# Current Opinion in Infectious Diseases Vancomycin Toxicity in Neonates: A Review of the Evidence --Manuscript Draft--

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# Vancomycin Toxicity in Neonates: A Review of the Evidence

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### **ABSTRACT**

# Purpose of review:

Vancomycin is a first-line agent in the treatment of serious Gram-positive infections in the neonatal population. The published evidence on vancomycin toxicity in neonates is limited. This review summarises pre-clinical studies and clinical trials describing vancomycin toxicity. We discuss proposed pathophysiology and summarise evidence supporting dose-response relationships, genetic and environmental determinants, and consider future research required to further define vancomycin toxicity.

# Recent findings:

Current dosing regimens for vancomycin result in sub-therapeutic levels in a large proportion of patients. Higher daily doses have been proposed, which have led to concerns regarding increased toxicity. Nephrotoxicity occurs in 1-9% of neonates receiving currently recommended doses. The incidence is highest in those receiving concomitant nephrotoxic drugs. Vancomycin-associated ototoxicity is rare in patients of all ages. Exposure-toxicity relationships in relation to nephro- and ototoxicity have not been clearly defined in neonates receiving vancomycin.

# **Summary**:

Current evidence supports the favourable safety profile of vancomycin in neonates. Further studies that address safety concerns relating to high-dose intermittent dosing regimens are needed. Such studies must include robust and standardised definitions of renal and hearing impairment, and include follow-up of sufficient length to establish the long-term implications of experimental findings.

### **KEYWORDS**

Vancomycin, toxicity, neonates, renal impairment, hearing loss

### INTRODUCTION

Vancomycin is a glycopeptide bactericidal antibiotic that disrupts cell wall synthesis in Gram-positive bacteria. (1) Impurities from early fermentation processes were associated with significant toxicity when the drug was first introduced in the early 1950s. The drug's poor safety profile, along with its thick brown appearance, led to the disparaging nickname 'Mississippi mud'. Refined purification methods improved the safety profile of vancomycin, and Food and Drug Administration approval was granted in 1958. Despite this, safety concerns lingered, leading to the drug's limited use. In recent decades clinical use has increased, however, owing to the rising incidence of infections caused by methicillin-resistant *Staphylococcus aureus* and other resistant Gram-positive pathogens that are susceptible to vancomycin. (2)

Vancomycin dosing strategies vary greatly and are generally based on a combination of post-menstrual age (PMA), post-natal age (PNA), weight and/or renal function (see table 1). Therapeutic drug monitoring (TDM) is widely advocated. Currently clinical guidelines recommend target trough concentrations of 10-15 µg/mL. (3, 4) However, recent studies suggest the need for target trough levels >15 µg/mL, based on the concern that lower trough concentrations may be selective for hetero-resistance. (5-7) In adults, higher daily doses of vancomycin (15 mg/kg 6 hourly) have yet to receive regulatory approval but have been proposed in recent consensus documents published by the Infectious Diseases Society of America, American Society of Health-System Pharmacists, and the Society of Infectious Diseases Pharmacists. (8-10) The use of higher vancomycin doses has raised concerns regarding the potentially increased risk

of toxicity. The significance of these concerns in paediatric populations is unclear. Here we summarise the current evidence relating to vancomycin toxicity including proposed pathophysiology, dose-toxicity relationships, genetic and environmental factors and future research needs. Studies pertaining to vancomycin-induced toxicity between March 1950 and March 2015 were identified from PubMed, Embase, Cochrane Controlled Trial Registry, and Medline databases. The following search terms were used "vancomycin", "toxicity", "adverse effects', "nephrotoxicity", "kidney disease", "renal impairment", "ototoxicity", "hearing loss", "neonate", "infant", "child", "paediatrics".

# **NEPHROTOXICITY**

# **Quantification of Nephrotoxicity**

Kidney function is difficult to define in the first month of life when renal physiology and fluid balance evolve rapidly. Serum creatinine (SCr) concentrations reflect maternal renal function in the first 48 hours after birth. (11, 12) In addition, preterm infants experience a rise in SCr due to tubular reabsorption, making this an imprecise proxy for glomerular filtration. (13) Clinical reports also often omit details of the laboratory methods used to quantify SCr (Jaffé colorimetry, enzymatic quantification, or isotope dilution mass spectroscopy), which can have an inter-test variability of up to 25% in neonates. (14, 15) Despite these limitations most studies define renal impairment as an increase in SCr. (16-19) Definitions for children and neonates have been reported based on several classifications systems, all of which were originally developed and validated in adults. (20, 21) Modified neonatal criteria exclude urine output or use a higher cut-off (< 1.0 mL/kg/h) in order to account for the non-oliguric

renal dysfunction that occurs in neonates due to fluid redistribution and impaired absorption in the immature renal tubule. (22, 23)

### **Pre-Clinical Studies**

Murine models have demonstrated that vancomycin has an affinity for biological membranes and accumulates in renal tissue. (24, 25) The energy-dependent transport mechanisms within the tubular epithelium render the kidney highly susceptible to toxin-induced cellular injury. Vancomycin enhances cellular ATP concentrations and stimulate oxygen consumption, resulting in oxidative phosphorylation. (26-31) However, oxidative stress of sufficient magnitude to cause clinically detectable renal impairment has not been demonstrated in animals, even at supra-therapeutic doses. (32) Histopathological changes and biochemical evidence of nephrotoxicity have been observed in an uninfected murine model in which high-dose vancomycin was co-administered with tobramycin. (25)

# **Clinical Studies**

The linear relationship between SCr and vancomycin exposure reflects the drugs primary route of elimination through renal excretion. As a result, a decrease in renal function from any cause will increase serum vancomycin concentrations, a fact that confounds the establishment of exposure-toxicity relationships. (33) Many studies that describe vancomycin-associated nephrotoxicity are conducted in patients otherwise at risk of renal impairment, such as those requiring intensive care and frequently include those receiving other potentially nephrotoxic drugs (e.g. NSAIDs, diuretics and aminoglycoside antibiotics) . However, irrespective of the difficulty in defining causal relationships, acute kidney injury from any cause is associated with a

significantly increased risk of mortality, hospital length of stay, and cost, even after adjustment for age, gender, chronic kidney disease, and co-morbidities at admission.

(34)

### Adult Studies

Nephrotoxicity in patients receiving vancomycin has been more systematically reported in adults than in neonates and children. The reported incidence of nephrotoxicity in adults receiving vancomycin is 10-15%. (35) In a recent metaanalysis, van Hal, et al. reviewed fifteen studies in which vancomycin was administered via intermittent infusion to adults >18 years of age. Randomised trials and observational studies in which data on renal function and trough vancomycin concentrations were available were included. The analysis found trough concentrations >15 µg/mL to be independently associated with an increased odds of nephrotoxicity (OR, 2.67; 95% CI, 1.95-3.65), and also suggested an incremental increase in nephrotoxicity risk with vancomycin administration beyond seven days. (36) The majority of reported cases were self resolving, with prolonged impairment requiring short-term dialysis occurring in 3% of cases. Similarly, a retrospective cohort study of 246 adults receiving intermittent-dosing of vancomycin for durations of >48 hours found toxicity to be correlated with exposure (based on dose). Patients receiving  $\ge 4$  g daily (n=26) had a significantly higher incidence of nephrotoxicity compared with those receiving <4 g daily (34.6% vs. 10.9%; p = 0.001). (18)

Concomitant aminoglycoside use has consistently been shown to increase the risk of nephrotoxicity in adults receiving vancomycin. In 2007, Fowler, *et al* reported a seminal open-label randomised trial of daptomycin versus either anti-staphylococcal

penicillin or vancomycin plus low-dose gentamicin in 236 patients with *S. aureus* bacteremia. The highest proportion of renal impairment occurred in patients receiving vancomycin plus gentamicin (20.4%), although this was not significantly higher than with penicillin plus gentamicin (18.6%).

Furthermore, the study found that the highest incidence of renal impairment in patients receiving vancomycin occurred later in treatment, peaking at 14-28 days. (37)

### Paediatric & Neonatal Studies

Vancomycin administration with intermittent dosing and in accordance with published guidance has been shown to result in sub-therapeutic trough concentrations in up to half of treated neonates. (4) Furthermore, antimicrobial point prevalence surveys that describe prescribing practices have demonstrated that significant variation in prescribing of vancomycin in neonates exists in clinical practice, with a large proportion of neonates receiving daily doses of vancomycin that are significantly below even the most conservative current recommendations. (38) Table 2 outlines the clinical studies that have reported nephrotoxicity associated with vancomycin use in neonates and infants. The majority of studies identified are heterogeneous observational or opportunistic pharmacokinetic studies based on routine TDM and are insufficiently powered to detect toxicity.

