Vitamin K₁ (Phytomenadione)

Newborn use only

Alert	Check ampoule carefully as an adult 10 mg ampoule (Konakion MM Adult) is also available.		
	USE ONLY Konakion MM Paediatric.		
	Vitamin K Deficiency Bleeding is also known as Haemorrhagic Disease of Newborn (HDN).		
Indication	Prophylaxis and treatment of vitamin K deficiency bleeding (VKDB)		
Action	Promotes the activation of blood coagulation Factors II, VII, IX and X in the liver		
Drug type	Fat soluble vitamin		
Trade name	Konakion MM Paediatric		
Presentation	2 mg/0.2 mL ampoule		
Dose	IM prophylaxis (Recommended route) ⁽¹⁾		
	 Birthweight ≥ 1500 g: 1 mg (0.1 mL of Konakion® MM) as a single dose at birth. 		
	Birthweight <1500 g: 0.5 mg (0.05 mL of Konakion® MM) as a single dose at birth.		
	Oral prophylaxis ⁽¹⁾		
	2 mg (0.2 mL of Konakion® MM) for 3 doses:		
	First dose: At birth		
	 Second dose: 3–5 days of age (at time of newborn screening) 		
	Third dose: During 4 th week (day 22-28 of life)		
	It is imperative that the third dose is given no later than 4 weeks after birth as the effect of		
	earlier doses decreases after this time		
	Repeat the oral dose if infant vomits within an hour of an oral dose or if diarrhoea occurs within		
	24 hours of administration		
	IV Prophylaxis ⁽⁵⁾		
	May be given in sick infants if unable to give IM or orally.		
	0.3 mg/kg (0.2-0.4 mg/kg) as a single dose as a slow bolus (maximum 1 mg/minute).		
	Dose may be repeated weekly.		
	Bose may be repeated weekly.		
	IV treatment of Vitamin K deficiency bleeding (VKDB)		
	1 mg IV as a slow bolus (maximum 1 mg/minute). Dose may be repeated in 4–6 hours if		
	required.		
	 Must be administered in the presence of a medical officer. 		
	May be given subcutaneously if venous access not available.		
Dose adjustment	No information		
Maximum dose			
Total cumulative			
dose			
Route	IM, Oral, IV, Subcutaneous		
Preparation	IM and Oral: Administer undiluted.		
ricparation	in and Oral. Administer difdirated.		
	IV: Draw up 0.2 mL (2 mg) of Konakion MM Paediatric and add 1.8 mL of glucose 5% or sodium chloride		
	0.9% to make a 1 mg/mL solution. (ANMF consensus)		
Administration	IM: Administer undiluted.		
	Oral: Injection solution can be administered orally via dispenser provided.		
	Repeated doses are advised if infant spits out or vomits within an hour of an oral dose or if diarrhoea		
	occurs within 24 hours of administration. Check with medical officer for advice.		
	IV: Slow bolus. Maximum rate 1 mg/minute.		
	Must be administered in the presence of a medical officer.		
	May be given subcutaneously if venous access not available.		
Monitoring	Prothrombin time when treating clotting abnormalities (a minimum of 2 to 4 hours is needed for		
	measurable improvement).		
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Contraindications	Oral prophylaxis is contraindicated in infants who are: preterm, unwell, on antibiotics, have cholestasis or
	have diarrhoea.
	Oral prophylaxis is contraindicated in infants of mothers who are on anticonvulsants including phenytoin,
	barbiturates and carbamazepine; rifampicin and the vitamin K antagonists including warfarin and phenindione.
Precautions	IV administration is associated with a possible risk of kernicterus in premature infants <2.5 kg.
Precautions	Efficacy of treatment is decreased in patients with liver disease.
Drug interactions	Co-administration of anticonvulsants can impair the action of vitamin K ₁ .
Adverse	Pain, swelling and erythema at IM injection site.
reactions	Severe hypersensitivity reactions, including death have been reported with rapid IV administration.
Compatibility	Fluids ^(8,9) : Glucose 5% (use immediately), glucose 10%, sodium chloride 0.9%, sodium chloride 0.45%.
Companionicy	Trains Torucose 378 (use infinediately), glacose 1078, sociali enionae 0.378, sociali enionae 0.4378.
	Y-site ⁽⁸⁾ : Amikacin, aminophylline, ascorbic acid, atracurium, atropine, azathioprine, aztreonam,
	benzylpenicillin, calcium chloride, calcium gluconate, cefazolin, cefotaxime, ceftazidime, ceftriaxone,
	cefuroxime, clindamycin, dexamethasone, dopamine, doxycycline, enalaprilat, adrenaline (epinephrine),
	epoietin alfa, erythromycin lactobionate, fentanyl, furosemide (frusemide), ganciclovir, gentamicin,
	heparin sodium, hydrocortisone, indomethacin, insulin regular, isoproterenol, labetalol, lidocaine,
	midazolam, morphine, naloxone, nitroglycerin, nitroprusside sodium, norepinephrine, oxacillin, penicillin
	G potassium, penicillin G sodium, phenobarbital (phenobarbitone), piperacillin, potassium chloride,
	propranolol, protamine, pyridoxine, ranitidine, sodium bicarbonate, streptokinase, succinylcholine,
	thiamine, ticarcillin, ticarcillin-clavulanate, tobramycin, tolazoline, urokinase, vancomycin, vasopressin,
	verapamil.
	Variable compatibility ⁽⁸⁾ : Amphotericin B conventional colloidal, ampicillin, dobutamine, hydralazine,
	methylprednisolone.
Incompatibility	Fluids: Fat emulsion (intravenous).
	Y-site ⁽⁸⁾ : Diazepam, diazoxide, magnesium sulfate, phenytoin, sulfamethoxazole-trimethoprim.
Stability	Use immediately.
Storage	Store below 25°C. Protect from light.
Excipients	Glycocholic acid, lecithin, sodium hydroxide, hydrochloric acid
Special	The risk of childhood cancer is not increased by IM administration of vitamin K ₁ .
comments Evidence	Destroyand
Evidence	Background
	All nowhern infants have a relative vitamin K deficiency at high Vitamin K, crosses the placenta nearly
	All newborn infants have a relative vitamin K deficiency at birth. Vitamin K ₁ crosses the placenta poorly
	resulting in low foetal plasma concentrations of the vitamin, with a 30:1 maternal-infant gradient.
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IV group when compared to 0.2 or 0.5 mg IM groups on day 5. By day 25, vitamin K_1 levels had declined in all the groups, but infants who received 0.5 mg IM had higher levels of vitamin K_1 than either the 0.2 mg IV group or the 0.2 mg IM group. Since there is no available evidence that vitamin K is harmful or ineffective and since vitamin K is an inexpensive drug, authors concluded to follow the recommendations of expert bodies and give vitamin K to preterm infants. (3)

