by penicillin and cephalosporin antibiotics. As commonly coprescribed, where feasible, give at separate sites or separate the administration time of the antibiotics. Unregistered products from overseas available during shortages may contain preservatives. Treatment of gram-negative infections. Bactericidal agent that acts by inhibiting protein synthesis in susceptible bacteria.					
Drug type	Aminoglycoside antibiotic					
Trade name	DBL gentamicin, Gentamicin BP (Pfizer)					
Presentation	10 mg/mL ampoule – paediatric strength 80 mg/2 mL ampoule – adult strength NOTE: SAS product may be considered in the event of a shortage. Consult the local pharmacy.					
Dose	Dose: 5 mg/kg as follows: (2-5)			, ,		
	Corrected Gestational Age/Postmenstrual Age*	Route	Dosing interval	Drug concentration to be performed at:		
	< 30 <sup>+0</sup> weeks*	IV/IM	48 hourly	22 hours after the 2 <sup>nd</sup> dose		
	30 <sup>+0</sup> –34 <sup>+6</sup> weeks*	IV/IM	36 hourly	22 hours after the 2 <sup>nd</sup> dose		
	≥ 35 <sup>+0</sup> weeks*	IV/IM	24 hourly	22 hours after the 2 <sup>nd</sup> dose		
	*Concurrent cyclo-oxygenase inhibitors (indomethacin or ibuprofen) (6-8)	IV/IM	Extend dosing interval by 12 hours Example: 48 hourly to 60 hourly			
	Therapeutic hypothermia (9-13)	IV/IM	36 hourly	Trough concentrations prior to every dose		
	Subsequent dose interval is based on a gentamicin concentration at 22 hours after the administration of the 2 <sup>nd</sup> dose as indicated in the table below.(3, 4)					
	22-hour Gentamicin concentration*		Interval			
	≤ 1.2 mg/L		Every 24 hours after pre			
	1.3 mg/L – 2.6 mg/L 2.7 mg/L – 3.5 mg/L		Every 36 hours after previous dose			
			Every 48 hours after previous dose			
	≥ 3.6 mg/L		Hold dose, repeat concentration 24 hours later			
	*Different to trough concentration performed prior to next dose – Refer to dose adjustment section.  Gentamicin monitoring is required ONCE only, except when the duration of gentamicin therapy is greater than 7 days or with the conditions described in dose adjustment and monitoring section.					
Dose adjustment	Therapeutic hypothermia –36 hourly interval(9-13). Measure trough concentrations before every dose.  ECMO - Renal dysfunction is the main determinant. Measure trough concentration before 2 <sup>nd</sup> dose.(14)  Renal impairment – Measure trough concentration before every dose.  Hepatic impairment – No specific dose adjustment.					
Maximum dose						