As with adults, cases of renal impairment are more frequently reported in neonates and children receiving concomitant nephrotoxic drugs. (19, 35) Eight of ten studies describing concomitant nephrotoxic drugs alongside vancomycin therapy demonstrated evidence of renal impairment, which was mild and transient in all reported cases (see table 2). Other reports have not found vancomycin to be an

independent predictor of nephrotoxicity. Constance, *et al*, for example, found no significant difference in the proportion of neonates developing nephrotoxicity in those receiving vancomycin plus gentamicin (12/533, 2.2%) versus those receiving gentamicin alone (7/533, 1.3%). Logistic regression demonstrated that while positive blood culture, low birth weight, patent ductus arteriosus, concomitant non-steroidal anti-inflammatory drug (NSAID) use, and illness severity were all independent risk factors for nephrotoxicity, vancomycin in conjunction with gentamicin was not. (23) Six studies report rates of nephrotoxicity in neonates receiving vancomycin alone. A transient rise in SCr, microproteinuria, and elevated NAG were reported in only three, two and one patient, respectively (see table 2). (51, 54, 59-61)

Exposure-toxicity relationships have not been clearly defined for vancomycin in neonates. Two case reports have described clinical outcomes in four neonates receiving accidental overdoses of up to 10-fold the maximum recommended dose.

(51, 59) All four patients developed transient renal impairment, though all had normal renal function at 6 months. It is noteworthy that these cases all involved single or brief exposures that were managed with immediate vancomycin withdrawal, and so these findings do not address the potential risk of cumulative exposure.

A number of recent studies have described the administration of vancomycin via continuous infusion in neonates. Continuous infusion has the theoretical advantage of maintaining constant plasma concentrations, meaning that overall drug exposure can be increased without a rise in peak concentrations. Studies investigating continuous vancomycin infusions in neonates have, to date, been based on studies defined toxicity differently and involved small patient cohorts that received varied dosing regimens. A loading dose of 7-15 mg/kg was given in three studies. Collectively, the

results of these studies suggest that continuous infusion may result in a higher proportion of patients achieving target concentrations between 15-20  $\mu$ g/mL. Reported rates of nephrotoxicity do not differ significantly from those in patients receiving intermittent dosing, and there is currently no evidence to suggest the use of a loading dose increases the risk of nephrotoxicity. (3, 45, 47, 60) Continuous infusions may, however, be impractical in the neonatal population where venous access if limited, and may lead to periods without effective antibiotic cover if access is lost. There have, as yet, been no systematic studies comparing the safety and efficacy of continuous infusion with high-dose intermittent regimens that specifically target trough concentrations of >15  $\mu$ g/mL.

### **OTOTOXICITY**

# **Quantification of Ototoxicity in Neonates**

The assessment of hearing in infants is challenging. Behavioural tests (e.g. visual reinforcement audiometry) are the gold standard, but cannot be performed reliably in children below ~8 months. (67) The methods currently employed to quantify auditory function in neonates are otoacoustic emissions (OAE) and auditory brainstem responses (ABR). Serial diagnostic OAE and ABR testing has been used to monitor ototoxicity in children receiving aminoglycosides. (68-71) There are no equivalent reports describing serial OAE or ABR following vancomycin use. Diagnostic OAE or ABR is labour-intensive and difficult to interpret in premature infants (PMA <34 weeks) due to immaturity of the cochlea and central auditory pathways (72-78).

Universal newborn hearing screening (UNHS) uses a combination of automated OAE and ABR, and is designed to identify severe permanent hearing impairment in term neonates. Standardised UNHS programmes are now being implemented in the USA

and Europe. (79, 80) UNHS is a single-point assessment and is not designed to detect and monitor subtle high-frequency hearing loss that typically occurs in drug-induced ototoxicity.

### **Pre-Clinical Studies**

The association between vancomycin use and hearing impairment is controversial. As in the case of nephrotoxicity, the mechanism of vancomycin-induced ototoxicity is thought to involve dose-dependent intracellular oxidative damage, which leads to the loss of cochlear sensory hair cells resulting in high frequency hearing loss. (81) Ototoxicity has not been consistently demonstrated with vancomycin in animal models. (25, 82-84) Studies in guinea pigs found no evidence of vancomycin ototoxicity, but found that vancomycin increases the probability of ototoxicity, measured by OAE, when co-administered with gentamicin. (85)

# **Clinical Studies**

The majority of vancomycin-associated ototoxicity was reported early in the drug's use in patients treated with indeterminate doses of an impure fermentation product who often received concomitant therapy with other severely ototoxic agents. (86, 87) Overall, the available recent literature on vancomycin-associated ototoxicity are heterogeneous and in many cases causation is uncertain.

### Adult Studies

Observational studies have suggested the risk of ototoxicity may be increased with vancomycin doses ≥4 g daily in adults. In a retrospective study of 89 adults receiving > 14 days of high-dose vancomycin, Forouzesh, *et al* reported high-frequency hearing

loss identified by pure-tone audiometry in ten patients (12%), a rate significantly higher than reported at standard doses. (88) However, the study included patients receiving concomitant aminoglycosides and diuretics, and identified no significant difference in mean trough vancomycin concentrations in patients with and without abnormal audiometry, suggesting the phenomenon was unlikely to be dose dependent.

### Paediatric & Neonatal Studies

Ototoxicity is infrequently reported in neonates and children treated with vancomycin in general. (89) Buckingham, *et al*, however, reported a high incidence of hearing loss in children with pneumococcal meningitis treated with vancomycin. (90) Patients aged 5-20 years (n=109) received vancomycin alongside a third-generation cephalosporin. Over half of those surviving had significant permanent hearing loss (n=37, 55%). Of note, hearing loss was independently associated with vancomycin administration within two hours of presentation. Whether these findings apply to other populations or reflect the specific pathophysiology of pneumococcal meningitis is uncertain. The findings do, however, highlight the need to consider population and disease-specific factors when reporting drug toxicity.

In neonates, reporting of vancomycin-associated ototoxicity comes almost exclusively from routine UNHS. De Hoog, *et al* studied 625 neonates over two years from a single NICU in the Netherlands. Exposure to vancomycin (alone or in combination with tobramycin or furosemide) was not associated with a significant rise in screening failure rates. (91) Similarly, in a retrospective study of >7000 infants, Gopel, *et al* found no association between vancomycin exposure and UNHS failure rates. (92) Vella-Brincat, *et al* described UNHS outcomes over five years from a single centre in

New Zealand. The cohort included 41 neonates who received vancomycin, and found higher failure rates (n=6; 22%), compared with those not exposed to vancomycin (n = 85/1,233; 7%). The significance of these finding is uncertain given the lack of systematic follow-up and many potential confounding factors.

### **CONCLUSION**

Despite historical concerns, a large body of evidence now exists to support the favourable safety profile of vancomycin in humans. In adults, nephrotoxicity is exposure-dependent, and more aggressive dosing may result in a significantly increased risk of toxicity. By comparison, current dosing regimens in neonates are conservative, and result in sub-therapeutic concentrations in a large proportion of patients. At present, insufficient evidence exists to inform conclusions about the safety of higher doses of vancomycin in neonates. Furthermore, the clinical difficulty in ascertaining a diagnosis of sepsis with certainty in neonates makes risk-benefit analyses challenging. The development and use of novel diagnostic biomarkers are likely to improve clinical decision-making and trial design in the years to come. (93) Dose modification based on trough vancomycin concentrations, which are, in turn, largely determined by renal function, adds complexity in determining exposure-toxicity relationships. These issues are probably best addressed with detailed prospective, observational data that define renal impairment according to standardised criteria and allow for a more clinically valid estimation of risk.

Ototoxicity appears rare in patients of all ages treated with vancomycin. Diagnostic OAE and ABR may detect subtle high-frequency hearing loss following vancomycin exposure. These tests can only reliably be carried out in babies >34 weeks before

which many preterm neonates will already have receive courses of vancomycin.

UNHS at term is unlikely to detect subtle hearing impairment and identify exposure-toxicity relationships. Prospective studies that use robust and standardised definitions of hearing impairment based on diagnostic testing, such as with visual reinforcement audiometry at 8-12 months, are needed to further clarify the risk clinically significant ototoxicity.

