## **Sodium acetate**

### **Newborn use only**

Alert	In Australia, it is available as sodium acetate 16.4% (2 mmol/mL of acetate). It has an osmolarity of 4000
	mOsm/L.
	Concentrated sodium acetate ampoules <b>MUST BE DILUTED</b> prior to use.(1)
	Calculated osmolarity of sodium acetate – half strength, standard strength and high strength in this
	formulary are 160 mOsm/L, 320 mOsm/L and 1000 mOsm/L respectively. These osmolarities are similar to
	sodium chloride 0.45%, 0.9% and 3% respectively.(2, 3) (Refer to special comments section).
Indication	Metabolic acidosis: Prevention and treatment
	2. Hyponatraemia: An alternative source of correction in the presence of acidosis.
	3. Maintenance of arterial line or central venous line patency
Action	Acetate is an alkalinising agent and can be used to increase plasma bicarbonate concentration and correct
	metabolic acidosis. (4) Acetate is metabolised in the liver to bicarbonate.
Drug type	Electrolyte
Trade name	DBL Sodium acetate concentrated injection
Presentation	Sodium acetate concentrated injection 10 mL glass ampoule: Contains 1.64 gram/10 mL sodium acetate.
	This is equivalent to sodium acetate 16.4%.(1) Each 1 mL contains 2 mmol acetate and 2 mmol sodium.
Dose	Intravenous correction for metabolic acidosis
	1-3 mmol/kg/day.
	Dose beyond 3 mmol/kg/day may be used at the discretion of treating team.
	Arterial line or central venous line patency (ANMF consensus)
	< 1 Kg: sodium acetate half strength* with heparin 1 unit/mL at 0.5 mL/hour.
	1-1.5 Kg: sodium acetate <b>standard strength</b> * with heparin 1 unit/mL at 0.5 mL/hour.
	>1.5 kg with metabolic acidosis: sodium acetate <b>standard strength</b> * with heparin 1 unit/mL up to 1 mL/hour.
	*Half strength and standard strengths are similar in osmolarity to sodium chloride 0.45% and 0.9%
	respectively.
Dose adjustment	No information.
Maximum dose	No information.
Total cumulative	No information.
dose	
Route	Intravenous, intra-arterial.
Preparation	Intravenous correction for metabolic acidosis
	Sodium acetate – Standard strength*
	Add 4 mL of sodium acetate (8 mmol) to 46 mL of water for injection to make a final volume of 50
	mL with a concentration of 0.16 mmol/mL.
	1 mmol/kg/day = 0.26 ml/kg/hour
	Sodium acetate – High strength* (central line preferred)
	Add 12.5 mL of sodium acetate (25 mmol) to 37.5 mL of water for injection to make a final volume
	of 50 mL with a concentration of 0.5 mmol/mL (25 mmol/ 50 ml).
	1 mmol/kg/day = 0.08 ml/kg/hour
	*standard and high strengths are similar in osmolarity to sodium chloride 0.9% and 3%
	respectively.
	Autorial line an eartest responsible anchorum (benesis added)
	Arterial line or central venous line patency (heparin added)
	Sodium acetate – Half strength* (for weight <1 Kg):
	Draw up 2 mL of sodium acetate (equivalent to 4 mmol of acetate), add 5 mL of Heparinised
	Saline (50 units), and add to 43 mL of water for injection to make a final volume of 50 mL with a
	concentration of 0.08 mmol/mL of sodium acetate.
	Sodium acetate – Standard strength* (for weight ≥1 kg):
	Sodium acetate – Standard strength* (for weight ≥1 kg):  Draw up 4 mL of sodium acetate (equivalent to 8 mmol of acetate), add 5 mL of Heparinised
	Sodium acetate – Standard strength* (for weight ≥1 kg):

