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Alert	Osmolarity: Sodium chloride 20%: 6846 mOsm/L ¹ . High risk of extravasation if administered undiluted.		
	Sodium supplementation is not always appropriate and fluid restriction may be appropriate in the		
	management of hyponatraemia. Treatm	ent should always be tailored to the cause.	
Indication	Treatment of hyponatraemia.		
Action	Sodium is the major cation in extracellula	ar fluid.	
Drug type	Sodium is the major cation in extracellular hald. Sodium chloride 20% contains 200 g/L sodium chloride, equivalent to 3.4 mmol/mL of sodium.		
Trade name	Sodium chloride 20%		
Presentation	Sodium chloride 20% Sodium chloride 20% – 10 mL ampoule. Can be used for both IV and oral routes. Refer to administration		
Presentation	section.	Lan be used for both iv and oral routes. Refer to administration	
Dose		cumptomatic hypopatraomia	
Dose	Severe hyponatraemia < 120 mmol/L or symptomatic hyponatraemia		
	IV: CAUTION—CANNOT BE GIVEN UNDILUTED. REFER TO PREPARATION/DILUTION SECTION FOR DETAILS. Infuse sodium chloride at 0.4 mmol/kg/hour until symptoms abate or sodium ≥ 120 mmol/L.		
	Then infuse sodium chloride at 0.15 mmol/kg/hour for 48 hours or until desired sodium is achieved.		
	Therapeutic goal is to increase sodium by 7 mmol/L/day.		
	IV supplementation		
	Start at 2–4 mmol/kg/day and increase as required. Oral supplementation		
		g/day) and increase as required, divided into 3–12 doses.	
Dose adjustment	Therapeutic hypothermia – No informati	on.	
	ECMO – No information.		
	Renal impairment – No information.		
	Hepatic impairment – No information.		
Maximum dose			
Total cumulative			
dose			
Route	IV, PO		
NOULE			
	IV infusion:		
Preparation		um chloride 20% and add 44 mL of water for injection to make a tration of 0.4 mmol/mL. Infusion at a rate of 1 mL/kg/hour = 0.4	
	Draw up 6 mL (20 mmol sodium) of sodium final volume of 50 mL with a final concer mmol/kg/hour (9.6 mmol/kg/day).	tration of 0.4 mmol/mL. Infusion at a rate of 1 mL/kg/hour = 0.4	
	Draw up 6 mL (20 mmol sodium) of sodium final volume of 50 mL with a final concer mmol/kg/hour (9.6 mmol/kg/day). Infusion Strength	tration of 0.4 mmol/mL. Infusion at a rate of 1 mL/kg/hour = 0.4 Prescribed amount	
	Draw up 6 mL (20 mmol sodium) of sodium final volume of 50 mL with a final concer mmol/kg/hour (9.6 mmol/kg/day).	Prescribed amount 6 mL of sodium chloride 20% and make up to 50 mL of	
	Draw up 6 mL (20 mmol sodium) of sodium final volume of 50 mL with a final concer mmol/kg/hour (9.6 mmol/kg/day). Infusion Strength 1 mL/kg/hour = 0.4 mmol/kg/hour	Prescribed amount 6 mL of sodium chloride 20% and make up to 50 mL of water for injection	
	Draw up 6 mL (20 mmol sodium) of sodium final volume of 50 mL with a final concer mmol/kg/hour (9.6 mmol/kg/day). Infusion Strength 1 mL/kg/hour = 0.4 mmol/kg/hour	Prescribed amount 6 mL of sodium chloride 20% and make up to 50 mL of	
	Draw up 6 mL (20 mmol sodium) of sodium final volume of 50 mL with a final concer mmol/kg/hour (9.6 mmol/kg/day). Infusion Strength 1 mL/kg/hour = 0.4 mmol/kg/hour *1 mL/kg of 0 .4 mmol/mL of sodium chl	Prescribed amount 6 mL of sodium chloride 20% and make up to 50 mL of water for injection	
	Draw up 6 mL (20 mmol sodium) of sodiu final volume of 50 mL with a final concer mmol/kg/hour (9.6 mmol/kg/day). Infusion Strength 1 mL/kg/hour = 0.4 mmol/kg/hour *1 mL/kg of 0 .4 mmol/mL of sodium chl Oral: Sodium chloride 20% oral solution	Prescribed amount 6 mL of sodium chloride 20% and make up to 50 mL of water for injection oride will raise serum sodium by 0.8 mmol/L.² (prepared in-house by pharmacy decanting 20% sodium chloride	
Preparation	Draw up 6 mL (20 mmol sodium) of sodium final volume of 50 mL with a final concer mmol/kg/hour (9.6 mmol/kg/day). Infusion Strength 1 mL/kg/hour = 0.4 mmol/kg/hour *1 mL/kg of 0 .4 mmol/mL of sodium chl	Prescribed amount 6 mL of sodium chloride 20% and make up to 50 mL of water for injection oride will raise serum sodium by 0.8 mmol/L.² (prepared in-house by pharmacy decanting 20% sodium chloride	