Source	PMA (weeks)	PNA (days)	Creatinine (mg/dL)	Dose (mg/kg)	Interval (hourly)
British National Formulary for Children	<29			15	24
	29–35			15	12
Manual of Childhood Infections: The Blue Book	>35			15	8
	≤29	0–14		10 (bactaeremia)	18
N. C. S.				15 (meningitis)	
Neofax®		>14		10	12
The Harriet Lane Handbook	30–36	0–14		10	12
The Harriet Lane Handbook		>14		10	8
The Sanford guide	37–44	0-7		10	12
The Samora guide		>7		10	8
	≥45	Any PNA		10	6
			<0.7	15	12

0.7–0.9	20	24
1–1.2	15	24
1.3–1.6	10	24
>1.6	15	48
	1-1.2	1–1.2 15 1.3–1.6 10

Table 1. Current dosing regimens for vancomycin in neonates

Γable 2. Studies descr	ribing nephrotoxicity in neo	nates receiving vancomy	yein	

Author (year)	Study population receiving	Dosing regimen	Concomitant	Timing/Target	Nephrotoxicity	Incidence of
Design	vancomycin		nephrotoxic	levels (μg/mL)	Definition	nephrotoxicity
	(mean ± SD unless		agents			
	otherwise stated)					
Constance (2015)	n = 533	Intermittent infusion: 12-40	Gentamicin	Timing and	SCr rise by	Transient rise in SCr in
Propensity-	PMA <25 weeks n = 35, 25-	mg/kg per day (not further	Ibuprofen	target	≥150% within 48	12/533 patients receiving
matched cohort	28 weeks $n = 73, 28-32$	described)		concentrations	hours or SCr	vancomycin with
study (23)	weeks $n = 158, 32-27$ weeks			not reported	≥1.5 mg/dL (132	gentamicin
	n = 152, >37 weeks $n = 115$				μmol/L)	
	weight 1649 (1060-2504) g				persisting ≥48 h	
	median (IQR)					
Moffett (2015)	n= 83 neonates with critical	Not specified	Gentamicin		Doubling of SCr	2 developed AKI. AKI
Retrospective	cardiac disease		Furosemide			more likely with
case-control			Amphotericin			concomitant nephrotoxic
study (39)						agents

Petrie	n = 83	Intermittent infusion:	Not reported	Timing not	Rise in SCr not	None detected
(2015)	PMA 30+3 (23+6-52+4)	15 mg/kg per dose 8-24		reported;	further defined	
Retrospective	weeks, weight 1.12 (0.56-	hourly according to PMA		trough 10-15		
observational	4.7) g median (IQR)					
study (40)						
Vandendriessche	n = 223	Intermittent infusion:	Not reported	Timing not	Difference in	None detected
(2014)	PMA 32 (24–41) weeks,	10-15 mg/kg 6-24 hourly		reported;	mean SCr	
Retrospective	median (range)	according to PMA, PNA		trough > 10		
observational	1650 (420–4680) g	and SCr				
study (41)						
Frymoyer	n = 249	15-20 mg/kg per dose 12-24	Not reported	After 3 <sup>rd</sup> dose;	Rise in SCr not	None detected
(2014)Prospectiv	PMA 39 (32–42) weeks,	hourly according to SCr and		trough 7-10	further defined	
e observational	weight 2900 (1600-3700) g	weight				
study (42)	median (IQR)					

Kim (2014)	n=50	15-40 mg/kg per dose 6–12	Not reported	Timing not	SCr >25% above	None detected
Retrospective	PMA $27.4 \pm 2.5$ weeks,	hourly according to PMA		reported;	baseline or urine	
observational	weight 1020 ± 420 g			trough ≤ 40	output <1 ml/kg/h	
study (43)						
Linder	n = 126	Intermittent infusion: 10	Not reported	1-2 days;	Not formally	None detected
(2013)	PMA $27.5 \pm 2.6$ weeks,	mg/kg per dose		Trough 10,	defined	
Retrospective	weight $1026 \pm 269 \text{ g}$	8-18 hourly according to		peak 30-40		
observational		PMA and PNA				
study (44)						
Patel	n = 32 (40 treatment	n = 20	Not reported	Before 3 <sup>rd</sup>	Rise in mean ±	No significant rise in SCr
(2013)	courses)	Intermittent infusion: 10-20		dose;	SD in SCr not	(mean $\pm$ SD) in either
Retrospective	PMA 36 (26–62) weeks,	mg/kg per dose 8-12 hourly		15-25	further defined	intermittent dosing or
observational	weight 2200 (620-6900) g	according to SCr				continuous infusion.
study (3)	median (IQR)	n = 20				

		Continuous infusion: 20–60				Two patients developed
		mg/kg daily according to				renal failure 2 <sup>nd</sup> to a
		PMA and SCr				decline in overall clinical
						status.
Zhao	n = 116	Loading dose: 10-15 mg/kg	Not reported	Timing not	SCr > 70 or 90	None detected
(2013)	PMA $33.8 \pm 5.3$ weeks,			reported; 15-	μmol/L based on	
Prospective	weight 1700 ± 964 g	Continuous infusion:		25	PNA	
observational		15-35 mg/kg daily				
study (45)		according to PMA, PNA				
		and SCr				
Irikura	n = 54	n = 21	Not reported	Timing and	Rise in SCr not	Transient rise in SCr in 3
(2011)	n = 21 (SCr-based dosing)	Intermittent infusion: 10-20		target	further defined	patients (two receiving
	PMA $29.65 \pm 5.27$ weeks	mg/kg per dose 12-48		concentrations		dose based on
	Birthweight 1322 ± 951 g	hourly according to SCr		not reported		weight/PNA and one

Prospective	n = 33 (dosing based on	n = 33				receiving dosing based
observational	weight & PNA)	Intermittent infusion: 10				on baseline SCr,
study	PMA 33.18 ± 5.94 weeks	mg/kg per dose 6-18 hourly				respectively).
(46)	Birthweight $1839 \pm 915 \text{ g}$	according to weight and				
		PNA				
Oudin	n = 47	Loading dose: 7 mg/kg	Not reported	Timing not	Rise in SCr not	Transient rise in SCr in 3
(2011)	PMA 29.5 $\pm$ 27, weight	Continuous infusion:		reported;	further defined	patients resolved by 3
PK study (47)	$1500 \pm 970 \text{ g}$	30mg/kg daily		trough 10-30		weeks after exposure.
Plan	n = 145	Intermittent infusion:	Diuretics, not	Timing not	Rise in SCr not	No significant rise in SCr
(2008)	PMA 28 (26–29) weeks,	15-30 mg/kg per dose 24	further defined	reported;	further defined	detected
Prospective	Weight 904 (780–1160) g	hourly according to SCr		trough 10-25		
observational	median (IQR)					
study (48)						

Giapros	n = 70	Intermittent infusion every	Gentamicin	Timing and	Increased urinary	5 patients with severe
(2007)	Weight <1000 g	12 – 48 h – not further		target not	excretion of	renal tubular disturbance
Retrospective		defined		reported	potassium,	(3 with raised SCr). All
observational					calcium and	abnormalities returned to
study (49)					phosphate; raised	baseline within 2 weeks
					SCr	of the last antibiotic
						course
Frattarelli	n = 153	Intermittent infusion not	Not reported	Timing not	Rise in SCr not	None detected
(2005)	SGA; PMA $48 \pm 8$ weeks,	further defined		reported;	further defined	
Retrospective	weight 641 ± 181 g			trough, target		
observational	Non-SGA; PMA 47 ± 8			not reported		
study (50)	weeks, weight $1158 \pm 765$ g					
Miner	n = 2	Intermittent infusion:	None	Timing not	Rise in SCr not	Transient rise SCr in one
(2004)				reported; peak	further defined	patient

Case series (51)	Case 1 – PMA 24 weeks at	150 mg/kg (10x accidental		concentration,		
	birth. Day 53 of life when	overdose)		>300		
	vancomycin commenced					
	Case 2 – PMA 28 weeks at					
	birth. Vancomycin					
	commenced day 9 of life					
Deville	n = 20	Intermittent infusion:	Gentamicin	Timing not	SCr twice the	Transient rise in SCr in
(2003)	PMA $38.6 \pm 7.3$ weeks	10-15 mg/kg per dose 6-24		reported;	upper limit of	one patient not further
Phase III, open	Population not further	hourly according to PMA		trough, target	normal or double	described
label RCT (52)	defined			not reported	baseline	
Tan	n = 101	Intermittent infusion:	Not reported	Timing not	Difference in	None detected
(2002)	PMA 30 (23–45) weeks	15 mg/kg per dose 12-18		reported;	mean SCr	
	Median (range)	hourly according to PNA		trough < 10		

Retrospective	Population not further					
observational	defined					
study (53)						
Feferbaum	n = 20	Intermittent infusion:	None	Timing not	Rise in SCr not	None detected
(2001)	PMA $42 \pm 3$ weeks	30-60 mg/kg per day (not		reported;	further defined	
Prospective	Population not further	further described)		trough 5-10,		
observational	defined			peak 30-40		
study (54)						
Machado	n = 25	10-20 mg/kg per dose 6-24	Gentamicin	Timing not	Rise in SCr and	Significant transient rise
(2001)	PMA $38 \pm 1$ weeks,	hourly according to PMA		reported;	urea 5 days after	in intra-individual SCr
Prospective	3130 ± 861.3 g			trough 5-10,	treatment started	but not urea. Not further
observational				peak 20-40	not otherwise	described.
study (55)					further defined	

De Hoog	n = 108	Intermittent infusion:	Not reported	Steady state	Increase in SCr or	None detected
(2000)	PMA 28.9 (24 – 41) weeks	30 mg/kg per dose 12		(not further	urea not further	
Retrospective	Birthweight 1002 g (485 –	hourly		defined); Peak	defined	
observational	4625 g)			≤40, trough 5-		
study (56)	Median (IQR)			15		
Bhatt-Mehta	n = 69	Intermittent infusion:	Gentamicin	After 3 <sup>rd</sup> dose;	SCr double	Transient rise in SCr in
(1999)	PMA $28.9 \pm 3.0$ weeks,	10-15 mg/kg per dose every		Peak ≤40	baseline or >0.6	6/61 patients with peak
Retrospective	weight 1219 ± 516 g	6-36 hourly according to			mg/dL (53	concentration ≤40 and
observational		PMA			μmol/L)	none with peak
study (57)						concentrations >40
Goebel	n = 1	100 mg/kg	Not reported	12 hours post	Anuria. Not	Commenced
(1999)	Term infant, PNA 6 d			dose; no target	otherwise further	hemofiltration due to
Case report (58)	Solitary dysplastic kidney				defined	persistently elevated
						vancomycin level (240