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	*Half strongt	and standard	strongths are simi	lar in asmalarity to sad	ium chlorido 0 4EV and 0 0V	
	*Half strength and standard strengths are similar in osmolarity to sodium chloride 0.45% and 0.9%					
	respectively.					
	Sodium and acetate in	mmol/kg/day	with the above inf	usions for intra-arterial	/central venous line	
	patency:				, 50.11. 4. 10.1045	
	Weight	Sodium ac	etate strength	Rate	mmol/kg/day	
	500 g				1.9 mmol/kg/day	
	750 g	Half	strength	0.5 mL/hour	1.2 mmol/kg/day	
	1000 g				0.9 mmol/kg/day	
	500 g				3.8 mmol/kg/day	
	750 g				2.5 mmol/kg/day	
	1000 g	Standa	rd strength	0.5 mL/hour	1.9 mmol/kg/day	
	2000 g				0.95 mmol/kg/day	
Administration	Continuous infusion				0.000., 1.8, 0.07	
Monitoring	Electrolytes, acid base	status (bicarbo	nate, base excess.	nCO2)		
Contraindications	Hypernatraemia	(10.000	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	, ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		
	Fluid overload					
Precautions	Renal impairment					
Drug interactions	,					
Adverse	Metabolic alkalosis					
reactions	Hypernatraemia					
	Fluid overload					
	Aluminium toxicity fro	m leaching of a	lluminium from gla	ass ampoules. (5)		
Compatibility	Fluids: Glucose 5%, so	dium chloride (	0.9%, Amino acid s	olutions, lipid emulsion	(6)	
	Y site: aciclovir, alfent	anil, allopurino	l, amifostine, amik	acin, aminophylline, am	picillin, anidulafungin,	
					busulfan, calcium folinate,	
	calcium gluconate, capreomycin, cefazolin, cefepime, cefotaxime, cefoxitin, ceftazidime, ceftriaxone, cefuroxime, clindamycin, dexamethasone, dexmedetomidine, digoxin, diltiazem, diphenhydramine,					
	dobutamine, dopamine, doxycycline, enalaprilat, ephedrine, adrenaline (epinephrine), erythromycin					
	lactobionate, esmolol, fentanyl, fluconazole, fluorouracil, foscarnet, fosphenytoin, furosemide, ganciclovir,					
	gentamicin, heparin, hydrocortisone, imipenem-cilastin, labetalol, levofloxacin, lidocaine (lignocaine),					
	linezolid, lorazepam, magnesium sulfate, methadone, methotrexate, methylprednisolone, metronidazole,					
	milrinone, morphine, naloxone, netilmicin, nitroprusside sodium, octreotide, ondansetron, pamidronate, pancuronium, pentobarbital, phenobarbital (phenobarbitone), phenylephrine, piperacillin-tazobactam,					
	potassium chloride, propranolol, ranitidine, remifentanil, rocuronium, sodium bicarbonate,					
	suxamethonium, sulfamethoxazole-trimethoprim, tacrolimus, theophylline, ticarcillin, tobramycin,					
	vancomycin, vasopres		•		,,	
Incompatibility	Fluids: No information		, ,	,		
	Y site: Amiodarone, ar	nphotericin B c	onventional colloid	dal and lipid complex, c	aspofungin, diazepam,	
	hydralazine, mycophe	nolate mofetil,	pantoprazole, phe	nytoin		
Stability						
Storage	Store below 30°C. Sing	le use only. Re	place syringe every	y 24 hours.		
Excipients	Water for injection					
Special						
comments	Solution	1	Flectroly	te (mmol/mL)	O       1 \	
	Joidtio		Liectioly		Osmolarity (mOsm/L)	
	Human Pla		Liectiony		280-300	
		sma	-	ol/mL of Na		
	Human Pla	sma e 16.4%	2 mm	ol/mL of Na mol/mL of Na	280-300	
	Human Pla Sodium acetat	sma e 16.4% e 0.45%	2 mmc	•	280-300 4000	
	Human Pla Sodium acetat Sodium chlorid	sma e 16.4% e 0.45% de 0.9%	2 mm 0.08 mr 0.15 mr	mol/mL of Na	280-300 4000 154	
	Human Pla Sodium acetate Sodium chlorid Sodium chlorid	sma e 16.4% e 0.45% de 0.9% de 3%	2 mm 0.08 mr 0.15 mr 0.51 mr	nol/mL of Na nol/mL of Na	280-300 4000 154 308	

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	Sodium acetate high strength	0.5 mmol/mL of Na and acetate	1000	
	Sodium bicarbonate 8.4%	1 mmol/mL of Na and bicarbonate	2000	
	Sodium bicarbonate 4.2%	0.5 mmol/mL of Na and bicarbonate	1000	
Evidence	Background  Sodium acetate is similar to bicarbonate in its ability to restore blood pH and plasma bicarbonate. (7) It can also be used as the source of sodium in parenteral nutrition solution in preterm neonates.  Efficacy  In a prospective study by Ekblad et al, 11 infants ≤ 34 weeks were supplemented with sodium acetate added to the daily intravenous fluids from day 1 of life. Sodium acetate was used as the sole source of sodium on day 1 of life and both sodium chloride and sodium acetate were used in equal amounts as the source of sodium from day 2 of life. Actual intakes of sodium acetate on day 1 and thereafter were 3 mmol/kg/day and 1.5 mmol/kg/day respectively. They demonstrated an improvement in metabolic acidosis (less number of infants with pH < 7.3) without any worsening in PCO₂. Serum sodium was normal in all infants.(8) In a double blind randomised controlled trial, Ali et al compared the parenteral nutrition (PN) solutions containing sodium acetate or sodium chloride on biochemical parameters and clinical outcomes in 52 infants < 33 weeks including 29 extremely low birth weight infants <1000 g. PN was prepared based on 2005 ESPGHAN guidelines. The intervention arm received sodium acetate as the entire source of sodium whereas the control arm received sodium chloride as the source of sodium. In the first 6 days of life, intervention arm received mean intake of sodium (and acetate) 4 mmol/kg/day. Blood pH and base excess rose to normal values after 3 days of PN in the acetate group. There was no significant difference in pCO₂ between groups. There was a significantly lower incidence of bronchopulmonary dysplasia in the acetate group. There was also a trend towards lower incidence of severe intraventricular haemorrhage.(7)			
	Pharmacokinetics			
	Following administration acetate is m	netabolised in liver to bicarbonate.		
Practice points				
References	kabi.us/Pls/Sodium_Ace_Inj_4582 2. 0.45% sodium chloride injection, U 3. 0.9% sodium chloride injection, U 4. DBL Sodium Acetate Concentrated 5. Sodium acetate. IBM Micromedex 6. Sodium acetate. Australian Injecta 7. Ali A, Ong E-Y, Singh BKS, Cheah F parenteral nutrition for very prete Gastroenterology, Hepatology & N 8. Ekblad H, Kero P, Takala J. Slow so	USP. Accessdata.fda.gov. SP. Accessdata.fda.gov. d Injection. Accessed via MIMS online on 8 c. Accessed online on 14 February 2022. able Drugs Handbook. Accessed online on -C. Comparison between sodium acetate actern infants on the acid-base status and ne	3 February 2022. [Internet].  14 February 2022.  and sodium chloride in conatal outcomes. Pediatric	

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