	Draw up 6 mL (20 mmol sodium) of sodius final volume of 50 mL with a final concer mmol/kg/hour (9.6 mmol/kg/day). Infusion Strength 1 mL/kg/hour = 0.4 mmol/kg/hour *1 mL/kg of 0 .4 mmol/mL of sodium chl Oral: Sodium chloride 20% oral solution from IV ampoules into bottles for oral do IV:	Prescribed amount 6 mL of sodium chloride 20% and make up to 50 mL of water for injection oride will raise serum sodium by 0.8 mmol/L.² (prepared in-house by pharmacy decanting 20% sodium chloride	
Preparation	Draw up 6 mL (20 mmol sodium) of sodiu final volume of 50 mL with a final concer mmol/kg/hour (9.6 mmol/kg/day). Infusion Strength 1 mL/kg/hour = 0.4 mmol/kg/hour *1 mL/kg of 0 .4 mmol/mL of sodium chl Oral: Sodium chloride 20% oral solution from IV ampoules into bottles for oral do	Prescribed amount 6 mL of sodium chloride 20% and make up to 50 mL of water for injection oride will raise serum sodium by 0.8 mmol/L.² (prepared in-house by pharmacy decanting 20% sodium chloride	
Preparation	Draw up 6 mL (20 mmol sodium) of sodium final volume of 50 mL with a final concer mmol/kg/hour (9.6 mmol/kg/day). Infusion Strength 1 mL/kg/hour = 0.4 mmol/kg/hour *1 mL/kg of 0 .4 mmol/mL of sodium chl Oral: Sodium chloride 20% oral solution from IV ampoules into bottles for oral do IV: IV infusion.	Prescribed amount 6 mL of sodium chloride 20% and make up to 50 mL of water for injection oride will raise serum sodium by 0.8 mmol/L.² (prepared in-house by pharmacy decanting 20% sodium chloride	
Preparation	Draw up 6 mL (20 mmol sodium) of sodium final volume of 50 mL with a final concer mmol/kg/hour (9.6 mmol/kg/day). Infusion Strength 1 mL/kg/hour = 0.4 mmol/kg/hour *1 mL/kg of 0 .4 mmol/mL of sodium chl Oral: Sodium chloride 20% oral solution from IV ampoules into bottles for oral do IV: IV infusion. Oral:	Prescribed amount 6 mL of sodium chloride 20% and make up to 50 mL of water for injection oride will raise serum sodium by 0.8 mmol/L.² (prepared in-house by pharmacy decanting 20% sodium chloride ising).	
Preparation	Draw up 6 mL (20 mmol sodium) of sodius final volume of 50 mL with a final concer mmol/kg/hour (9.6 mmol/kg/day). Infusion Strength 1 mL/kg/hour = 0.4 mmol/kg/hour *1 mL/kg of 0 .4 mmol/mL of sodium chl Oral: Sodium chloride 20% oral solution from IV ampoules into bottles for oral do IV: IV infusion. Oral: To be given mixed with feeds.	Prescribed amount 6 mL of sodium chloride 20% and make up to 50 mL of water for injection oride will raise serum sodium by 0.8 mmol/L.² (prepared in-house by pharmacy decanting 20% sodium chloride ising).	
Preparation	Draw up 6 mL (20 mmol sodium) of sodiu final volume of 50 mL with a final concer mmol/kg/hour (9.6 mmol/kg/day). Infusion Strength 1 mL/kg/hour = 0.4 mmol/kg/hour *1 mL/kg of 0 .4 mmol/mL of sodium chl Oral: Sodium chloride 20% oral solution from IV ampoules into bottles for oral do IV: IV infusion. Oral: To be given mixed with feeds. IV: Signs of extravasation at IV insertion	Prescribed amount 6 mL of sodium chloride 20% and make up to 50 mL of water for injection oride will raise serum sodium by 0.8 mmol/L.² (prepared in-house by pharmacy decanting 20% sodium chloride using).	
Preparation	Draw up 6 mL (20 mmol sodium) of sodius final volume of 50 mL with a final concer mmol/kg/hour (9.6 mmol/kg/day). Infusion Strength 1 mL/kg/hour = 0.4 mmol/kg/hour *1 mL/kg of 0 .4 mmol/mL of sodium chl Oral: Sodium chloride 20% oral solution from IV ampoules into bottles for oral do IV: IV infusion. Oral: To be given mixed with feeds. IV: Signs of extravasation at IV insertion Oral: Watch for signs of gastric irritation Serum sodium as per clinical team's reco	Prescribed amount 6 mL of sodium chloride 20% and make up to 50 mL of water for injection oride will raise serum sodium by 0.8 mmol/L.² (prepared in-house by pharmacy decanting 20% sodium chloride using).	