						μg/mL). Normal renal
						function at PNA 3 weeks.
Müller	n = 2	Intermittent infusion:	None	9 hours post	Reduced GFR or	Transient
(1999)	PMA 35 weeks, weight	Single dose 35 mg/kg and		dose; no target	microproteinuria	microproteinuria in both
Case series (59)	1985 and 2390 g	38 mg/kg (accidental		reported	(urinary protein	patients but no change in
		overdose)			electrophoresis)	GFR
Pawlotsky	n = 53	n = 29	None	Post loading	Rise in SCr and	Transient rise in SCr in
(1998)	PMA $33.5 \pm 3.7$ weeks,	Loading dose: 7 mg/kg		dose; peak ≤40	urea not further	one patient with
Non-randomised	weight $1500 \pm 300 \text{ g (n =}$	Continuous infusion:			defined	klebsiella septicaemia
un-blinded trial	29)	10-40 mg/kg per over 24		Steady state		
(60)	PMA $33.9 \pm 4.8$ weeks,	hours according to PMA		(not further		
	weight $1800 \pm 800 \text{ g (n =}$	and weight		defined);		
	24)	n = 24		10-30		
		Continuous infusion:				

		10-30 mg/kg per over 24				
		hours according to PMA				
		and weight				
Sakata	n = 20	Intermittent infusion: 9-11	None	Timing and	Rise in SCr,	Transient rise in NAG
(1996)	PMA $26.3 \pm 1.4$ weeks	mg/kg per dose 12 hourly		target	fractional	index and FENa after
Prospective	Population not further			concentrations	excretion of	treatment in one patient
observational	defined			not reported	sodium (FENa)	
study (61)					and NAG index	
					(NAG:creatinine	
					ratio)	
McDougal	n = 44	Intermittent infusion:	Not reported	Steady state	Increase in SCr or	None detected
(1995)	PMA $29.4 \pm 0.9$ weeks,	15-18 mg/kg per dose 12-36		(not further	urea not further	
	weight $972 \pm 178 \text{ g (n = 16)}$	hourly according to PMA		defined); peak	defined	

Prospective	PMA $32.9 \pm 1.8$ weeks,			25-35, trough		
observational	weight $1379 \pm 382 \text{ g (n = }$			5-10		
study (62)	15)					
	PMA $39.2 \pm 2$ weeks,					
	weight $2616 \pm 753 \text{ g (n = }$					
	13)					
Tissing (1993)	n = 1	Intermittent infusion:	Tobramycin	Timing not	Rise in SCr and	Transient rise in SCr and
Case report (63)	PMA 29.3 weeks, weight	15 mg/kg per dose every 12		reported; no	oliguria not	persistently elevated
	1520 g	hours		target	further defined	vancomycin level.
				(reported level		Resolution at 2 months
				63.3)		of age.
Lisby-Sutch	n = 13	Intermittent infusion:	Not reported	Timing not	Increase in SCr or	None detected
(1988)	PMA $29.8 \pm 3.4$ weeks,			reported; peak	urea not further	
Prospective	weight 1350 ± 500 g				defined	

PK study (64)		10 mg/kg per dose 6-12		25–35, trough,		
		hourly according to PMA		5–10		
		and weight				
Nahata	n = 61 < 1 year	Intermittent infusion:	Gentamicin	Timing not	Doubling of SCr	None detected
(1987)	Further clinical information	20-60 mg/kg/day (mean 35)		reported;	concentration	
Prospective	not available.			trough 2-18		
observational						
study (65)						
James	n = 20	Intermittent infusion:	Not reported	Timing not	Not formally	None reported although
(1987)	PMA $26.5 \pm 2.6$ weeks,	9-18 mg/kg per dose 12		reported; peak	defined	positive linear correlation
Prospective PK	weight 880 ± 340 g	hourly		30, trough 6		between vancomycin
study (33)						level and SCr identified

Dean	n = 28	Intermittent infusion:	Gentamicin	Timing not	Increase in SCr	Transient rise in SCr in	
(1985)	Population not further	11-55 mg/kg (mean 30		reported; peak	by >0.5 mg/dl (45	2/19 and 2/9 patients	
Retrospective	defined	mg/kg) daily not further		20-40, trough	μmol/L)	receiving vancomycin	
observational		defined		5-10		alone and with	
study (66)						gentamicin, respectively.	

# REFERENCES

Papers of special note have been highlighted as:

- \* of special interest
- \*\* of outstanding interest
- Sheldrick GM, Jones PG, Kennard O, Williams DH, Smith GA. 1978.
   Structure of vancomycin and its complex with acetyl-D-alanyl-D-alanine.
   Nature 271:223-225.
- 2. **Moellering RC, Jr.** 2006. Vancomycin: a 50-year reassessment. Clinical infectious diseases: an official publication of the Infectious Diseases Society of America **42 Suppl 1:**S3-4.
- 3. **Patel AD, Anand D, Lucas C, Thomson AH.** 2013. Continuous infusion of vancomycin in neonates. Archives of disease in childhood **98:**478-479.
- 4. **Sinkeler FS, de Haan TR, Hodiamont CJ, Bijleveld YA, Pajkrt D, Mathot RA.** 2014. Inadequate vancomycin therapy in term and preterm neonates: a retrospective analysis of trough serum concentrations in relation to minimal inhibitory concentrations. BMC pediatrics **14:**193.
- 5. **Jones RN.** 2006. Microbiological features of vancomycin in the 21st century: minimum inhibitory concentration creep, bactericidal/static activity, and applied breakpoints to predict clinical outcomes or detect resistant strains. Clinical infectious diseases: an official publication of the Infectious Diseases Society of America **42 Suppl 1:**S13-24.
- Hidayat LK, Hsu DI, Quist R, Shriner KA, Wong-Beringer A. 2006. Highdose vancomycin therapy for methicillin-resistant Staphylococcus aureus infections: efficacy and toxicity. Archives of internal medicine 166:2138-2144.

- 7. **American Thoracic S, Infectious Diseases Society of A.** 2005. Guidelines for the management of adults with hospital-acquired, ventilator-associated, and healthcare-associated pneumonia. American journal of respiratory and critical care medicine **171:**388-416.
- 8. Liu C, Bayer A, Cosgrove SE, Daum RS, Fridkin SK, Gorwitz RJ,
  Kaplan SL, Karchmer AW, Levine DP, Murray BE, M JR, Talan DA,
  Chambers HF, Infectious Diseases Society of A. 2011. Clinical practice
  guidelines by the infectious diseases society of america for the treatment of
  methicillin-resistant Staphylococcus aureus infections in adults and children.
  Clinical infectious diseases: an official publication of the Infectious Diseases
  Society of America 52:e18-55.
- 9. **Rybak M, Lomaestro B, Rotschafer JC, Moellering R, Jr., Craig W, Billeter M, Dalovisio JR, Levine DP.** 2009. Therapeutic monitoring of vancomycin in adult patients: a consensus review of the American Society of Health-System Pharmacists, the Infectious Diseases Society of America, and the Society of Infectious Diseases Pharmacists. American journal of health-system Pharmacy: AJHP: official journal of the American Society of Health-System Pharmacists **66:**82-98.
- 10. **Rybak MJ, Lomaestro BM, Rotschafer JC, Moellering RC, Craig WA, Billeter M, Dalovisio JR, Levine DP.** 2009. Vancomycin therapeutic guidelines: a summary of consensus recommendations from the infectious diseases Society of America, the American Society of Health-System Pharmacists, and the Society of Infectious Diseases Pharmacists. Clinical infectious diseases: an official publication of the Infectious Diseases Society of America **49:**325-327.