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Total cumulative				
dose	IV			
Route	IM – only if IV access is not available.			
Preparation	10mg/mL – paediatric strength			
rieparation	Draw up 1mL (10mg) gentamicin and add to 4mL of sodium chloride 0.9% to make a final volume			
	of 5mL with a concentration of 2mg/mL solution.			
	80mg/2 mL – adult strength			
	Draw up 1mL (40mg) gentamicin and add to 19mL of sodium chloride 0.9% to make a final			
	volume of 20mL with a concentration of 2mg/mL solution.			
Administration	IV - Inject slowly over 5 minutes as an IV injection.(15)			
, (4	IM- only given when IV route is not available as the IM absorption is variable. Administer			
	required dose undiluted, deeply into anterolateral thigh muscle.			
Monitoring	Urine output, urine analysis, blood urea, nitrogen and creatinine			
Womtoning	Monitor for anaphylaxis			
	Trough concentrations – Target trough concentration: <2 mg/L. Repeat trough concentrations			
	are not required routinely unless: (4)			
	(1) duration of therapy is ≥ 7 days – In this scenario, prior to dose on day 7 and then weekly			
	thereafter.			
	(2) renal impairment or perinatal hypoxia with Apgar <5 at 5 minutes and/or concomitant use			
	of other nephrotoxic agents or therapeutic hypothermia In these scenarios, perform trough			
	concentration prior to every dose.			
	concentration prior to every asser			
	If trough concentration ≥2 mg/L, withhold the dose, repeat trough concentrations before the			
	subsequent dosing and discuss with infectious disease specialist/clinical microbiologist for either			
	extended dosing interval or alternate antibiotic.			
	Peak concentrations - Not required routinely. Target peak concentrations: 5-12 mg/L. Peak			
	concentration should be drawn at 30 minutes post dose.			
Contraindications	Hypersensitivity to aminoglycosides			
Precautions	CAUTION in patients with pre-existing renal impairment, auditory or vestibular impairment,			
	hypocalcaemia, depressed neuromuscular transmission.			
Drug interactions	Gentamicin should not be mixed with penicillins or cephalosporins as inactivation occurs. (15)			
<b>.</b>	Ensure line is adequately flushed between antibiotics and if possible, stagger the time of			
	administration of each drug so that they are separated by several hours.			
	Avoid use with other potent diuretics, neurotoxic, nephrotoxic and neuromuscular blocking			
	Avoid use with other potent diuretics, neurotoxic, nephrotoxic and neuromuscular blocking agents.(16)			
Adverse reactions				
Adverse reactions	agents.(16)			
Adverse reactions	agents.(16)  Toxicity is rare in the newborn but can include:			
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Adverse reactions	agents.(16)  Toxicity is rare in the newborn but can include:  1. Nephrotoxicity- Associated with excessive accumulation of gentamicin. The initial symptoms may be due to renal tubular concentrating defect. These include excessive losses of sodium, calcium and magnesium. This may progress to proteinuria, increased urea, oliguria, increased serum creatinine. Renal impairment is usually reversible.  2. Ototoxicity.  Primarily vestibular but also auditory toxicity. Associated with excessive accumulation of gentamicin and duration of therapy. Effects often irreversible.  3. Neuromuscular blockade- Muscular paralysis and respiratory failure may occur particularly when used with other neuromuscular blockers such as pancuronium.  4. Hypersensitivity-			