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Precautions	Impaired renal function, cardiac insufficiency, pre-existing oedema with sodium retention.
Drug interactions	No information.
Adverse reactions	Hypernatraemia, volume overload, congestive heart failure, respiratory distress
	Hyperchloraemia, hypercalciuria
	Disseminated intravascular coagulation (DIC) is associated with inadvertent injections of sodium chloride
	into blood vessels of the uterus or placenta due to hypernatraemic shock. Not reported in infants.
	Osmotic demyelinating syndrome.
	Fever
	IV site: Extravasation, phlebitis, venous thrombosis.
Camanatibility	Oral: Gastric irritation.
Compatibility	IV Fluids: Glucose 5%, glucose 10%, glucose 5% in sodium chloride 0.9%, glucose 5% in sodium chloride
	0.45%, sodium chloride 0.9%, sodium chloride 0.45%. Y site: No information.
Incompatibility	IV Fluids: Fat emulsion.
meompationity	Y site: No information.
	Amino Acid solutions – No information.
Stability	PO: Expiry 8 days from manufacture.
Storage	IV: Store at room temperature, 20–25°C.
	PO: Refrigerate (2–8°C).
Excipients	
Special comments	Osmolarity of undiluted hypertonic sodium chloride is >1000 mOsm/L, posing the risk of extravasation for
	peripheral IV solutions. ^{3,4} So, local consensus was to bring the osmolarity of IV preparation to 2.4% sodium
	chloride that has 0.4 mmol/L of sodium and an estimated osmolarity of 855 mOsm/L.
	Total body water is traditionally calculated as weight x 0.6 in children. Greater total body water content in
Evidence	newborns should be considered and therefore should be calculated as weight x 0.75. ^{2,5} IV correction for severe and/or symptomatic hyponatraemia
Evidence	The body of evidence to base recommendations in this clinical setting is extremely limited, particularly in
	neonatal populations. Recommendations are based on expert opinion, which have been extrapolated from
	adult consensus guidelines ^{6,7} and take into account specific neonatal safety concerns (see Safety below). In
	acute hyponatraemia, where the risk of sequelae is greater than that of osmotic demyelination, the
	correction should be rapid. ⁸
	Aim to increase sodium by 1–2 mmol/L per hour until symptoms abate or a safe level of sodium is
	achieved (≥ 120 mmol/L). Once the safe level is achieved, suggested subsequent goals are 6–8 mmol/L in
	24 hours, 12–14 mmol/L in 48 hours and 14–16 mmol/L in 72 hours. 10 (LOE IV, GOR C)
	Dosage and infusion rate recommendations in this formulary are extrapolated from the rate of rise
	expected with sodium chloride 3% ² and are as follows:
	0.5 mmol/mL of sodium chloride (i.e. sodium chloride 3%), when administered at 1 mL/kg, will raise serum
	sodium by 1 mmol/L.
	0.4 mmol/mL of sodium chloride (i.e. diluted sodium chloride in this formulary), when administered at 1
	mL/kg, will raise serum sodium by 0.8 mmol/L.
	Sodium deficit calculation
	Deficit in mmol = (desired sodium – serum sodium) x total body water
	Total body water is traditionally calculated as weight x 0.6 in children. Greater total body water content in
	newborns should be considered and therefore should be calculated as weight x 0.75. 2.5 (LOE IV, GOR C)
	(22.1) 33.6.6
	<u>Oral supplementation</u>
	A randomised, controlled trial of 4 mmol/kg/d (0.4 mL/kg per dose of 2.5 mmol/mL sodium chloride) of
	sodium versus placebo from DOL 7 to 35 in infants born 24–31 weeks (53 infants) showed higher sodium
	levels and increased weight gain in the intervention group. 11 A randomised, controlled trial of 4 mmol/kg/d
	(concentration not specified) of sodium versus placebo from DOL 4 to 14 in infants born at 29–34 weeks
	(20 infants) showed higher sodium levels and increased weight gain in the intervention group. 12 There are
	also three case–control studies that report similar findings with respect to sodium levels and growth in

