Alert	Most often given in conjunction with calcium for the prevention and treatment of metabolic bone	
	disease in preterm infants.	
	1 mmol phosphorus/phosphate (P) = 31 mg elemental phosphorus.	
	1 mmol elemental calcium (Ca) = 40 mg elemental calcium.	
	Separate oral doses from calcium supplements by at least 1 hour.	
	When using IV preparation, always check plasma sodium and potassium concentrations to assist in	
	choosing the right phosphate preparation (e.g. sodium or potassium phosphate preparation).	
Indication	Treatment of Metabolic Bone Disease.	
	Treatment of hypophosphataemia.	
	Supplementation to meet the recommended daily intakes.	
Action	Phosphorus is a major intracellular mineral and is important in bone mineralisation and energy	
	production.	
Drug Type	Mineral	
Trade Name	IV .	
	Glycophos® Concentrated injection solution for infusion (Fresenius-Kabi) (recommended organic	
	preparation)	
	Each 1 mL of Glycophos® corresponds to 1 mmol phosphate and 2 mmol sodium.	
	Sodium dihydrogen phosphate Phebra IV (Preferred inorganic preparation)	
	Each 1 mL vial corresponds to 1 mmol phosphate, 1 mmol sodium and 2 mmol hydrogen.	
	Potassium dihydrogen phosphate concentrated injection DBL IV	
	Potassium dihydrogen phosphate concentrated injection Phebra IV	
	Each 1 mL ampoule corresponds to 1 mmol phosphate, 1 mmol potassium and 2 mmol hydrogen.	
	ORAL	
	Phosphate-Phebra® oral effervescent tablets	
	Each tablet contains: 16.1 mmol phosphate (equivalent to 500 mg elemental phosphorus); 20.4 mmol	
	Sodium; 3.1 mmol potassium Sodium dihydrogen phosphate Phebra IV (preferred IV preparation) Each 10 mL vial (sodium dihydrogen phosphate 1.56 g) contains: 10 mmol phosphate; 10 mmol sodium;	
	20 mmol hydrogen	
	Potassium dihydrogen phosphate concentrated injection DBL IV Potassium dihydrogen phosphate concentrated injection Phebra IV Forb 40 order to the concentrated injection Phebra IV	
	Each 10 mL ampoule (potassium dihydrogen phosphate 1.361 g) contains: 10 mmol phosphate; 10 mmo	
Presentation	potassium; 20 mmol hydrogen IV: Glycophos 20 mL ampoule; Sodium dihydrogen phosphate 10 mL vial; Potassium dihydrogen	
Presentation	phosphate concentrated injection 10 mL ampoule.	
	Oral: 500 mg effervescent tablets; IV preparation (e.g. sodium or potassium dihydrogen phosphate) can	
	be given orally.	
	be given ordiny.	
Dose	Treatment of metabolic bone disease (MBD)	
	(= /	
	PO: 1 to 3 mmol/kg/day in 2-4 divided doses as an addition to intake from milk and other	
	sources to a maximum intake of 4.5 mmol/kg/day.	
	Use either Sodium dihydrogen phosphate Phebra IV preparation or Phosphate-Sandoz tablets.	
	General principles of treatment of MBD:	
	A. Commence at low dose (e.g. 1 mmol/kg/day) and titrate the dose up as tolerated.	
	B. Given in conjunction with calcium supplementation (but not together - example: Calcium 8	
	AM, 2 PM, 8 PM and Phosphorus 6 AM, 12 MD, 6 PM)	
	C. Aim to reach the upper end of the recommended intake: Ca 5 mmol/kg/day and P 4.5	
	mmol/kg/day. ⁸	

