

# METABOLIC BONE DISEASE (MBD) • 1/2

## RECOGNITION AND ASSESSMENT

### Definition

- Decreased mineralisation of bones due to deficient phosphate (PO<sub>4</sub>), calcium (Ca) or vitamin D in preterm babies
- Also known as osteopenia of prematurity

### Causes

- Inadequate postnatal intake or absorption to **match** intrauterine mineral accretion rate

### Risk factors

- <32 weeks' gestation **or** <1500 g birth weight
- Male gender
- **Delay in establishing enteral feeds**/enteral feeds with low mineral content/bioavailability [unfortified expressed breast milk (EBM), term formula]
- **PO<sub>4</sub> deficiency** (primary nutritional reason) **or** vitamin D deficiency
- Prolonged parenteral nutrition (PN) (**>2 weeks**)
- Chronic drug **use** that increases mineral excretion (diuretics, **steroids**, sodium bicarbonate)
- Lack of mechanical stimulation e.g. sedation/paralysis
- **Chronic lung disease**
- Cholestatic jaundice
- Short gut syndrome (malabsorption of vitamin D and Ca)

### Symptoms and signs

- ≤6 weeks – most babies are asymptomatic and normal on examination
- Usually presents aged 6–12 weeks
- Poor weight gain or faltering growth
- Respiratory difficulties: failure to **extubate** due to **increased** chest wall compliance
- Fractures with minor or no trauma; may manifest as pain on handling
- Craniotables (softening of skull bones)

### Later clinical consequences

- Marked dolicocephaly (long and narrow skull)
- Reduced linear growth

## DIAGNOSIS

### Serum biomarkers

- Low serum PO<sub>4</sub> (<1.8 mmol/L) with elevated **alkaline phosphatase (ALP)** (>900 IU/L) is 100% sensitive and 70% specific for diagnosing low bone mineral density
- Low serum PO<sub>4</sub> concentrations (<1.8 mmol/L) have 96% specificity but only 50% sensitivity. **Routine PO<sub>4</sub> supplementation in high risk babies could lead to secondary hyperparathyroidism, and thus worsen MBD**
- **Serum Ca levels may remain normal until late in the disease despite bone losses of Ca**
- **In suspected MBD with elevated ALP and low PO<sub>4</sub>, serum parathormone (PTH) measurement will help in establishing if there is underlying Ca or PO<sub>4</sub> deficiency to provide correct supplementation**
- **Ca deficiency causes increased PTH to maintain normocalcaemia**
- **in PO<sub>4</sub> deficiency there is no compensatory mechanism – PTH remains normal**

### Urinary biomarkers:

- Urinary excretion of Ca >1.2 mmol/L and PO<sub>4</sub> >0.4 mmol/L signifies slight surplus of supply and correlates with highest bone mineral accretion rate
- **phosphaturia** can occur due to aminoglycoside, indomethacin and **steroid** therapy
- **calciuria** can occur due to diuretics, **steroids** and theophylline
- Tubular reabsorption percent (TRP) of PO<sub>4</sub> is also a guide to adequacy of PO<sub>4</sub> supplementation. TRP of >95% indicates inadequate supplementation
- $TRP (\%TRP) = [1 - (\text{urine PO}_4 / \text{urine creatinine}) (\text{plasma creatinine} / \text{plasma PO}_4)] \times 100$

### Radiological

- **Low bone density on X-rays (rachitic changes, cortical thinning, periosteal elevation) or fractures of long bones or ribs**
- Dual-energy X-ray absorptiometry/qualitative ultrasound to assess bone mineral density