- 11. **Strauss J, Zilleruelo G, Gorman HM, Baker R, Galindez R.** 1982. [Renal function in the fetus and the newborn infant]. Boletin medico del Hospital Infantil de Mexico **39:**243-252.
- 12. **Bueva A, Guignard JP.** 1994. Renal function in preterm neonates. Pediatric research **36:**572-577.
- 13. **Guignard JP, Drukker A.** 1999. Why do newborn infants have a high plasma creatinine? Pediatrics **103:**e49.
- 14. Allegaert K, Kuppens M, Mekahli D, Levtchenko E, Vanstapel F, Vanhole C, van den Anker JN. 2012. Creatinine reference values in ELBW infants: impact of quantification by Jaffe or enzymatic method. The journal of maternal-fetal & neonatal medicine: the official journal of the European Association of Perinatal Medicine, the Federation of Asia and Oceania Perinatal Societies, the International Society of Perinatal Obstet 25:1678-1681.
- 15. Delanghe JR, Cobbaert C, Harmoinen A, Jansen R, Laitinen P,
  Panteghini M. 2011. Focusing on the clinical impact of standardization of creatinine measurements: a report by the EFCC Working Group on Creatinine Standardization. Clinical chemistry and laboratory medicine: CCLM / FESCC 49:977-982.
- 16. Hermsen ED, Hanson M, Sankaranarayanan J, Stoner JA, Florescu MC, Rupp ME. 2010. Clinical outcomes and nephrotoxicity associated with vancomycin trough concentrations during treatment of deep-seated infections. Expert opinion on drug safety 9:9-14.
- 17. **Jeffres MN, Isakow W, Doherty JA, Micek ST, Kollef MH.** 2007. A retrospective analysis of possible renal toxicity associated with vancomycin in patients with health care-associated methicillin-resistant Staphylococcus aureus pneumonia. Clinical therapeutics **29:**1107-1115.

- 18. **Lodise TP, Lomaestro B, Graves J, Drusano GL.** 2008. Larger vancomycin doses (at least four grams per day) are associated with an increased incidence of nephrotoxicity. Antimicrobial agents and chemotherapy **52:**1330-1336.
- Ragab AR, Al-Mazroua MK, Al-Harony MA. 2013. Incidence and predisposing factors of vancomycin-induced nephrotoxicity in children.
   Infectious diseases and therapy 2:37-46.
- 20. Askenazi DJ, Halloran B, Patil N, Keeling S, Saeidi B, Koralkar R,
  Ambalavanan N. 2015. Genetic polymorphisms of heme-oxygenase 1 (HO-1)
  may impact on acute kidney injury, bronchopulmonary dysplasia, and
  mortality in premature infants. Pediatric research 77:793-798.
- 21. **Lopes JA, Jorge S.** 2013. The RIFLE and AKIN classifications for acute kidney injury: a critical and comprehensive review. Clin Kidney J **6:**8-14.
- 22. Bezerra CT, Vaz Cunha LC, Liborio AB. 2013. Defining reduced urine output in neonatal ICU: importance for mortality and acute kidney injury classification. Nephrology, dialysis, transplantation: official publication of the European Dialysis and Transplant Association European Renal Association 28:901-909.
- 23. Constance JE, Balch AH, Stockmann C, Linakis MW, Korgenski EK, Roberts JK, Ward RM, Sherwin CM, Spigarelli MG. 2015. A propensitymatched cohort study of vancomycin-associated nephrotoxicity in neonates. Arch Dis Child Fetal Neonatal Ed. [Epub ahead of print]
  - \* Well designed prospective cohort study of nephrotoxicity in neonates receiving vancomycin.
- 24. **Marre R, Schulz E, Anders T, Sack K.** 1984. Renal tolerance and pharmacokinetics of vancomycin in rats. The Journal of antimicrobial chemotherapy **14:**253-260.

- 25. Wood CA, Kohlhepp SJ, Kohnen PW, Houghton DC, Gilbert DN. 1986.
  Vancomycin enhancement of experimental tobramycin nephrotoxicity.
  Antimicrobial agents and chemotherapy 30:20-24.
- 26. Celik I, Cihangiroglu M, Ilhan N, Akpolat N, Akbulut HH. 2005.
  Protective effects of different antioxidants and amrinone on vancomycin-induced nephrotoxicity. Basic & clinical pharmacology & toxicology 97:325-332.
- 27. Cetin H, Olgar S, Oktem F, Ciris M, Uz E, Aslan C, Ozguner F. 2007.
  Novel evidence suggesting an anti-oxidant property for erythropoietin on vancomycin-induced nephrotoxicity in a rat model. Clinical and experimental pharmacology & physiology 34:1181-1185.
- 28. Hodoshima N, Nakano Y, Izumi M, Mitomi N, Nakamura Y, Aoki M, Gyobu A, Shibasaki S, Kurosawa T. 2004. Protective effect of inactive ingredients against nephrotoxicity of vancomycin hydrochloride in rats. Drug metabolism and pharmacokinetics 19:68-75.
- 29. King DW, Smith MA. 2004. Proliferative responses observed following vancomycin treatment in renal proximal tubule epithelial cells. Toxicology in vitro: an international journal published in association with BIBRA 18:797-803.
- 30. Oktem F, Arslan MK, Ozguner F, Candir O, Yilmaz HR, Ciris M, Uz E. 2005. In vivo evidences suggesting the role of oxidative stress in pathogenesis of vancomycin-induced nephrotoxicity: protection by erdosteine. Toxicology 215:227-233.
- 31. **Toyoguchi T, Takahashi S, Hosoya J, Nakagawa Y, Watanabe H.** 1997. Nephrotoxicity of vancomycin and drug interaction study with cilastatin in rabbits. Antimicrobial agents and chemotherapy **41:**1985-1990.

- 32. **Dieterich C, Puey A, Lin S, Swezey R, Furimsky A, Fairchild D, Mirsalis JC, Ng HH.** 2009. Gene expression analysis reveals new possible mechanisms of vancomycin-induced nephrotoxicity and identifies gene markers candidates.

  Toxicological sciences: an official journal of the Society of Toxicology **107:**258-269.
- 33. James A, Koren G, Milliken J, Soldin S, Prober C. 1987. Vancomycin pharmacokinetics and dose recommendations for preterm infants.
  Antimicrobial agents and chemotherapy 31:52-54.
- 34. Chertow GM, Burdick E, Honour M, Bonventre JV, Bates DW. 2005.

  Acute kidney injury, mortality, length of stay, and costs in hospitalized patients. Journal of the American Society of Nephrology: JASN 16:3365-3370.
- 35. McKamy S, Hernandez E, Jahng M, Moriwaki T, Deveikis A, Le J. 2011.

  Incidence and risk factors influencing the development of vancomycin

  nephrotoxicity in children. The Journal of pediatrics 158:422-426.
- 36. van Hal SJ, Paterson DL, Lodise TP. 2013. Systematic review and metaanalysis of vancomycin-induced nephrotoxicity associated with dosing schedules that maintain troughs between 15 and 20 milligrams per liter.
  Antimicrobial agents and chemotherapy 57:734-744.
  - \*\* Comprehensive meta-analysis of studies of toxicity associated with increased target trough vancomycin concentrations in adults.
- 37. Fowler VG, Jr., Boucher HW, Corey GR, Abrutyn E, Karchmer AW, Rupp ME, Levine DP, Chambers HF, Tally FP, Vigliani GA, Cabell CH, Link AS, DeMeyer I, Filler SG, Zervos M, Cook P, Parsonnet J, Bernstein JM, Price CS, Forrest GN, Fatkenheuer G, Gareca M, Rehm SJ, Brodt HR, Tice A, Cosgrove SE, Endocarditis Sa, Bacteremia Study G. 2006.

- Daptomycin versus standard therapy for bacteremia and endocarditis caused by Staphylococcus aureus. The New England journal of medicine **355:**653-665.
- 38. Metsvaht T, Nellis G, Varendi H, Nunn AJ, Graham S, Rieutord A, Storme T, McElnay J, Mulla H, Turner MA, Lutsar I. 2015. High variability in the dosing of commonly used antibiotics revealed by a Europewide point prevalence study: implications for research and dissemination.

  BMC pediatrics 15:41.
  - \*\* First systematic survey identifying widespread systematic underdosing of vancomycin in infants and children.
- 39. **Moffett BS, Hilvers PS, Dinh K, Arikan AA, Checchia P, Bronicki R.**2015. Vancomycin-associated acute kidney injury in pediatric cardiac intensive care patients. Congenital heart disease **10:**E6-10.
- 40. **Petrie K, O'Brien C, Bhushan S, Tonna A.** 2015. Neonatal vancomycin trough level audit using British National Formulary for Children dosing.

  Archives of disease in childhood. Fetal and neonatal edition **100:**F278-279.
- 41. Vandendriessche A, Allegaert K, Cossey V, Naulaers G, Saegeman V, Smits A. 2014. Prospective validation of neonatal vancomycin dosing regimens is urgently needed. Current therapeutic research, clinical and experimental 76:51-57.
- 42. Frymoyer A, Hersh AL, El-Komy MH, Gaskari S, Su F, Drover DR, Van Meurs K. 2014. Association between vancomycin trough concentration and area under the concentration-time curve in neonates. Antimicrobial agents and chemotherapy **58**:6454-6461.
- 43. Kim J, Walker SA, Iaboni DC, Walker SE, Elligsen M, Dunn MS, Allen VG, Simor A. 2014. Determination of vancomycin pharmacokinetics in

- neonates to develop practical initial dosing recommendations. Antimicrobial agents and chemotherapy **58:**2830-2840.
- 44. **Linder N, Lubin D, Hernandez A, Amit L, Ashkenazi S.** 2013. Duration of vancomycin treatment for coagulase-negative Staphylococcus sepsis in very low birth weight infants. British journal of clinical pharmacology **76:**58-64.
- Zhao W, Lopez E, Biran V, Durrmeyer X, Fakhoury M, Jacqz-Aigrain E.
   2013. Vancomycin continuous infusion in neonates: dosing optimisation and therapeutic drug monitoring. Archives of disease in childhood 98:449-453.
   \* Well designed clinical study describing outcomes and PKPD modelling for vancomycin administered via continuous infusion in neonates.
- 46. Irikura M, Fujiyama A, Saita F, Fukushima S, Kitaoka H, Fukuda T, Kawase A, Kondo Y, Ishitsuka Y, Kondo G, Maeda T, Yukawa E, Irie T. 2011. Evaluation of the vancomycin dosage regimen based on serum creatinine used in the neonatal intensive care unit. Pediatrics international: official journal of the Japan Pediatric Society 53:1038-1044.
- 47. **Oudin C, Vialet R, Boulamery A, Martin C, Simon N.** 2011. Vancomycin prescription in neonates and young infants: toward a simplified dosage.

