Vitamin E Newborn use only

Alert This formulary covers oral vitamin E. Vitamin E 1 International Unit (hereafter referred to as "units") = 0.67 mg d-alpha-tocopherol.¹ Penta-Vite, a commonly used multi-vitamin supplement doesn't contain vitamin E. Indication Prevention and treatment of vitamin E deficiency. Neonatal cholestasis Action Fat soluble vitamin. It is an antioxidant protecting cell membranes from oxidative stress. Active isomer is α -tocopherol. Drug type Fat soluble vitamin. Trade name Micel-E oral liquid (Oral liquid SAS product may be available – water soluble liquid, Aqua-E containing 16 mg/mL (20 units/mL). Presentation Micel-E oral liquid: d-alpha-tocopherol 104.7 mg/mL (vitamin E 156 units/mL); 50 mL bottle. Dose Supplementation in preterm neonates* 8 units/kg daily (6-12 units/kg/day)² Neonatal cholestasis: Refer to vitamins in cholestasis formulary. *Preterm human milk + Human milk fortifier (HMF) at 170 mL/kg/day provides an average 8 units/kg/day. Dose adjustment Therapeutic hypothermia – No information. ECMO - No information. Renal impairment – No information. Hepatic impairment – No information. Maximum dose Doses exceeding 25 units/kg/day ORAL may pose more risk than benefit for preterm neonates.³ **Total cumulative** dose Route Oral Preparation No preparation is required. Administration Administer undiluted. Monitoring Serum vitamin E levels – Not routinely required. Target 1.0-2.0 mg/dL.^{4,5} Contraindications Hypersensitivity to vitamin E or any component Precautions Interacts with iron and other oxidants or any polyunsaturated fatty acids. Increases serum bilirubin. **Drug interactions** Iron - Lowers bioavailability of Vitamin E. Vitamin E may increase the effects of vitamin K antagonists and antiplatelet agents. **Adverse reactions** Sepsis. Intracranial haemorrhage (IV dosing). Necrotising enterocolitis. Compatibility Not applicable. Incompatibility Not applicable. Stability Micel E oral liquid: Store below 25°C (room temperature). Storage Micel-E: Potassium sorbate, citric acid anhydrous, glycerol, PEG-35 castor oil, ethanol, water. Excipients Special comments Evidence Efficacy Cochrane review by Brion et al 2003 assessed the effects of routine vitamin E supplementation on morbidity and mortality in preterm infants. Twenty-six randomized clinical trials with over 2000 preterm infants < 37 weeks or < 2500 g were analysed. In very low birth weight (VLBW) infants≤ 1500 g, vitamin E supplementation significantly reduced the risk of severe retinopathy and blindness but significantly increased the risk of sepsis. Subgroup analyses demonstrated (1) an association between intravenous, high-dose vitamin E supplementation and increased risk of sepsis and cerebral haemorrhage; (2) an association between non-intravenous vitamin E route and reduced risk of any or severe intraventricular haemorrhage and (3) an association between serum tocopherol levels greater than 3.5 mg/dl and increased risk of sepsis and reduced risk for severe retinopathy. Author's conclusions: Vitamin E supplementation in preterm infants reduced the risk of intracranial haemorrhage but increased the risk of sepsis. In VLBW infants, vitamin E increased the risk of sepsis, and reduced the risk of severe retinopathy

	and blindness among those examined. Evidence does not support the routine use of vitamin E supplementation by intravenous route at high doses or aiming at serum tocopherol levels greater than 35		
	mg/L (81 μmol/L). ⁶ (LOE I GOR A)		
	Safety		
	Routine vitamin E supplementation significantly reduced the risk of intraventricular haemorrhage but increased the risk of sepsis in preterm neonates. In VLBW infants (≤ 1500 g), vitamin E supplementation significantly increased the risk for sepsis and cerebral haemorrhage. (LOE I GOR A) A retrospective analysis has shown a significant association between pharmacologic oral doses of vitami E in VLBW infants and necrotizing enterocolitis ⁷ but this effect was not evident in meta-analysis. ⁶		
Practice points	Vitamin E content in preterm human milk: 0.64 units/dL (0.43 mg/dL)		
	Average human milk fortifier (HMF) at 80 kcal/100 mL provides additional 4-4.5 units/dL.		
	Preterm human milk + HMF at 170 mL/kg/day provides an average 8 units/kg/day. Recommended dietary allowances Colostrum and preterm human milk contains 2-3 times more alpha-tocopherol in mature milk. ^{2,8} Vitamin supplements for the preterm infant less than 1000 g birth weight are recommended to be 2.8 to 3.5 units/kg/day parenterally and 6 to 12 units/kg/day enterally. ^{2,3,9,10} (LOE III-3 GOR B) Recommended parenteral vitamin E for preterm neonates: 3 units/kg/day (2.8-3.5 units/kg/day). ^{2,10} SMOFlipid 20% contains 163 – 225 mg dl-alpha-tocopherol per 1000 mL. Vitalinid-N Infant contains 0.64 mg dl-alpha-tocopherol per 1 ml. ¹¹		
	The current Australasian consensus parenteral nutrition provides 2.8 units/kg/day at 150 mL/kg/day. ¹²		
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