	 D. Dose can be adjusted with a goal of slight excess supply aiming for urinary calcium ≥1.2mmol/L and phosphate ≥0.4 mmol/L.
	Treatment of acute hypophosphataemia IV: 0.2 mmol/kg/dose [range 0.15–0.33 mmol/kg/dose] over 6 hours. Repeat as necessary. Aim to maintain normophosphataemia of 1.8–2.6 mmol/L (5.6–8.1 mg/dl).
	Daily enteral Supplementation to meet the recommended daily intakes (RDI) 2–4.5 mmol/kg/day (62–140 mg/kg/day of phosphorus) ^{7,8}
	 Calculate intake from parenteral and enteral sources Supplement the difference via IV or oral route.
Dose adjustment	
Maximum dose	
Total cumulative dose	
Route	PO IV
Preparation	IV infusion for treatment of acute hypophosphataemia: IV infusion (Glycophos): Draw up 1 mL (1 mmol phosphate) and add 19 mL sodium chloride 0.9% or water for injection to make a final volume of 20 mL with a concentration of 0.05 mmol/mL. Draw up 4 mL/kg (0.2 mmol/kg).
	IV infusion (sodium dihydrogen phosphate): Draw up 1 mL (1 mmol phosphate) and add 19 mL sodium chloride 0.9% or glucose 5% to make a final volume of 20 mL with a concentration of 0.05 mmol/mL. Draw up 4 mL/kg (0.2 mmol/kg).
	IV infusion (potassium dihydrogen phosphate): Draw up 1 mL (1 mmol phosphate) and add 24 mL sodium chloride 0.9% or glucose 5% to make a final volume of 25 mL with a concentration of 0.04 mmol/mL. Draw up 5 mL/kg (0.2 mmol/kg).
	Oral Option 1 (preferred option for infants going home or when a long storage time is required in the NICU): Disperse 500 mg (16.1 mmol) Phosphate-Sandoz in 16 mL of water for injection to make a solution with a concentration of 1 mmol/mL.
	Option 2 (can be used where preparation with low osmolality is preferred e.g. infants with history of feed intolerance): IV sodium dihydrogen phosphate decanted into a bottle and given orally undiluted (expiry time: 7 days).
Administration	Oral Can be administered with feeds (refer to evidence summary section). Separate calcium supplements by at least 2 hours.
	IV As part of parenteral nutrition fluid – refer to individual parenteral nutrition formulations.
	IV infusion for treatment of acute hypophosphataemia: IV glycophos: Infuse over at least 8 hours. IV sodium dihydrogen phosphate or IV potassium dihydrogen phosphate: Infuse over at least 6 hours.
	For severe hypophosphataemia infuse over 8–12 hours. Maximum infusion rate of 0.2 mmol/kg/h.
Monitoring	Phosphate, calcium, magnesium, alkaline phosphatase concentrations are required at least fortnightly or more often if required. Once these concentrations normalise, serum analysis may be performed once monthly for 6 months or at the discretion of the clinician. ¹⁰ Urinary calcium and phosphate and Tubular Reabsorption Phosphate (TRP)%, parathormone, and vitamin D concentrations may be useful under certain circumstances.

