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	Enoxaparin is one type	e of low molecular weight hepari	ı (LMWH).				
	Commonly known as Clexane.						
	High risk medication. An overdose can be fatal.						
	Treatment must be dis	Treatment must be discussed with the Haematologist on-call before commencement.					
	LMWH is not a suitable choice of anticoagulant in patients with significant bleeding risk (unfractionated						
	heparin (UFH) is preferred), who are clinically unstable or about to have invasive procedures. This is du						
	longer half-life than UFH and only partial reversal with protamine. Monitoring is performed with anti-factor Xa levels. The APTT is not useful in monitoring LMWH therapy. Please check with your local pathology department on what time of the day/night anti-factor Xa sample processing is performed.						
Indication	Prophylaxis of thromb						
	,						
	(Note: Enoxaparin/heparin does not treat the clot that has already occurred but rather its role is to						
	prevent clot extension, i.e. secondary prophylaxis)						
Action	It binds to and potentiates anti-thrombin III activity leading to irreversible inactivation of factor Xa, and						
Action	a lesser degree inactivation of factor IIa; in turn, inhibiting thrombin and fibrinogen generation.						
Drug tuno		anticoagulant; LMWH	ing thrombin and histinogen generation.				
Drug type							
Trade name	Clevane, Clexane Forte		st automotic cofoty lock auctors and street				
Presentation	Clexane (enoxaparin sodium) prefilled syringes, with/out automatic safety lock system, solution for						
	injection*:						
	20 mg/0.2mL						
	40 mg/0.4mL						
	60 mg/0.6mL						
	80 mg/0.8mL						
	100 mg/1mL						
	*containing 10 000 and	ti-Xa unit/mL					
	Clexane Forte, with/out automatic safety lock system, solution for injection [∆] :						
	120mg/0.8mL						
	150mg/1mL						
	△ containing 15 000 anti-Xa unit/mL						
	[△] containing 15 000 an	ti-Xa unit/mL					
	[∆] containing 15 000 an	ti-Xa unit/mL					
			aseptically prepared by local pharmacy.				
Dose		for patient specific doses can be	aseptically prepared by local pharmacy.				
Dose	Enoxaparin injections f	for patient specific doses can be	aseptically prepared by local pharmacy.				
Dose	Enoxaparin injections f	for patient specific doses can be	aseptically prepared by local pharmacy. ≥2 months				
Dose	Enoxaparin injections f Subcutaneous (SC) inje	for patient specific doses can be ection: ¹ 2 months of age	≥2 months				
Dose	Enoxaparin injections f	for patient specific doses can be ection: ¹					
Dose	Enoxaparin injections f Subcutaneous (SC) inje	for patient specific doses can be ection: <pre> <2 months of age 0.75 mg/kg/dose 12 hourly </pre>	≥2 months 0.5 mg/kg/dose 12 hourly				
Dose	Enoxaparin injections for Subcutaneous (SC) injections for Prophylactic dose	for patient specific doses can be ection: <pre> <2 months of age 0.75 mg/kg/dose 12 hourly <2 months of age</pre>	≥2 months 0.5 mg/kg/dose 12 hourly ≥2 months				
Dose	Enoxaparin injections f Subcutaneous (SC) inje	for patient specific doses can be ection: <pre> <2 months of age 0.75 mg/kg/dose 12 hourly </pre>	≥2 months 0.5 mg/kg/dose 12 hourly				
Dose	Enoxaparin injections for Subcutaneous (SC) injections for Prophylactic dose Treatment dose	for patient specific doses can be ection: ¹ <2 months of age 0.75 mg/kg/dose 12 hourly <2 months of age 1.5 mg/kg/dose 12 hourly	≥2 months 0.5 mg/kg/dose 12 hourly ≥2 months 1 mg/kg/dose 12 hourly	3 to			
Dose	Enoxaparin injections of Subcutaneous (SC) injections of Subcu	for patient specific doses can be ection: <pre> <2 months of age 0.75 mg/kg/dose 12 hourly <2 months of age 1.5 mg/kg/dose 12 hourly tion is as per anti-Xa levels. The features </pre>	≥2 months 0.5 mg/kg/dose 12 hourly ≥2 months 1 mg/kg/dose 12 hourly rst anti-Xa measurement is usually done after	3 to			
