# trimETHOPRIM and Sulfamethoxazole

## **Newborn use only**

Alant	Not to be seed to see				
Alert	Not to be used in preterm infants until 4 weeks corrected gestational age.				
	Not to be used in term infants <4 weeks of age.				
	Term infants 4-8 weeks age: Watch for risk of kernicterus in high risk group or babies wi				
	1 -	jaundice.			
	Dose is expressed as trimethoprim (TMP) component.  The Antimicrobial Stewardship Team recommends this drug is listed under the following cate				
			wing category:		
Indiantian	Also known as co-trir				
Indication	Prophylaxis of urinary tract infections (UTI).				
		evere infections including UTI and acute otitis media.			
A -41	Prophylaxis in HIV-exposed infants.				
Action	Sulfamethoxazole is a sulfonamide that prevents the formation of dihydrofolic acid, a bacterial				
	compound necessary for survival. Trimethoprim is a synthetic antibiotic that interferes with the production of folic acid by inhibiting the action of dihydrofolate reductase.				
David trans		and by inhibiting the action of dinydroloiate reductase.			
Drug type	Antibiotic.				
Trade name	Oral: Septrin Sugar Free Oral liquid [Arrow]				
	IV: DBL Sulfamethoxazole 400 mg and Trimethoprim 80 mg Concentrate Injection BP [Pfizer]				
Presentation	-	prim 8 mg/mL and sulfamethoxazole 40 mg/mL, 100 mL bottle			
	IV: Trimethoprim 16 mg/mL and sulfamethoxazole 80 mg/mL, 5mL ampoule				
Dose	Dosage recommendations are based on trimethoprim component.				
	UTI prophylaxis				
	_	MP/kg/dose daily or 5 mg TMP/kg/dose twice weekly.			
		xposed infants <6 months of age	ana datha (nat nan la		
		ce from 4–6 weeks of age at a dose of 20 mg trimethoprim	once daily (not per kg		
		tes to 2.5 mL oral liquid daily)			
		evere infections (e.g. UTI, acute otitis media)			
		lerate infections			
		: 3–6 mg TMP/kg/dose 12 hourly (AAP Guidelines 2011).			
	Severe infections				
Dose adjustment	IV.	2–3 mg TMP/kg/dose 6 hourly.	1		
Dose aujustillelit		Renal Impairment Dose Adjustments			
	CrCl (mL/min)	Dosage			
	Above 25	Standard regimen			
	15 to 25	50% of the standard regimen			
	Below 15	Not recommended	•		
Maximum dose					
Total cumulative					
dose					
Route	Oral, IV				
Preparation	Oral: Oral liquid does not require preparation.				
	IV. Draw up 2 ml (22 mg trimothonnim and 00 mg sulfamath averally and add 40 ml of and in the column like it				
	-	mg trimethoprim and 80 mg sulfamethoxazole) and add 48 ml			
	0.9%, glucose 5% or glucose 10% to make a final volume of 50mL with a concentration of 0.64				
TMP.					
	For somewhy flyid restricted as supplies				
	For severely fluid restricted neonates:				
		g trimethoprim and 80 mg sulfamethoxazole) and add 18 mL of	_		
	final volume of 20mL with a final concentration of 1.6 mg/mL of TMP and infuse ONLY VIA A CENT				
	LINE as it is an alkaline solution. Flush the line with sufficient volume of sodium chloride 0.9% to ensure				
A 1	total dose is given.				
Administration					
	IV: Infuse over 60–90 minutes. Flush the line with sufficient volume of sodium chloride 0.9% to ensure				
	total dose is given.				
Monitoring	Watch for skin reactions and blood dyscrasias.  Monitor renal function and full blood count.				
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ANMF consensus group JHCH\_NICU\_19.084