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preterm infants supplemented with oral sodium.¹³⁻¹⁵ A systematic review comparing higher versus lower sodium intake for preterm infants is in progress.¹⁶ These findings support the use of oral sodium supplements to correct hyponatraemia and potentially improve growth. (LOE II, GOR B)

Safety

An historical, case-control study identified 42/350 ELBW NICU admissions with an episode of hyponatraemia (Na < 125 mmol/L [range 113–124]) that lasted > 6 hours (median 1.5 days). Rates of abnormal head ultrasound (IVH or PVL) and abnormal neurological examination were higher in the hyponatraemic group (p < 0.03; p < 0.001 respectively). Correction \geq 0.5 mmol/L/h showed a trend toward higher rates of abnormal neurological examination. In paediatric and adult populations, multiple cohort studies and reviews have concluded that in patients with chronic hyponatraemia (\geq 48 hours), neurologic sequelae due to osmotic demyelination are associated with more rapid rates of correction. Page 113–124]

In summary, rapid correction of hyponatraemia may be detrimental to neurological outcome during myelination of the newborn brain. 17 In adult populations, osmotic demyelination syndrome can usually be avoided by limiting correction of chronic hyponatraemia to < 10 to 12 mmol/L in 24 hours and to < 18 mmol/L in 48 hours. These estimates should be regarded as approximate limits and not goals of therapy. 7 (LOE IV, GOR C)

Osmolarity and Osmolar load

A retrospective, matched-cohort study of 352 children \leq 18 years evaluated the incidence of phlebitis or infiltration associated with peripheral administration of parenteral nutrition with an osmolarity > 1000 mOsm/L vs \leq 1000 mOsm/L. There were 151 neonates in the study. There were no differences between patients who did or did not develop adverse events in terms of age or weight. Administration of PPN with osmolarity > 1000 mOsm/L vs \leq 1000 mOsm/L significantly increased infiltration (17% vs 7%; odds ratio [OR, 2.47]; 95% confidence interval [CI], 1.24–4.94; p = 0.01) and the combined composite end point of phlebitis or infiltration (45% vs 34%; OR, 1.65; 95% CI, 1.07–2.54; p = 0.02). In multivariate analysis, osmolarity > 1000 mOsm/L vs \leq 1000 mOsm/L was an independent risk factor for developing complications (OR, 1.67; 95% CI, 1.08–2.52; p = 0.02). (LOE III, GOR C)

A prospective, observational study in adults suggests that osmolar load (i.e. number of milliosmoles per hour, calculated as osmolarity x infusion rate) is a better predictor than osmolarity alone for phlebitis. They found an osmolarity rate of 84–99 mOsm/hour was associated with 4–27% rate of phlebitis. They did not report on other injuries such as extravasation. The infusion rates suggested in our formulary have low osmolar load and are considered to carry minimal risk of phlebitis (Consensus opinion).

Practice points

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VERSION/NUMBER	DATE
Original 1.0	06/09/2017
Current 2.0	15/12/2020
REVIEW (5 years)	15/12/2025

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