Dose	Enoxaparin injections of Subcutaneous (SC) injections of Subcu	for patient specific doses can be ection: <2 months of age 0.75 mg/kg/dose 12 hourly <2 months of age 1.5 mg/kg/dose 12 hourly tion is as per anti-Xa levels. The fa hours after the commencement	≥2 months 0.5 mg/kg/dose 12 hourly ≥2 months 1 mg/kg/dose 12 hourly rst anti-Xa measurement is usually done after	3 to			
Dose	Enoxaparin injections of Subcutaneous (SC) injections of Subcu	for patient specific doses can be ection: <2 months of age 0.75 mg/kg/dose 12 hourly <2 months of age 1.5 mg/kg/dose 12 hourly tion is as per anti-Xa levels. The factors after the commencement age: 0.5 to 1.0 units/mL to be me	≥2 months 0.5 mg/kg/dose 12 hourly ≥2 months 1 mg/kg/dose 12 hourly rst anti-Xa measurement is usually done after asured 4 hours (3-5 hours) after the last	3 to			
Dose	Enoxaparin injections of Subcutaneous (SC) injections of Subcu	for patient specific doses can be ection: <2 months of age 0.75 mg/kg/dose 12 hourly <2 months of age 1.5 mg/kg/dose 12 hourly tion is as per anti-Xa levels. The fa hours after the commencement	≥2 months 0.5 mg/kg/dose 12 hourly ≥2 months 1 mg/kg/dose 12 hourly rst anti-Xa measurement is usually done after asured 4 hours (3-5 hours) after the last	3 to			
Dose	Enoxaparin injections of Subcutaneous (SC) injections of Subcutaneous (SC) injections of Subcutaneous (SC) injections of Subcutaneous injections of Subcutan	ction: <2 months of age 0.75 mg/kg/dose 12 hourly <2 months of age 1.5 mg/kg/dose 12 hourly tion is as per anti-Xa levels. The factor of the commencement ange: 0.5 to 1.0 units/mL to be men. Refer to dose adjustment below	≥2 months 0.5 mg/kg/dose 12 hourly ≥2 months 1 mg/kg/dose 12 hourly rst anti-Xa measurement is usually done after assured 4 hours (3-5 hours) after the last ow: ⁸	3 to			
Dose	Enoxaparin injections of Subcutaneous (SC) injections of Subcutaneous (SC) injections of Subcutaneous of Subsequent dose Treatment dose Subsequent dose titrated doses, i.e. around 48 Target peak anti-Xa rar subcutaneous injection Anti-factor Xa	for patient specific doses can be ection: <2 months of age 0.75 mg/kg/dose 12 hourly <2 months of age 1.5 mg/kg/dose 12 hourly tion is as per anti-Xa levels. The factors after the commencement age: 0.5 to 1.0 units/mL to be me	≥2 months 0.5 mg/kg/dose 12 hourly ≥2 months 1 mg/kg/dose 12 hourly rst anti-Xa measurement is usually done after asured 4 hours (3-5 hours) after the last	3 to			
Dose	Enoxaparin injections of Subcutaneous (SC) injections of Subcutaneous (SC) injections of Subcutaneous of Subsequent dose Treatment dose Subsequent dose titrated doses, i.e. around 48 Target peak anti-Xa rais subcutaneous injection Anti-factor Xa concentration	ction: <2 months of age 0.75 mg/kg/dose 12 hourly <2 months of age 1.5 mg/kg/dose 12 hourly tion is as per anti-Xa levels. The factor of the commencement ange: 0.5 to 1.0 units/mL to be men. Refer to dose adjustment below	≥2 months 0.5 mg/kg/dose 12 hourly ≥2 months 1 mg/kg/dose 12 hourly rst anti-Xa measurement is usually done after assured 4 hours (3-5 hours) after the last ow: ⁸	3 to			
Dose	Enoxaparin injections of Subcutaneous (SC) injections of Subcutaneous (SC) injections of Subcutaneous of Subsequent dose Treatment dose Subsequent dose titrated doses, i.e. around 48 Target peak anti-Xa rais subcutaneous injection Anti-factor Xa concentration unit/mL	for patient specific doses can be ection: <2 months of age 0.75 mg/kg/dose 12 hourly <2 months of age 1.5 mg/kg/dose 12 hourly tion is as per anti-Xa levels. The factors after the commencement ange: 0.5 to 1.0 units/mL to be men.¹ Refer to dose adjustment below	≥2 months 0.5 mg/kg/dose 12 hourly ≥2 months 1 mg/kg/dose 12 hourly rst anti-Xa measurement is usually done after assured 4 hours (3-5 hours) after the last ow: Next anti-factor Xa measurement	3 to			