Treatment of VKDB: Any infant suspected of VKDB should receive immediate intravenous vitamin K replacement. It is standard practice to administer a dose of 1 mg which will usually result in correction within a few hours. (LOE IV; GOR C) Intravenous vitamin K can be associated with anaphylactoid reactions and should be administered by slow intravenous injection; if venous access cannot be established it can be given subcutaneously, the intramuscular route being avoided in the presence of a coagulopathy. (4)

Pharmacokinetics

In healthy, fully breast-fed, newborn babies, significantly higher plasma vitamin K_1 concentrations were reported several weeks after IM as compared to oral vitamin K_1 . Half-life of oral and intramuscular vitamin K_1 were considerably longer in newborn infants (median 76 hours; range 26 to 193 hours)^(5, 6) compared to adults (6 hours; range 2–26 hours)⁽⁷⁾. Re-dosing of oral vitamin K_1 is recommended by 1 month in breast fed infants.⁽⁶⁾ (LOE II GOR B)

In preterm infants and sick infants unable to receive intramuscular vitamin K_1 , 0.3 mg/kg intravenously resulted in similar serum concentrations as oral administration of 3 mg vitamin K_1 and intramuscular administration of 1.5 mg vitamin K_1 supports recommendation for intravenous 0.4 mg/kg phytomenadione - vitamin K_1 - Konakion MM Paediatric in infants unable to receive oral or intramuscular vitamin K_1 .⁽⁵⁾ (LOE IV, GOR B).

Practice points

Australian NHMRC Guidelines 2010 position statement(1):

- All newborn infants should receive vitamin K prophylaxis.
- Healthy newborn infants should receive vitamin K₁ either:
 - o By intramuscular injection of 1 mg (0.1 mL) of Konakion® MM Paediatric at birth. This is the preferred route for reliability of administration and level of compliance **OR**
 - Three 2 mg (0.2 mL) oral doses of Konakion® MM Paediatric, given at birth, at the time of newborn screening (usually at 3-5 days of age) and in the fourth week.
- Newborns who are too unwell and are unable to take oral vitamin K₁ (or whose mothers have taken medications that interfere with vitamin K metabolism) should be given 1 mg of Konakion® MM Paediatric by intramuscular injection at birth. A smaller intramuscular dose of 0.5 mg (0.05 mL) should be given to infants with a birth weight of less than 1.5 kg.

References

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Authors Contribution

Original author/s	Srinivas Bolisetty, Nilkant Phad
Evidence Review	Srinivas Bolisetty
Expert review	
Nursing Review	Eszter Jozsa, Kirsty Minter
Pharmacy Review	Cecilia Peng
ANMF Group contributors	Nilkant Phad, Bhavesh Mehta, John Sinn, Rebecca Barzegar, Mohammad Irfan Azeem, Kate
	Dehlsen, Michelle Jenkins, Helen Huynh, Stephanie Halena
Final editing	Thao Tran
Electronic version	Cindy Chen, Ian Callander
Facilitator	Srinivas Bolisetty