Contraindications	Hyperphosphataemia, dehydration, se	vere renal insufficiency, sh	ock.
Precautions	Hypernatraemia (avoid sodium dihydro	ogen phosphate).	
	Hyperkalaemia (avoid potassium dihyd		
Orug Interactions	Calcium and magnesium antacids (e.g.		e. hvdroxide etc.) reduce phosphate
rug meeraations	absorption — separate doses by at lea		e, ilyaroxiae etci, redace priospriate
	Additive effects with other drugs that		
	Potassium dihydrogen phosphate prep		risk of hynerkalaemia when used in
	conjunction with potassium sparing di	-	
Adverse	Diarrhoea (oral use only), hypocalcaen		
Reactions	hypomagnesaemia.	ma, mepin ocomorey, protong	sea Qt meer val, mypotension,
	Hyperphosphataemia – carpopedal spa	asm. seizures. ²	
Compatibility	Glycophos		
,	Fluids: Sodium chloride 0.9%, water fo	r injection, glucose 5%.	
	Y-site: No iformation.		
	Potassium dihydrogen phosphate		
	Fluids: Glucose 5%, glucose 10%, gluco	se in sodium chloride solut	tions, sodium chloride 0.45%, sodiun
	chloride 0.9%, sodium chloride 3%.		
	Y-site: No information.		
	Sodium dihydrogen phosphate		
	Fluids: Glucose 5%, sodium chloride 0.	9%.	
	Y-site: No information		
ncompatibility	Potassium dihydrogen phosphate		
, ,	Fluids: No information		
	Drugs: Aciclovir, amiodarone, calcium salts, ketamine, lorazepam, magnesium salts, rocuronium.		
	Solutions that contain other cations such as calcium, magnesium, iron and aluminium may also		
	precipitate.		
	Sodium dihydrogen phosphate		
	Fluids : No information		
	Drugs: Aciclovir, amiodarone, calcium	salts, calcium, aluminium c	or magnesium, iron and magnesium
	containing solutions.		
Stability	Preparation from oral effervescent tablets: It is to be used immediately after preparation and discard		
	unused portion.		
	Oral preparation from IV sodium dihyc	Irogen phosphate: 7 days	
	Glycophos: To be used within 24 hours	after reconstitution.	
Storage	Store below 25°C.		
Excipients	Phosphate-Phebra® oral effervescent t	tablets: Sodium bicarbonat	e, potassium bicarbonate, macrogol
-	4000, citric acid, sucrose, orange 5257		
	Glycophos: Hydrochloric acid and water		
Special			
Comments			
vidence	Recommended daily intakes (RDI)		
	Phosphorus absorption is typically 80% to 90% of dietary intake. ³		
	Parenteral intake: Previously, the recommended doses of parenteral Ca and P in preterm infants varied		
	from 1.3–3 mmol Ca/kg/day and 1.0–2.3 mmol P/kg/day, with a Ca:P ratio in the range of 1.3–1.7. ^{1,4-6} ESPGHAN 2018 updated guidelines on parenteral nutrition recommends the following Ca and		
	Phosphate:12		
		Parenteral Ca	Parenteral Ph
		mmol (mg)/kg/day	mmol (mg)/kg/day
	Preterm during the first days of life	0.8-2.0 (32-80)	1.0-2.0 (31-62)
	<u> </u>		<u> </u>
	Growing preterm	1.6-3.5 (100-140)	1.6-3.5 (77-108)
	Growing preterm Term neonate	1.6-3.5 (100-140) 0.8-1.5 (30-60)	1.6-3.5 (77-108) 0.7-1.3 (20-40)

Newborn use only

Enteral intake: ESPGHAN 2010 Guidelines for enteral nutrition recommend 2–3 mmol/kg/day of a highly absorbable phosphate source in a ratio with calcium (Ca:P) of 1.5–2.0.⁷ American Academy of Pediatrics Committee on Nutrition 2013 Guidelines recommend Ca 150-200 mg/kg/day (3.8-5 mmol/kg/day) and P 75-140 mg/kg/day (2.4-4.5 mmol/kg/day) and 200-400 IU/day of vitamin D for enteral nutrition in preterm neonates.⁸

The exact serum phosphorus concentration at which to commence supplementation of phosphate is not known and recommendations vary from 1.3 mmol/L⁸ to 1.8 mmol/L.⁹

Metabolic bone disease

Goal: Aim for the upper end of the recommended range to prevent fractures and clinical symptoms of osteopenia: Ca and P of around 4-4.5 mmol/kg/day. Adjust the mineral intake with a goal of achieving a slight excess of urinary mineral excretion: Urinary calcium \geq 1.2mmol/L and phosphate \geq 0.4 mmol/L.

Step 1: Calculate the mineral intake from enteral feed:

Example: 150 ml/kg/day of mature preterm EBM contains: Ca 1 mmol/kg/day and P 0.6 mmol/kg/day. 150 ml/kg/day preterm EBM+24kcal HMF contains: Ca 4.5 mmol/kg/day and P 2.7 mmol/kg/day.