Dose	Enoxaparin injections of Subcutaneous (SC) injections of Subcutaneous (SC) injections of Subsequent dose Treatment dose Subsequent dose titrated doses, i.e. around 48 Target peak anti-Xa rais subcutaneous injection Anti-factor Xa concentration unit/mL <0.35	for patient specific doses can be ection: <2 months of age 0.75 mg/kg/dose 12 hourly <2 months of age 1.5 mg/kg/dose 12 hourly tion is as per anti-Xa levels. The fa hours after the commencement age: 0.5 to 1.0 units/mL to be men.¹ Refer to dose adjustment below Dose adjustment increase next dose by 25%	≥2 months 0.5 mg/kg/dose 12 hourly ≥2 months 1 mg/kg/dose 12 hourly rst anti-Xa measurement is usually done after assured 4 hours (3-5 hours) after the last ow: Next anti-factor Xa measurement 4 hr following dose adjustment	3 to			
Dose	Enoxaparin injections of Subcutaneous (SC) injections of Subcutaneous (SC) injections of Subsequent dose Treatment dose Subsequent dose titrated doses, i.e. around 48 Target peak anti-Xa rare subcutaneous injection Anti-factor Xa concentration unit/mL <0.35 0.35 - 0.49	ction: Camonths of age	≥2 months 0.5 mg/kg/dose 12 hourly ≥2 months 1 mg/kg/dose 12 hourly rst anti-Xa measurement is usually done after assured 4 hours (3-5 hours) after the last ow:8 Next anti-factor Xa measurement 4 hr following dose adjustment 4 hr following dose adjustment	3 to			
Dose	Enoxaparin injections of Subcutaneous (SC) injections of Subcutaneous (SC) injections of Subsequent dose Treatment dose Subsequent dose titrated doses, i.e. around 48 Target peak anti-Xa rais subcutaneous injection Anti-factor Xa concentration unit/mL <0.35	for patient specific doses can be ection: <2 months of age 0.75 mg/kg/dose 12 hourly <2 months of age 1.5 mg/kg/dose 12 hourly tion is as per anti-Xa levels. The fa hours after the commencement age: 0.5 to 1.0 units/mL to be men.¹ Refer to dose adjustment below Dose adjustment increase next dose by 25%	≥2 months 0.5 mg/kg/dose 12 hourly ≥2 months 1 mg/kg/dose 12 hourly rst anti-Xa measurement is usually done after assured 4 hours (3-5 hours) after the last ow: Next anti-factor Xa measurement 4 hr following dose adjustment 4 hr following dose adjustment Weekly 4 hr following a dose	3 to			
Dose	Enoxaparin injections of Subcutaneous (SC) injections of Subcutaneous (SC) injections of Subsequent dose Treatment dose Subsequent dose titrated doses, i.e. around 48 Target peak anti-Xa rare subcutaneous injection Anti-factor Xa concentration unit/mL <0.35 0.35 - 0.49	ction: Camonths of age	≥2 months 0.5 mg/kg/dose 12 hourly ≥2 months 1 mg/kg/dose 12 hourly rst anti-Xa measurement is usually done after assured 4 hours (3-5 hours) after the last ow: Next anti-factor Xa measurement 4 hr following dose adjustment 4 hr following dose adjustment Weekly 4 hr following a dose If change in renal function, addition	3 to			
Dose	Enoxaparin injections of Subcutaneous (SC) injections of Subcutaneous (SC) injections of Subsequent dose Treatment dose Subsequent dose titrated doses, i.e. around 48 Target peak anti-Xa rare subcutaneous injection Anti-factor Xa concentration unit/mL <0.35 0.35 - 0.49	ction: Camonths of age	≥2 months 0.5 mg/kg/dose 12 hourly ≥2 months 1 mg/kg/dose 12 hourly rst anti-Xa measurement is usually done after assured 4 hours (3-5 hours) after the last ow: Next anti-factor Xa measurement 4 hr following dose adjustment 4 hr following dose adjustment Weekly 4 hr following a dose	3 to			

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		1.1-1.5		decrease	next dose by 20	%	Before n		d 4 h following	
	1.6 to 2.0 hold dose until anti-factor Xa level <1 then decrease next dose by 30% 4 hr following dose adjustments				ndjustment					
		>2.0			until anti-factor decrease next d				Xa level <0.5, einstitution of	
Dose adjustment	Re im	nal impairmen pairment. Disc	t – Moni uss with	tor anti-Xa haematol	rin is not the pro a factor closely. I ogist. nent is not establ	Oose adjust	_		vere renal	
Maximum dose		-		-						
Total cumulative dose										
Route	Su	bcutaneous inj	ection.							