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F		
Compatibility	Fluids: Glucose 5%, glucose 10%, Hartmann's, sodium chloride 0.9%, Ringer's (15)	
	Y-Site: Amino acid solutions, amifostine, amiodarone, anidulafungin, atracurium, aztreonam, bivalirudin, calcium chloride, calcium gluconate, caspofungin, ciprofloxacin, cisatracurium,	
	clindamycin, dexmedetomidine, digoxin, dobutamine, esmolol, fentanyl, fluconazole, foscarnet,	
	granisetron, hydromorphone, labetalol, linezolid, magnesium sulfate, meropenem,	
	methylprednisolone, metronidazole, midazolam, morphine sulfate, , pancuronium, pethidine,	
	phenobarbital sodium, potassium chloride, remifentanil, rocuronium, suxamethonium,	
	tigecycline, vancomycin, vecuronium, zidovudine.	
Incompatibility	Fluids: Fat emulsions.	
	Y-site: Azathioprine, azithromycin, chloramphenicol, dexamethasone, flucloxacillin, folic acid,	
	frusemide, ganciclovir, heparin sodium, indomethacin, pentamidine, propofol, teicoplanin.	
a. 1 !!!:	Note: Do not mix together with penicillins or cephalosporins.	
Stability	Administer immediately, discard unused portion.	
Storage	Protect from light. Store below 25°C	
Excipients	DBL Gentamicin: Disodium edetate	
	Pfizer Gentamicin: Disodium edetate, sodium hydroxide, sulfuric acid.	
Special comments		
Evidence	Efficacy	
	Extended interval dosing for gentamicin in neonates provides a superior pharmacokinetic profile	
	compared to multiple doses a day dosing. However, there is insufficient evidence to conclude	
	whether a 'once a day' or a 'multiple doses a day' regimen of gentamicin is clinically superior in	
	treating proven neonatal sepsis. (17, 18) (Rao SC 2016, Nestaas E 2005)	
	Current dosing recommendations are based on 4 prospective observational studies using	
	extended-interval dosing interval with a single drug concentration at 22 hours after the first	
	dose.(2-5) Three of them were consecutive Canadian studies. First of the studies evaluated the	
	extended interval dosing (EID) regimen in neonates ≤28-week gestation. The dosing interval was	
	based on a 22 h level after the first dose of 5mg/kg. All neonates, except one, achieved	
	therapeutic peak and trough levels. Based on the 22 h level, dosing interval was 36 h in 61% of	
	neonates and 48 h in 39% of neonates. In their second prospective, observational study, similar	
	findings were noted in 104 neonates ≤7 days of life, gestational age 23 weeks to full term.	
	Appropriate peak and trough concentrations were attained in all neonates. A third prospective	
	observational study by the group assessed extended-interval dosing of gentamicin in neonates	
	>7 days old and found appropriate peak and trough concentrations in all neonates. (2-4) Fourth	
	observational study by Matinkova et al, in which 4 mg/kg/dose was given at various intervals	
	based on gestational age groups (<34 weeks-48 hourly; 34-38 weeks – 36 hourly; >38 weeks – 24	
	hourly). The initial dose of gentamicin 4mg/kg during the first week of life was high enough to	
	reach bactericidal Cmax within 6–10mg/L. However, C <sub>max</sub> <6 mg/L occurred in 13% of neonates.	
	The inter-dose interval modified according to the recommendation resulted in C <sub>trough</sub> values	
	within the target range of 0.5–2.0mg/L in all but 2 neonates.(5)	
	Patients who have early (I-hr post-infusion) peak plasma aminoglycoside levels that are >5	
	ug/mL for gentamicin and tobramycin and >20 ug/ml for amikacin are less likely to die from	
	gram-negative bacteraemia. Moore et al reported a 2.4% mortality rate in adults who achieved	
	1-hour post-infusion gentamicin or tobramycin peak concentrations above 5 µg/mL. Mortality	
	rate increased to 20.9% for patients failing to achieve peak concentrations above 5 μg/mL within	
	24–48 hours of starting therapy. (19, 20)	
	Therapeutic hypothermia (TH): Gentamicin clearance is decreased in neonates receiving	
	hypothermia treatment. Modified gentamicin dosing regimens are required to avoid potential	
	toxicity related to higher concentrations.(13)	
	<b>ECMO</b> : During ECMO, gentamicin has an increased volume of distribution (Vd), and decreased	
	clearance (CI), leading to a prolonged elimination half-life. The renal dysfunction, which is a	
	common multifactorial condition during ECMO, can be considered as the main determinant of	

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the prolonged elimination half-life of gentamicin. Given the concentration dependent antimicrobial activity of aminoglycosides, it is recommended to perform therapeutic drug monitoring (TDM) to ensure adequate antimicrobial exposure. (14)

Cyclo-oxygenase inhibitors: Renal drug clearance of aminoglycosides is lower in infants on cyclo-oxygenase inhibitors. (6-8)

#### Safety

Ototoxicity: There is no clear association between peak or trough levels and ototoxicity in neonates. (21-23) The chance of gentamicin ototoxicity is reported to be greater in those who receive the drug for a longer duration.(21)

Nephrotoxicity: Nephrotoxicity does not seem to be related to peak or trough levels and more related to drug concentration and longer duration. (24) Among neonates with PDA and receiving gentamicin, non-steroid anti-inflammatory drugs (ibuprofen, indomethacin) therapy increases the risk of acute kidney injury.(25)

MT-RNR1 genotype: MT-RNR1 gene mutation is one of the common causes of hereditary hearing loss, particularly in Asian population. In individuals who carry mutations in MT-RNR1 gene, a single dose of gentamicin can result in hearing loss. (26, 27)

Intraventricular antibiotics: In infants with meningitis and ventriculitis, intraventricular antibiotics in combination resulted in a three-fold increase in mortality compared to standard treatment with intravenous antibiotics alone and should be avoided. (28)