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Contraindications	Hunarcancitivity to cultanamidae ar trimathanrim
Contraindications	Hypersensitivity to sulfonamides or trimethoprim.  Infants < 4 weeks of age
Precautions	Use with caution in renal impairment. Refer to dose adjustment section.
riecautions	In individuals with glucose-6-phosphate dehydrogenase deficiency, haemolysis may occur.
	Sulfamethoxazole may interfere with the serum albumin binding of bilirubin to produce kernicterus.
Drug interactions	Risk of prolonged QT interval with concurrent use of chloral hydrate, erythromycin and fluconazole.
Drug interactions	Increased effects and side effects of phenytoin (folate deficiencies) could occur when sulfamethoxazole/
	trimethoprim is given concurrently. Sulfamethoxazole/trimethoprim may inhibit the hepatic metabolism
	of phenytoin.
	Concomitant use of other agents that increase serum potassium, such as angiotensin converting enzyme
	inhibitors, angiotensin receptor blockers, potassium sparing diuretics and prednisolone can lead to
	hyperkalaemia.
	Increased sulfamethoxazole blood levels may occur in patients who are also receiving indomethacin.
	Cross sensitisation may exist between sulfamethoxazole/trimethoprim and some antithyroid agents,
	diuretics (thiazides) and oral hypoglycaemic drugs.
Adverse reactions	Gastrointestinal upset (vomiting, diarrhoea).
	Severe dermatologic reactions, blood dyscrasias, hepatotoxicity.
	Prolonged use may result in fungal or bacterial superinfection.
	Prolonged QT interval, torsades de pointes, ventricular tachycardias have been reported in adults.
	Severe cases of thrombocytopenia have been reported in adults.
Compatibility	Fluids <sup>17</sup> : Glucose 5%, glucose 10%, sodium chloride 0.9%, sodium chloride 0.45%.
	Y site (for dilutions of 1 in 25 only): Aciclovir, atracurium, dexmedetomidine, filgrastim, magnesium
	sulfate, morphine sulfate, piperacillin-tazobactam, vecuronium, and zidovudine.
	Y-site <sup>18</sup> (at 0.8 and 4mg/mL in glucose 5%): Aciclovir, amphotericin B liposome, azithromycin, cefepime,
	dexmedetomidine, filgrastim, linezolid, metronidazole, milrinone, octreotide, pamidronate,
	pancuronium, piperacillin-tazobactam, potassium acetate, remifentanil, sodium acetate, vecuronium,
	voriconazole, zidovudine.
Incompatibility	Fluids: No information. 17,18
	Y site <sup>17,18</sup> : Amikacin, aminophylline, amiodarone, amphotericin b lipid complex, ampicillin, atropine,
	benzylpenicillin, calcium chloride, calcium gluconate, cefazolin, cefotaxime, ceftazidime, ceftriaxone,
	chloramphenicol, clindamycin, dexamethasone, diazepam, diazoxide, digoxin, dobutamine, dopamine,
	adrenaline (epinephrine), epoetin alfa, erythromycin, fentanyl, fluconazole, folic acid, furosemide,
	ganciclovir, gentamicin, glycopyrrolate, hydralazine, hydrocortisone, imipenem-cilastatin, indomethacin,
	insulin, isoprenaline, ketamine, lidocaine (lignocaine), linezolid, methylprednisolone, metoclopramide,
	midazolam, multiple vitamins injection, nitroprusside sodium, noradrenaline (norepinephrine),
	phenobarbital (phenobarbitone), phenytoin, potassium chloride, propranolol, protamine, pyridoxine,
	ranitidine, sodium bicarbonate, ticarcillin-clavulanate, tobramycin, urokinase, vancomycin.
Stability	IV: infusion must be completed within 2 hours of preparation. Monitor for precipitation, particularly with
,	concentrated solutions.
Storage	Store IV and oral preparations below 30°C. Do not refrigerate. Protect from light.
	IV preparation: If stored at low temperatures precipitation may occur and solutions in which
	precipitation has occurred should be discarded.
Excipients	IV: diethanolamine, propylene glycol, alcohol, hydrochloric acid, sodium methabisulphate, sodium
	hydroxide.
	Oral: sorbitol, preservatives methyl hydroxybenzoate and sodium benzoate, ethanol, Cherry Flavour Artif
	F1242 (PI 286), sunset yellow, allura red, citric acid, cellulose, glycerol, polysorbate 80, sodium
	carmellose, saccharin sodium.
Cupaint some	
Special comments	Describedade in contract and have
Evidence	Prophylaxis in vesicoureteric reflux  The appearation of infente with high grade vesicoureteric reflux (VUR) are one all infente with fabrille UTIs is
	The proportion of infants with high grade vesicoureteric reflux (VUR) among all infants with febrile UTIs is
	small. There is no statistically significant benefit of prophylaxis in preventing recurrence of febrile
	UTI/pyelonephritis in infants without reflux. <sup>1</sup>