Preterm milk	Ca, mmol (mg)/100 mL	P, mmol (mg)/100 mL
1 st week	0.7 (26)	0.4 (11)
2 nd week	0.6 (25)	0.5 (15)
Week 3/4	0.6 (25)	0.5 (14)
Week 10/12	0.7 (29)	0.4 (12)
Term milk		
1 st week	0.7 (26)	0.4 (12)
2 nd week	0.7 (28)	0.6 (17)
Week 3/4	0.7 (27)	0.5 (16)
Week 10/12	0.7 (26)	0.5 (16)

Elemental Ca, 1 mmol = 40 mg. Elemental Phosphorus, 1 mmol = 31 mg. Adapted from Gidrewicz and Fenton BMC Pediatrics 2014, 14:216. 15

Step 2: Calculate the gap in Ca and P intake/requirement: This will be the dose required.

Step 3: Prescribe 50% of the required dose of Ca and P in 2-3 divided doses alternatively but not together. (example: Ca 8 AM, 2 PM, 8 PM and P 6 AM, 12 MD, 6 PM).

Step 4: Once 50% dose is tolerated for 1 week, increase to 100% required dose.

ORAL preparation during NICU stay: Sodium dihydrogen phosphate Phebra IV is the preferred preparation for oral administration due to its low osmolality.

ORAL preparation at discharge or stable neonates: Phosphate-Sandoz tablets can be used.

American Academy of Pediatrics Committee on nutrition 2013 Guidelines on management for Enterally Fed Preterm Infants With Radiologic Evidence of Rickets: 1. Maximize nutrient intake. 2. If no further increases in these can be made, add elemental calcium and phosphorus as tolerated. Usually beginning at 20 mg/kg per day of elemental calcium and 10–20 mg/kg per day elemental phosphorus and increasing, as tolerated, usually to a maximum of 70–80 mg/kg per day of elemental calcium and 40–50 mg/kg per day elemental phosphorus. May consider targeting 25-OH-D concentration of >20 ng/mL (50 nmol/L).8 However, breast milk content of phosphorus is variable and harder to estimate the intakes accurately. A more pragmatic approach suggested by our consensus group: start with P 0.5-1.0 mmol/kg/day in divided doses and increase as tolerated to a maximum of P 3 mmol/kg/day.

Efficacy and safety

An ideal oral form of phosphate for use in preterm infants does not exist. Administering the intravenous preparations orally can be considered, because they are lower in osmolarity than are commercially available phosphorus-containing liquids. For example, potassium dihydrogen phosphate provides 31 mg

Newborn use only

	of elemental phosphorus per millimole. A dose of 10 to 20 mg/kg per day of elemental phosphorus is reasonable and will likely resolve hypophosphataemia in most preterm infants. ⁸	
	Oral phosphorus and feeds	
	It is recommended to separate oral doses from calcium and antacids containing agents such as	
	aluminium hydroxide, calcium or magnesium salts, as these may reduce the bioavailability of phosphate.	
	Oral phosphate preparation has high osmolality and administration with feeds may have theoretical	
	benefit of reducing the osmolality (consensus opinion).	
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Authors Contribution

Original author/s	Chris Wake, Srinivas Bolisetty
Expert review	
Current version author	Srinivas Bolisetty

Evidence Review	David Osborn
Nursing Review	Eszter Jozsa
Pharmacy Review	Jing Xiao, Cindy Chen
ANMF Group contributors	Nilkant Phad, Bhavesh Mehta, John Sinn, Michelle Jenkins, Thao Tran,
	Helen Huynh, Simarjit Kaur, Jessica Mehegan
Final content and editing review of the original	Ian Whyte
Electronic version	Cindy Chen, Ian Callander
Facilitator	Srinivas Bolisetty