#### **Pharmacokinetics**

Aminoglycosides display concentration-dependent killing, suggesting higher peaks provide greater efficacy.(29, 30) While a peak aminoglycoside concentration to minimum inhibitory concentration (MIC) ratio of 8–10:1 is considered ideal, based on the usual MICs of Escherichia coli (range 0.25-1 mg/L) a peak of at least 5 mg/L has a high likelihood of being effective.(4, 30) Aminoglycosides display a post-antibiotic effect whereby bacterial growth is suppressed despite negligible drug concentrations. (31) Aminoglycosides have poor CNS penetration when administered intravenously.(32)

### **Practice points**

### Dose

There is insufficient evidence whether a 'once a day' regimen of gentamicin is optimal in treating proven neonatal sepsis, however, pharmacokinetic data suggests 'once a day' gentamicin regimens are superior to a 'multiple doses a day' regimens.(17) (LOE I, GOR B)

The recommended dose regimen in this formulary is a pragmatic adaptation of the dosing used in 4 prospective observational studies.(2-5) (LOE III-3, GOR B)

### Dose adjustment

An increased dosing interval is recommended in therapeutic hypothermia. (9-13) (LOE IV, GOR B) An increased dosing interval is recommended in infants on cyclo-oxygenase inhibitors.(6) (LOE IV, GOR B)

### Monitoring

The evidence suggests a serum gentamicin concentration performed 22 hours after the 1st dose is useful to guide dosing intervals. (2-4)(LOE III-3, GOR B). However, in daily practice, gentamicin is most often discontinued within 36-48 hours of commencement (once the neonate is deemed no longer at risk of sepsis and septic screen remain negative). Therefore, measurement of drug concentrations is recommended only after the 2nd dose to limit the burden of blood sampling. (ANMF consensus recommendation).

Subsequent concentrations are not routinely required. (2-4) (LOE III-3, GOR B)

Routine peak concentrations are not necessary as high dose extended interval dosing regimens are able to achieve target peak concentrations in the majority of infants (2-4, 17, 18) (LOE III-3, GOR B)

Consider performing peak concentrations if there is poor clinical response in gram negative infections, oedema or macrosomia.(5) (LOE IV, GOR C).

A peak concentration, if required, can be performed after the 2nd or 3rd dose. (29)

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### **Newborn use only**

Target peak concentrations of 5–12 mg/L. (17-19, 29) (LOE IV, GOR C)

Target trough concentrations of < 2 mg/L to reduce risk of ototoxicity and nephrotoxicity. (33, 34) (LOE IV, GOR C - adult)

Duration of therapy  $\geq$  7 days – Perform trough concentration prior to dose on day 7 and then weekly thereafter. (4, 35) (LOE IV, GOR B)

Perinatal hypoxia – Perform trough concentrations prior to every dose. (4, 35) (LOE IV, GOR B) Renal impairment – Perform trough concentrations prior to every dose. (4, 35) (LOE IV, GOR B) Concomitant use of other nephrotoxic agents – Perform trough concentrations prior to every dose. (4, 35) (LOE IV, GOR B)

ECMO – Perform trough concentration before 2<sup>nd</sup> dose.(14) (LOE IV, GOR B)

#### Route

Intraventricular antibiotics are associated with increased mortality and should be avoided.(28) (LOE II, GOR B)

#### General

Aim to minimise aminoglycoside toxicity by (1) avoiding gentamicin to patients at elevated risk (i.e. on indomethacin, history of hypoxia and/or significant renal dysfunction), (2) minimising the duration of treatment and (3) prescribing a dose in a way that minimizes risk (i.e. EID with dose adjustment as necessary). (ANMF consensus recommendations)