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There is benefit of prophylaxis in reducing the recurrence of infections in children with VUR but no change in renal scarring and concerns regarding multi-resistant strains among treated children.<sup>2,3</sup> **Treatment duration of infections** McMullan et al reviewed the evidence for minimum intravenous and total antibiotic duration in children younger than 18 years with bacterial infections, comparing shorter courses with traditionally longer durations. In many infections, especially when clinical improvement is rapid, emerging data suggest that traditional long durations of intravenous antibiotics might be unnecessary and that intravenous to oral switch can occur earlier. In most of the other infections, evidence for routine longer courses is sparse.<sup>4</sup> In a Cochrane review of childhood lower urinary tract infection, no difference in persistent bacteriuria or recurrence was noted between 2-4 days and 7-14 days of oral antibiotics. Results from a subsequent Cochrane review showed that a single-dose antibiotic was associated with more persistent bacteriuria than was 10 days of antibiotics, although there was no difference in symptom duration or recurrence. A large retrospective study of infants younger than 6 months found no difference in treatment failure between intravenous antibiotics for 3 days or less and 4 days or more.<sup>4-7</sup> Prophylaxis in HIV-exposed infants All HIV-exposed infants born to mothers living with HIV must receive co-trimoxazole prophylaxis, commencing at 4-6 weeks of age (or at first encounter with the healthcare system) and continued until HIV infection can be excluded.8 **Practice points** Roberts KB; Subcommittee on Urinary Tract Infection, Steering Committee on Quality Improvement and Management. References Urinary tract infection: clinical practice guideline for the diagnosis and management of the initial UTI in febrile infants and children 2 to 24 months. Pediatrics 128(3), 595-610(2011). de Bessa J Jr, de Carvalho Mrad FC, Mendes EF, Bessa MC, Paschoalin VP, Tiraboschi RB, Sammour ZM, Gomes CM, Braga LH, Bastos Netto JM. Antibiotic prophylaxis for prevention of febrile urinary tract infections in children with vesicoureteral reflux: a meta-analysis of randomized, controlled trials comparing dilated to nondilated vesicoureteral reflux. J Urol 2015; 193(5 Suppl):1772-7.  $P\'{e}rez-Gaxiola~G.~Antibiotic~prophylaxis~reduced~symptomatic~urinary~tract~infection~in~children~with~vesicoureteral~reflux,~but$ not scarring. Arch Dis Child Educ Pract Ed 2015; 100:52. McMullan BJ, Andresen D, Blyth CC, Avent ML, Bowen AC, Britton PN, Clark JE, Cooper CM, Curtis N, Goeman E, Hazelton B, Haeusler GM, Khatami A, Newcombe JP, Osowicki J, Palasanthiran P, Starr M, Lai T, Nourse C, Francis JR, Isaacs D, Bryant PA, ANZPID-ASAP group. 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Lieberthal AS, Carroll AE, Chonmaitree T, et al: The diagnosis and management of acute otitis media. Pediatrics 2013; 131(3):e964-e999. 13. Markowitz N & Saravolatz LD: Use of trimethoprim-sulfamethoxazole in a glucose-6-phosphate dehydrogenase-deficient population. Rev Infect Dis 1987; 9(suppl 2):S218-S225. Bell TAL, Foster JN, & Townsend ML: Trimethoprim-sulfamethoxazole-induced hepatotoxicity in a pediatric patient. Pharmacotherapy 2010; 30(5):539. Oliver RM, Rickenbach MA, Thomas MR, et al: Intrahepatic cholestasis associated with co-trimoxazole. Br J Clin Pract 1987; 41:975-976. 16. Paap CM & Nahata MC: Trimethoprim/sulfamethoxazole dosing during renal dysfunction. Ann Pharmacother 1995; 29:1300. 17. Australian injectable drugs handbook. Accessed online on 20 May 2021. 18. Micromedex online. Accessed on 20 May 2021.

VERSION/	/NUMBER
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## 2021

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Original 1.0	24/08/2016
Version 2.0	20/05/2021
Current 3.0	16/09/2021
REVIEW	16/09/2026

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