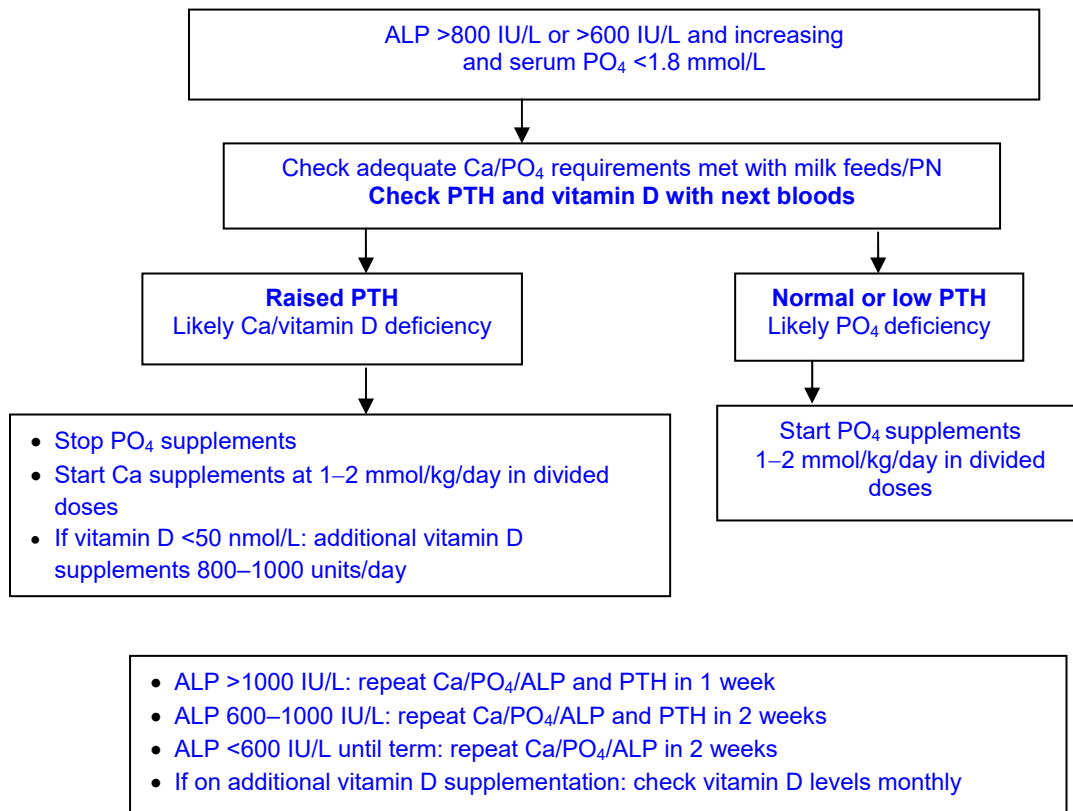
# METABOLIC BONE DISEASE (MBD) • 2/2

## PREVENTION

- Optimal nutritional intake
- early PN with optimised Ca and PO<sub>4</sub> content – see **Parenteral nutrition** guideline
- early enteral feeds
- optimal ratio of enteral Ca:PO<sub>4</sub> of 1.5:1 to 1.7:1 mmol-to-mmol basis to avoid secondary hyperparathyroidism
- adequate Ca (3–5.5 mmol/kg/day) and PO<sub>4</sub> (2.3–3.7 mmol/kg/day) intake by using fortified EBM or preterm formula. ≥140 mL/kg/day fortified EBM or preterm formula is needed to provide this
- daily intake of ≥400–700 units/kg/day vitamin D
- Ensure appropriate handling and position using deep boundaries to promote active bone loading

## INVESTIGATION AND TREATMENT:

For all high risk babies, measure serum Ca, PO<sub>4</sub> and ALP levels from third week of life and follow this guidance:



### NOTE:

- Do not give Ca and PO<sub>4</sub> at the same time as they may precipitate; give at alternate feeds
- Ca supplementation can cause intestinal obstruction and hypercalcinosis
- Regular monitoring of urinary Ca/urinary creatinine ratio necessary to detect hypercalciuria (urinary Ca/urinary creatinine) >0.6
- Consider other nutritional deficiencies e.g. zinc, in a baby with faltering growth with evidence of significant bone disease

## MONITORING AND FOLLOW-UP

- Stop additional vitamin D when vitamin D and PTH are normal
- Adjust Ca dose based on serum Ca levels and stop only once PTH levels normalise
- Adjust PO<sub>4</sub> dose based on serum PO<sub>4</sub> levels and ALP
- Continue treatment until biochemical indices are normal and radiographic evidence of healing, usually until term corrected gestation