#### References

- 1. Rhodes A, Evans LE, Alhazzani W, Levy MM, Antonelli M, Ferrer R, et al. Surviving Sepsis Campaign: International Guidelines for Management of Sepsis and Septic Shock: 2016. Intensive care medicine. 2017;43(3):304-77.
- Alshaikh B, Dersch-Mills D, Taylor R, Akierman AR, Yusuf K. Extended interval dosing of gentamicin in premature neonates ≤ 28-week gestation. Acta Paediatrica. 2012;101(11):1134-9.
- 3. Dersch-Mills D, Akierman A, Alshaikh B, Yusuf K. Validation of a Dosage Individualization Table for Extended-Interval Gentamicin in Neonates. Annals of Pharmacotherapy. 2012;46(7-8):935-42.
- 4. Dersch-Mills D, Akierman A, Alshaikh B, Sundaram A, Yusuf K. Performance of a dosage individualization table for extended interval gentamicin in neonates beyond the first week of life. The Journal of Maternal-Fetal & Neonatal Medicine. 2016;29(9):1451-6.
- 5. Martínková J, Pokorná P, Záhora J, Chládek J, Vobruba V, Selke-Krulichová I, et al. Tolerability and outcomes of kinetically guided therapy with gentamicin in critically ill neonates during the first week of life: an open-label, prospective study. Clinical therapeutics. 2010;32(14):2400-14.
- 6. Allegaert K. The impact of ibuprofen or indomethacin on renal drug clearance in neonates. The Journal of Maternal-Fetal & Neonatal Medicine. 2009;22(sup3):88-91.
- 7. Smits A, De Cock RFW, Allegaert K, Vanhaesebrouck S, Danhof M, Knibbe CAJ. Prospective Evaluation of a Model-Based Dosing Regimen for Amikacin in Preterm and Term Neonates in Clinical Practice. Antimicrobial Agents and Chemotherapy. 2015;59(10):6344.
- 8. Smits A, Kulo A, Van Den Anker J, Allegaert K. The amikacin research program: a stepwise approach to validate dosing regimens in neonates. 2016:1-10.
- 9. Frymoyer A, Lee S, Bonifacio SL, Meng L, Lucas SS, Guglielmo BJ, et al. Every 36-h gentamicin dosing in neonates with hypoxic–ischemic encephalopathy receiving hypothermia. Journal of Perinatology. 2013;33(10):778-82.
- 10. Bijleveld YA, De Haan TR, Van Der Lee HJH, Groenendaal F, Dijk PH, Van Heijst A, et al. Altered gentamicin pharmacokinetics in term neonates undergoing controlled hypothermia. British Journal of Clinical Pharmacology. 2016;81(6):1067-77.
- 11. Lutz IC, Allegaert K, de Hoon JN, Marynissen H. Pharmacokinetics during therapeutic hypothermia for neonatal hypoxic ischaemic encephalopathy: a literature review. BMJ Paediatr Open. 2020;4(1):e000685-e.