  Archives of disease in childhood. Fetal and neonatal edition **96:**F365-370.
- 48. Plan O, Cambonie G, Barbotte E, Meyer P, Devine C, Milesi C, Pidoux O, Badr M, Picaud JC. 2008. Continuous-infusion vancomycin therapy for preterm neonates with suspected or documented Gram-positive infections: a new dosage schedule. Archives of disease in childhood. Fetal and neonatal edition 93:F418-421.
- 49. **Giapros VI, Papadimitriou FK, Andronikou SK.** 2007. Tubular disorders in low birth weight neonates after prolonged antibiotic treatment. Neonatology **91:**140-144.

- 50. **Frattarelli DA, Ergun H, Lulic-Botica M, Lehr VT, Aranda JV.** 2005. Vancomycin elimination in human infants with intrauterine growth retardation. The Pediatric infectious disease journal **24:**979-983.
- 51. **Miner LJ, Faix RG.** 2004. Large vancomycin overdose in two premature infants with minimal toxicity. American journal of perinatology **21:**433-438.
- 52. Deville JG, Adler S, Azimi PH, Jantausch BA, Morfin MR, Beltran S, Edge-Padbury B, Naberhuis-Stehouwer S, Bruss JB. 2003. Linezolid versus vancomycin in the treatment of known or suspected resistant grampositive infections in neonates. The Pediatric infectious disease journal 22:S158-163.
- 53. **Tan WH, Brown N, Kelsall AW, McClure RJ.** 2002. Dose regimen for vancomycin not needing serum peak levels? Archives of disease in childhood. Fetal and neonatal edition **87:**F214-216.
- 54. Feferbaum R, Kobol Machado JK, de Albuquerque Diniz EM, Okay TS, Santos SR, Ceccon ME, Krebs VL, de Araujo MC, Costa Vaz A. 2001.

  Vancomycin monitoring in term newborns: comparison of peak and trough serum concentrations determined by high performance liquid chromatography and fluorescence polarization immunoassay. Revista do Hospital das Clinicas 56:149-152.
- 55. Machado JK, Feferbaum R, Diniz EM, Okay TS, Ceccon ME, Costa Vaz FA. 2001. Monitoring the treatment of sepsis with vancomycin in term newborn infants. Revista do Hospital das Clinicas 56:17-24.
- 56. de Hoog M, Schoemaker RC, Mouton JW, van den Anker JN. 2000.
  Vancomycin population pharmacokinetics in neonates. Clinical pharmacology and therapeutics 67:360-367.

- 57. **Bhatt-Mehta V, Schumacher RE, Faix RG, Leady M, Brenner T.** 1999. Lack of vancomycin-associated nephrotoxicity in newborn infants: a casecontrol study. Pediatrics **103:**e48.
- 58. **Goebel J, Ananth M, Lewy JE.** 1999. Hemodiafiltration for vancomycin overdose in a neonate with end-stage renal failure. Pediatric nephrology **13:**423-425.
- Muller D, Hufnagel M, Suttorp M. 1999. Accidental overdose of vancomycin in preterm twins. The Pediatric infectious disease journal 18:744-745.
- 60. **Pawlotsky F, Thomas A, Kergueris MF, Debillon T, Roze JC.** 1998. Constant rate infusion of vancomycin in premature neonates: a new dosage schedule. British journal of clinical pharmacology **46:**163-167.
- 61. Sakata H, Maruyama S, Ishioka T, Shirai M, Taketazu M, Taketazu G.
  1996. Change of renal function during vancomycin therapy in extremely low birthweight infants. Acta paediatrica Japonica; Overseas edition 38:619-621.
- 62. **McDougal A, Ling EW, Levine M.** 1995. Vancomycin pharmacokinetics and dosing in premature neonates. Therapeutic drug monitoring **17:**319-326.
- 63. Tissing WJ, Umans-Eckenhausen MA, van den Anker JN. 1993.
  Vancomycin intoxication in a preterm neonate. European journal of pediatrics
  152:700.
- 64. **Lisby-Sutch SM, Nahata MC.** 1988. Dosage guidelines for the use of vancomycin based on its pharmacokinetics in infants. European journal of clinical pharmacology **35:**637-642.
- 65. **Nahata MC.** 1987. Lack of nephrotoxicity in pediatric patients receiving concurrent vancomycin and aminoglycoside therapy. Chemotherapy **33:**302-304.

- 66. **Dean RP, Wagner DJ, Tolpin MD.** 1985. Vancomycin/aminoglycoside nephrotoxicity. The Journal of pediatrics **106:**861-862.
- 67. Widen JE, Folsom RC, Cone-Wesson B, Carty L, Dunnell JJ, Koebsell K, Levi A, Mancl L, Ohlrich B, Trouba S, Gorga MP, Sininger YS, Vohr BR, Norton SJ. 2000. Identification of neonatal hearing impairment: hearing status at 8 to 12 months corrected age using a visual reinforcement audiometry protocol. Ear and hearing 21:471-487.
- 68. **Mulheran M, Degg C.** 1997. Comparison of distortion product OAE generation between a patient group requiring frequent gentamic in therapy and control subjects. British journal of audiology **31:**5-9.
- 69. Katbamna B, Homnick DN, Marks JH. 1998. Contralateral suppression of distortion product otoacoustic emissions in children with cystic fibrosis: effects of tobramycin. Journal of the American Academy of Audiology 9:172-178.
- 70. Stavroulaki P, Vossinakis IC, Dinopoulou D, Doudounakis S,
  Adamopoulos G, Apostolopoulos N. 2002. Otoacoustic emissions for monitoring aminoglycoside-induced ototoxicity in children with cystic fibrosis. Archives of otolaryngology--head & neck surgery 128:150-155.
- 71. **Kohelet D, Usher M, Arbel E, Arlazoroff A, Goldberg M.** 1990. Effect of gentamicin on the auditory brainstem evoked response in term infants: a preliminary report. Pediatric research **28:**232-234.
- 72. **Tognola G, Parazzini M, de Jager P, Brienesse P, Ravazzani P, Grandori F.** 2005. Cochlear maturation and otoacoustic emissions in preterm infants: a time-frequency approach. Hearing research **199:**71-80.
- 73. Ubbink SW, van Dijk P, de Kleine E, Brienesse P, Chenault MN, Tan FE,

  Anteunis LJ. 2011. Frequency shifts with age in click-evoked otoacoustic

- emissions of preterm infants. The Journal of the Acoustical Society of America **129:**3788-3796.
- 74. Smurzynski J, Jung MD, Lafreniere D, Kim DO, Kamath MV, Rowe JC, Holman MC, Leonard G. 1993. Distortion-product and click-evoked otoacoustic emissions of preterm and full-term infants. Ear and hearing 14:258-274.
- 75. **Smurzynski J.** 1994. Longitudinal measurements of distortion-product and click-evoked otoacoustic emissions of preterm infants: preliminary results. Ear and hearing **15:**210-223.
- 76. **Chuang SW, Gerber SE, Thornton AR.** 1993. Evoked otoacoustic emissions in preterm infants. International journal of pediatric otorhinolaryngology **26:**39-45.
- 77. **Brienesse P, Maertzdorf W, Anteunis L, Manni J, Blanco C.** 1998. Longterm and short-term variations in amplitude and frequency of spontaneous otoacoustic emissions in pre-term infants. Audiology: official organ of the International Society of Audiology **37:**278-284.
- 78. Norton SJ, Gorga MP, Widen JE, Folsom RC, Sininger Y, Cone-Wesson B, Vohr BR, Mascher K, Fletcher K. 2000. Identification of neonatal hearing impairment: evaluation of transient evoked otoacoustic emission, distortion product otoacoustic emission, and auditory brain stem response test performance. Ear and hearing 21:508-528.
- 79. **Uloziene I, Grandori F.** 2003. The European project AHEAD II on newborn hearing screening. International Congress Series **1240**:329-332.
- 80. Grandori F, Lutman M. 1999. The European Consensus Development Conference on Neonatal Hearing Screening (Milan, May 15-16, 1998).American journal of audiology 8:19-20.

- 81. **Bailie GR, Neal D.** 1988. Vancomycin ototoxicity and nephrotoxicity. A review. Medical toxicology and adverse drug experience **3:**376-386.
- 82. **Wold JS, Turnipseed SA.** 1981. Toxicology of vancomycin in laboratory animals. Reviews of infectious diseases **3 suppl:**S224-229.
- 83. **Tange RA, Kieviet HL, von Marle J, Bagger-Sjoback D, Ring W.** 1989. An experimental study of vancomycin-induced cochlear damage. Archives of oto-rhino-laryngology **246:**67-70.
- 84. **Aronoff GR, Sloan RS, Dinwiddie CB, Jr., Glant MD, Fineberg NS, Luft FC.** 1981. Effects of vancomycin on renal function in rats. Antimicrobial agents and chemotherapy **19:**306-308.
- 85. **Brummett RE, Fox KE, Jacobs F, Kempton JB, Stokes Z, Richmond AB.**1990. Augmented gentamicin ototoxicity induced by vancomycin in guinea
  pigs. Archives of otolaryngology--head & neck surgery **116:**61-64.
- 86. **Brummett RE, Fox KE.** 1989. Vancomycin- and erythromycin-induced hearing loss in humans. Antimicrobial agents and chemotherapy **33:**791-796.
- 87. Elting LS, Rubenstein EB, Kurtin D, Rolston KV, Fangtang J, Martin CG, Raad, II, Whimbey EE, Manzullo E, Bodey GP. 1998. Mississippi mud in the 1990s: risks and outcomes of vancomycin-associated toxicity in general oncology practice. Cancer 83:2597-2607.
- 88. **Forouzesh A, Moise PA, Sakoulas G.** 2009. Vancomycin ototoxicity: a reevaluation in an era of increasing doses. Antimicrobial agents and chemotherapy **53:**483-486.
- 89. **H.D. R, N.J. R.** 1960. Treatment of Severe Staphylococcal Infections in Infancy and Childhood with Vancomycin. Antibiotics Annual:908-916.
- 90. Buckingham SC, McCullers JA, Lujan-Zilbermann J, Knapp KM,Orman KL, English BK. 2006. Early vancomycin therapy and adverse

outcomes in children with pneumococcal meningitis. Pediatrics **117:**1688-1694.

- 91. **de Hoog M, van Zanten BA, Hop WC, Overbosch E, Weisglas-Kuperus N, van den Anker JN.** 2003. Newborn hearing screening: tobramycin and vancomycin are not risk factors for hearing loss. The Journal of pediatrics **142:**41-46.
- 92. Gopel W, Berkowski S, Preuss M, Ziegler A, Kuster H, Felderhoff-Muser U, Gortner L, Mogel M, Hartel C, Herting E, German Neonatal N. 2014.

  Mitochondrial mutation m.1555A>G as a risk factor for failed newborn hearing screening in a large cohort of preterm infants. BMC pediatrics 14:210.
- 93. Smith CL, Dickinson P, Forster T, Khondoker M, Craigon M, Ross A, Storm P, Burgess S, Lacaze P, Stenson BJ, Ghazal P. 2007. Quantitative assessment of human whole blood RNA as a potential biomarker for infectious disease. The Analyst 132:1200-1209.

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## **CONFLICTS OF INTEREST**

None declare.

PMA (weeks)	PNA (days)	Creatinine (mg/dL)	Dose (mg/kg)	Interval (hourly)
<29			15	24
29–35			15	12
>35			15	8
≤29	0–14		10 (bactaeremia)	18
			15 (meningitis)	
	>14		10	12
30–36	0-14		10	12
	>14		10	8
37–44	0-7		10	12
	>7		10	8
≥45	Any PNA		10	6
		<0.7	15	12
		0.7–0.9	20	24
	<29 29–35 >35 <229  30–36  37–44			<29

	1–1.2	15	24
	1.3–1.6	10	24
	>1.6	15	48

Table 1. Current dosing regimens for vancomycin in neonates

Table 2

Table 2. Studies describing nephrotoxicity in neonates receiving vancomycin

Author (year)	Study population receiving	Dosing regimen	Concomitant	Timing/Target	Nephrotoxicity	Incidence of
Design	vancomycin		nephrotoxic	levels (μg/mL)	Definition	nephrotoxicity
	(mean ± SD unless		agents			
	otherwise stated)					
Constance (2015)	n = 533	Intermittent infusion: 12-40	Gentamicin	Timing and	SCr rise by	Transient rise in SCr in
Propensity-	PMA <25 weeks n = 35, 25-	mg/kg per day (not further	Ibuprofen	target	≥150% within 48	12/533 patients receiving
matched cohort	28 weeks n = 73, 28-32	described)		concentrations	hours or SCr	vancomycin with
study (23)	weeks n = 158, 32-27 weeks			not reported	≥1.5 mg/dL (132	gentamicin
	n = 152, >37 weeks $n = 115$				μmol/L)	
	weight 1649 (1060-2504) g				persisting ≥48 h	
	median (IQR)					
Moffett (2015)	n= 83 neonates with critical	Not specified	Gentamicin		Doubling of SCr	2 developed AKI. AKI
Retrospective	cardiac disease		Furosemide			more likely with
case-control			Amphotericin			concomitant nephrotoxic
study (39)						agents

Petrie	n = 83	Intermittent infusion:	Not reported	Timing not	Rise in SCr not	None detected
(2015)	PMA 30+3 (23+6-52+4)	15 mg/kg per dose 8-24		reported;	further defined	
Retrospective	weeks, weight 1.12 (0.56-	hourly according to PMA		trough 10-15		
observational	4.7) g median (IQR)					
study (40)						
Vandendriessche	n = 223	Intermittent infusion:	Not reported	Timing not	Difference in	None detected
(2014)	PMA 32 (24–41) weeks,	10-15 mg/kg 6-24 hourly		reported;	mean SCr	
Retrospective	median (range)	according to PMA, PNA		trough > 10		
observational	1650 (420–4680) g	and SCr				
study (41)						
Frymoyer	n = 249	15-20 mg/kg per dose 12-24	Not reported	After 3 <sup>rd</sup> dose;	Rise in SCr not	None detected
(2014)Prospectiv	PMA 39 (32–42) weeks,	hourly according to SCr and		trough 7-10	further defined	
e observational	weight 2900 (1600-3700) g	weight				
study (42)	median (IQR)					

Kim (2014)	n=50	15-40 mg/kg per dose 6–12	Not reported	Timing not	SCr >25% above	None detected
Retrospective	PMA 27.4 $\pm$ 2.5 weeks,	hourly according to PMA		reported;	baseline or urine	
observational	weight 1020 ± 420 g			trough ≤ 40	output <1 ml/kg/h	
study (43)						
Linder	n = 126	Intermittent infusion: 10	Not reported	1-2 days;	Not formally	None detected
(2013)	PMA $27.5 \pm 2.6$ weeks,	mg/kg per dose		Trough 10,	defined	
Retrospective	weight 1026 ± 269 g	8-18 hourly according to		peak 30-40		
observational		PMA and PNA				
study (44)						
Patel	n = 32 (40 treatment	n = 20	Not reported	Before 3 <sup>rd</sup>	Rise in mean ±	No significant rise in SCr
(2013)	courses)	Intermittent infusion: 10-20		dose;	SD in SCr not	(mean $\pm$ SD) in either
Retrospective	PMA 36 (26–62) weeks,	mg/kg per dose 8-12 hourly		15-25	further defined	intermittent dosing or
observational	weight 2200 (620-6900) g	according to SCr				continuous infusion.
study (3)	median (IQR)	n = 20				Two patients developed
						renal failure 2 <sup>nd</sup> to a

		Continuous infusion: 20–60				decline in overall clinical
		mg/kg daily according to				status.
		PMA and SCr				
Zhao	n = 116	Loading dose: 10-15 mg/kg	Not reported	Timing not	SCr > 70 or 90	None detected
(2013)	PMA $33.8 \pm 5.3$ weeks,			reported; 15-	μmol/L based on	
Prospective	weight 1700 ± 964 g	Continuous infusion:		25	PNA	
observational		15-35 mg/kg daily				
study (45)		according to PMA, PNA				
		and SCr				
Irikura	n = 54	n = 21	Not reported	Timing and	Rise in SCr not	Transient rise in SCr in 3
(2011)	n = 21 (SCr-based dosing)	Intermittent infusion: 10-20		target	further defined	patients (two receiving
Prospective	PMA $29.65 \pm 5.27$ weeks	mg/kg per dose 12-48		concentrations		dose based on
observational	Birthweight 1322 ± 951 g	hourly according to SCr		not reported		weight/PNA and one
study	n = 33 (dosing based on	n = 33				receiving dosing based
(46)	weight & PNA)	Intermittent infusion: 10				on baseline SCr,
	PMA 33.18 ± 5.94 weeks	mg/kg per dose 6-18 hourly				respectively).