ANMF consensus group Gentamicin Page 5 of 7

### Newborn use only

- 12. Cristea S, Smits A, Kulo A, Knibbe CAJ, van Weissenbruch M, Krekels EHJ, et al. Amikacin Pharmacokinetics To Optimize Dosing in Neonates with Perinatal Asphyxia Treated with Hypothermia. Antimicrobial Agents and Chemotherapy. 2017;61(12):e01282-17.
- 13. Choi D, Park J, Lee S, An S. Effect of hypothermia treatment on gentamicin pharmacokinetics in neonates with hypoxic-ischaemic encephalopathy: A systematic review and meta-analysis. Journal of Clinical Pharmacy and Therapeutics. 2018;43(4):484-92.
- 14. Raffaeli G, Pokorna P, Allegaert K, Mosca F, Cavallaro G, Wildschut E, et al. Drug disposition and pharmacotherapy in neonatal ECMO: from fragmented data to integrated knowledge. Frontiers in pediatrics. 2019;7:360.
- 15. Gentamicin. Australian Injectable drug handbook 8th edition. (18 Spetember 2020).
- 16. . MIMS online. (14 September 2020).
- 17. Rao SC, Srinivasjois R, Moon K. One dose per day compared to multiple doses per day of gentamicin for treatment of suspected or proven sepsis in neonates. Cochrane Database of Systematic Reviews. 2016.
- 18. Nestaas E. Aminoglycoside extended interval dosing in neonates is safe and effective: a meta-analysis. 2005;90(4):F294-f300.
- 19. Moore RD, Smith CR, Lietman PS. The association of aminoglycoside plasma levels with mortality in patients with gram-negative bacteremia. Journal of infectious diseases. 1984;149(3):443-8.
- 20. Moore RD, Smith CR, Lietman PS. Association of aminoglycoside plasma levels with therapeutic outcome in gram-negative pneumonia. The American journal of medicine. 1984;77(4):657-62.
- 21. Kent A, Turner MA, Sharland M, Heath PT. Aminoglycoside toxicity in neonates: something to worry about? Expert Review of Anti-infective Therapy. 2014;12(3):319-31.
- 22. Setiabudy R, Suwento R, Rundjan L, Yasin FH, Louisa M, Dwijayanti A, et al. Lack of a relationship between the serum concentration of aminoglycosides and ototoxicity in neonates. Int J Clin Pharmacol Ther. 2013;51(5):401-6.
- 23. El-Barbary MN, Ismail RIH, Ibrahim AAA. Gentamicin extended interval regimen and ototoxicity in neonates. 2015;79(8):1294-8.
- 24. Quiros Y, Vicente-Vicente L, Morales AI, López-Novoa JM, López-Hernández FJ. An integrative overview on the mechanisms underlying the renal tubular cytotoxicity of gentamicin. Toxicological sciences. 2011;119(2):245-56.
- 25. Constance JE, Reith D, Ward R, Balch A, Stockmann C, Korgenski EK, et al. Risk of nonsteroidal anti-inflammatory drug-associated renal dysfunction among neonates diagnosed with patent ductus arteriosus and treated with gentamicin. Journal of Perinatology. 2017;37(10):1093-102.
- 26. Wang X, Hong Y, Cai P, Tang N, Chen Y, Yan T, et al. Rapid and Reliable Detection of Nonsyndromic Hearing Loss Mutations by Multicolor Melting Curve Analysis. Scientific Reports. 2017;7(1):42894.
- 27. Dean L. Gentamicin Therapy and MT-RNR1 Genotype: National Center for Biotechnology Information (US), Bethesda (MD); 2012 2012.
- 28. Shah SS, Ohlsson A, Shah VS. Intraventricular antibiotics for bacterial meningitis in neonates. Cochrane Database of Systematic Reviews. 2012(7).
- 29. Touw DJ, Westerman EM, Sprij AJ. Therapeutic Drug Monitoring of Aminoglycosides in Neonates. Clinical Pharmacokinetics. 2009;48(2):71-88.
- 30. Kearns GL, Abdel-Rahman SM, Alander SW, Blowey DL, Leeder JS, Kauffman RE.

  Developmental pharmacology—drug disposition, action, and therapy in infants and children.

  New England Journal of Medicine. 2003;349(12):1157-67.
- 31. Lacy MK, Nicolau DP, Nightingale CH, Quintiliani R. The pharmacodynamics of aminoglycosides. Clinical infectious diseases. 1998;27(1):23-7.

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- 32. Sullins AK, Abdel-Rahman SM. Pharmacokinetics of antibacterial agents in the CSF of children and adolescents. Pediatric Drugs. 2013;15(2):93-117.
- 33. Dahlgren JG, Anderson ET, Hewitt WL. Gentamicin blood levels: a guide to nephrotoxicity. Antimicrobial agents and chemotherapy. 1975;8(1):58-62.
- 34. Goodman EL, Van Gelder J, Holmes R, Hull AR, Sanford JP. Prospective comparative study of variable dosage and variable frequency regimens for administration of gentamicin. Antimicrobial Agents and Chemotherapy. 1975;8(4):434-8.
- 35. de Hoog M, Mouton JW, van den Anker JN, editors. New dosing strategies for antibacterial agents in the neonate. Seminars in Fetal and Neonatal Medicine; 2005: Elsevier.

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