	Birthweight 1839 ± 915 g	according to weight and				
		PNA				
Oudin	n = 47	Loading dose: 7 mg/kg	Not reported	Timing not	Rise in SCr not	Transient rise in SCr in 3
(2011)	PMA $29.5 \pm 27$ , weight	Continuous infusion:		reported;	further defined	patients resolved by 3
PK study (47)	$1500 \pm 970 \text{ g}$	30mg/kg daily		trough 10-30		weeks after exposure.
Plan	n = 145	Intermittent infusion:	Diuretics, not	Timing not	Rise in SCr not	No significant rise in SCr
(2008)	PMA 28 (26–29) weeks,	15-30 mg/kg per dose 24	further defined	reported;	further defined	detected
Prospective	Weight 904 (780–1160) g	hourly according to SCr		trough 10-25		
observational	median (IQR)					
study (48)						
Giapros	n = 70	Intermittent infusion every	Gentamicin	Timing and	Increased urinary	5 patients with severe
(2007)	Weight <1000 g	12 – 48 h – not further		target not	excretion of	renal tubular disturbance
Retrospective		defined		reported	potassium,	(3 with raised SCr). All
observational					calcium and	abnormalities returned to
study (49)					phosphate; raised	baseline within 2 weeks
					SCr	

						of the last antibiotic
						course
Frattarelli	n = 153	Intermittent infusion not	Not reported	Timing not	Rise in SCr not	None detected
(2005)	SGA; PMA $48 \pm 8$ weeks,	further defined		reported;	further defined	
Retrospective	weight 641 ± 181 g			trough, target		
observational	Non-SGA; PMA $47 \pm 8$			not reported		
study (50)	weeks, weight $1158 \pm 765$ g					
Miner	n = 2	Intermittent infusion:	None	Timing not	Rise in SCr not	Transient rise SCr in one
(2004)	Case 1 – PMA 24 weeks at	150 mg/kg (10x accidental		reported; peak	further defined	patient
Case series (51)	birth. Day 53 of life when	overdose)		concentration,		
	vancomycin commenced			>300		
	Case 2 – PMA 28 weeks at					
	birth. Vancomycin					
	commenced day 9 of life					

Deville	n = 20	Intermittent infusion:	Gentamicin	Timing not	SCr twice the	Transient rise in SCr in
(2003)	PMA $38.6 \pm 7.3$ weeks	10-15 mg/kg per dose 6-24		reported;	upper limit of	one patient not further
Phase III, open	Population not further	hourly according to PMA		trough, target	normal or double	described
label RCT (52)	defined			not reported	baseline	
Tan	n = 101	Intermittent infusion:	Not reported	Timing not	Difference in	None detected
(2002)	PMA 30 (23–45) weeks	15 mg/kg per dose 12-18		reported;	mean SCr	
Retrospective	Median (range)	hourly according to PNA		trough < 10		
observational	Population not further					
study (53)	defined					
Feferbaum	n = 20	Intermittent infusion:	None	Timing not	Rise in SCr not	None detected
(2001)	PMA $42 \pm 3$ weeks	30-60 mg/kg per day (not		reported;	further defined	
Prospective	Population not further	further described)		trough 5-10,		
observational	defined			peak 30-40		
study (54)						
Machado	n = 25	10-20 mg/kg per dose 6-24	Gentamicin	Timing not	Rise in SCr and	Significant transient rise
(2001)	PMA $38 \pm 1$ weeks,	hourly according to PMA		reported;	urea 5 days after	in intra-individual SCr

Prospective	3130 ± 861.3 g			trough 5-10,	treatment started	but not urea. Not further
observational				peak 20-40	not otherwise	described.
study (55)					further defined	
De Hoog	n = 108	Intermittent infusion:	Not reported	Steady state	Increase in SCr or	None detected
(2000)	PMA 28.9 (24 – 41) weeks	30 mg/kg per dose 12		(not further	urea not further	
Retrospective	Birthweight 1002 g (485 –	hourly		defined); Peak	defined	
observational	4625 g)			≤40, trough 5-		
study (56)	Median (IQR)			15		
Bhatt-Mehta	n = 69	Intermittent infusion:	Gentamicin	After 3 <sup>rd</sup> dose;	SCr double	Transient rise in SCr in
(1999)	PMA $28.9 \pm 3.0$ weeks,	10-15 mg/kg per dose every		Peak ≤40	baseline or >0.6	6/61 patients with peak
Retrospective	weight 1219 ± 516 g	6-36 hourly according to			mg/dL (53	concentration ≤40 and
observational		PMA			μmol/L)	none with peak
study (57)						concentrations >40
Goebel	n = 1	100 mg/kg	Not reported	12 hours post	Anuria. Not	Commenced
(1999)	Term infant, PNA 6 d			dose; no target	otherwise further	hemofiltration due to
Case report (58)	Solitary dysplastic kidney				defined	persistently elevated

						vancomycin level (240
						μg/mL). Normal renal
						function at PNA 3 weeks.
Müller	n = 2	Intermittent infusion:	None	9 hours post	Reduced GFR or	Transient
(1999)	PMA 35 weeks, weight	Single dose 35 mg/kg and		dose; no target	microproteinuria	microproteinuria in both
Case series (59)	1985 and 2390 g	38 mg/kg (accidental		reported	(urinary protein	patients but no change in
		overdose)			electrophoresis)	GFR
Pawlotsky	n = 53	n = 29	None	Post loading	Rise in SCr and	Transient rise in SCr in
(1998)	PMA $33.5 \pm 3.7$ weeks,	Loading dose: 7 mg/kg		dose; peak ≤40	urea not further	one patient with
Non-randomised	weight $1500 \pm 300 \text{ g (n =}$	Continuous infusion:			defined	klebsiella septicaemia
un-blinded trial	29)	10-40 mg/kg per over 24		Steady state		
(60)	PMA $33.9 \pm 4.8$ weeks,	hours according to PMA		(not further		
	weight $1800 \pm 800 \text{ g (n =}$	and weight		defined);		
	24)	n = 24		10-30		
		Continuous infusion:				

		10-30 mg/kg per over 24				
		hours according to PMA				
		and weight				
Sakata	n = 20	Intermittent infusion: 9-11	None	Timing and	Rise in SCr,	Transient rise in NAG
(1996)	PMA 26.3 ± 1.4 weeks	mg/kg per dose 12 hourly		target	fractional	index and FENa after
Prospective	Population not further			concentrations	excretion of	treatment in one patient
observational	defined			not reported	sodium (FENa)	
study (61)					and NAG index	
					(NAG:creatinine	
					ratio)	
McDougal	n = 44	Intermittent infusion:	Not reported	Steady state	Increase in SCr or	None detected
(1995)	PMA $29.4 \pm 0.9$ weeks,	15-18 mg/kg per dose 12-36		(not further	urea not further	
Prospective	weight $972 \pm 178 \text{ g (n = 16)}$	hourly according to PMA		defined); peak	defined	
observational	PMA $32.9 \pm 1.8$ weeks,			25-35, trough		
study (62)	weight $1379 \pm 382$ g (n =			5-10		
	15)					

	PMA $39.2 \pm 2$ weeks,					
	weight $2616 \pm 753$ g (n =					
	13)					
Tissing (1993)	n = 1	Intermittent infusion:	Tobramycin	Timing not	Rise in SCr and	Transient rise in SCr and
Case report (63)	PMA 29.3 weeks, weight	15 mg/kg per dose every 12		reported; no	oliguria not	persistently elevated
	1520 g	hours		target (reported	further defined	vancomycin level.
				level 63.3)		Resolution at 2 months
						of age.
Lisby-Sutch	n = 13	Intermittent infusion:	Not reported	Timing not	Increase in SCr or	None detected
(1988)	PMA 29.8 $\pm$ 3.4 weeks,	10 mg/kg per dose 6-12		reported; peak	urea not further	
Prospective	weight $1350 \pm 500 \text{ g}$	hourly according to PMA		25–35, trough,	defined	
PK study (64)		and weight		5–10		
Nahata	n = 61 < 1 year	Intermittent infusion:	Gentamicin	Timing not	Doubling of SCr	None detected
(1987)	Further clinical information	20-60 mg/kg/day (mean 35)		reported;	concentration	
	not available.			trough 2-18		

Prospective						
observational						
study (65)						
James	n = 20	Intermittent infusion:	Not reported	Timing not	Not formally	None reported although
(1987)	PMA $26.5 \pm 2.6$ weeks,	9-18 mg/kg per dose 12		reported; peak	defined	positive linear correlation
Prospective PK	weight 880 ± 340 g	hourly		30, trough 6		between vancomycin
study (33)						level and SCr identified
Dean	n = 28	Intermittent infusion:	Gentamicin	Timing not	Increase in SCr	Transient rise in SCr in
(1985)	Population not further	11-55 mg/kg (mean 30		reported; peak	by >0.5 mg/dl (45	2/19 and 2/9 patients
Retrospective	defined	mg/kg) daily not further		20-40, trough	μmol/L)	receiving vancomycin
observational		defined		5-10		alone and with
study (66)						gentamicin, respectively.