Preparation	Enoxaparin injections for patient specific administration can be aseptically prepared by local pharmac follows: Draw 0.8 mL of sodium chloride 0.9% into a 2 mL syringe. Inject the contents of enoxaparin 20 mg/0.2 pre-filled syringe into the sodium chloride syringe to make a final volume of 1 mL. The resulting solut contains 20 mg/mL.				.2 mL					
	D	ose	1.5 mg	3	2 mg	3 mg	4	4 mg	5 mg	
	V	olume	0.075	mL	0.1 mL	0.15 mL	_ (0.2 mL	0.25 mL	
Administration	_	scard remaining			remove the air b	ubble in th	ne prefilled	svringe Ro	tate the site of	
	subcutaneous injection. Enoxaparin may also be administered via an Insuflon catheter placed into the subcutaneous tissue. However, Insuflon catheters are not recommended for infants less than 3kg. When administering enoxaparin via an Insuflon catheter, the air bubble in the syringe should be removed. Do not rub the injection site after administration. Note: Injection in low birth weight infants with little subcutaneous fat may enter intramuscular rather than subcutaneous which can impact anti-Xa level due to different absorption rate and pharmacokinetics. Significant tissue oedema at injection sites may also impact absorption.									
Monitoring	Anti-factor Xa levels Platelet count every 2-3 days Potassium levels Renal function									
Contraindications	Hypersensitivity to enoxaparin, heparin or other low molecular weight heparins Active uncontrollable bleeding Severe thrombocytopenia (MIMS) Haemorrhagic stroke Acute bacterial endocarditis (MIMS) History of heparin-induced thrombocytopenia (HIT) within the past 100 days (MIMS)									
Precautions	Risk of haemorrhage – example, acquired or congenital bleeding disorders Concomitant medical conditions: Hepatic insufficiency, uncontrolled hypertension, a history of gastrointestinal ulceration, recent neuro- or ophthalmologic surgery and haemorrhage. Heparin-induced thrombocytopenia (HIT) Spinal anaesthesia									
Drug interactions	Dri An	ugs affecting ha ticoagulants, tl stemic glucocol	aemosta hrombol rticoids.	ytics, non- If the com	be discontinued steroidal anti-in bination is indica emostatic factor	flammator ated, enox	y agents, a aparin shou	spirin, antip uld be used	latelet agents o	r

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	Drugs that increase serum notacsium levels may be administered consurrently with enevanerin sedium
	Drugs that increase serum potassium levels may be administered concurrently with enoxaparin sodium under careful clinical and laboratory monitoring.
Adverse reactions	Elevated liver enzymes, anaemia, diarrhoea, peripheral oedema, fever, allergic reaction, urticarial,
	bruising/ pain at injection site, bleeding, hyperkalaemia
	Rare: Thrombocytopenia, hyperkalaemia, cholestasis, bullous dermatitis, osteoporosis, allergic reaction
Compatibility	Glucose 5%, sodium chloride 0.9%
Incompatibility	No information available
Stability	Discard any unused contents of syringes.
,	
	Aseptically prepared product by local pharmacy is stored refrigerated at 2-8°C with an expiry date of 7
	days.
Storage	Store below 25°C. Do not freeze.
	Aseptically prepared product by local pharmacy is stored refrigerated at 2-8°C.
Excipients	Water for injections
Special comments	Protamine may be used to reverse anticoagulant effect of enoxaparin but the reversal is partial.
Evidence	Efficacy
	A review of published reports between 1980 and 2007 comprising of 240 neonates from 13 studies
	showed that the mean enoxaparin dose that resulted in therapeutic plasma anti-factor Xa levels of 0.5-1.0
	units/mL varied between 1.48 and 2.27 mg/kg subcut every 12 hours for all infants. The mean length of
	therapy for neonatal thrombosis fluctuated from 12 days to 3 months. Higher doses in preterm neonates
	have been suggested to maintain therapeutic anti-Xa levels. ^{6,7} However, clinical trials have not been
	performed to confirm the safety and efficacy of a higher-dose approach.
	American College of Chest Physicians Evidence-Based Clinical Practice Guidelines 2012 recommend a
	prophylactic dose of 0.75 mg/kg/dose subcut every 12 hours and 0.5 mg/kg/dose subcut every 12 hours
	for infants <2 months and ≥2 months respectively. For treatment, doses of 1.5 mg/kg/dose subcut every
	12 hours and 1 mg/kg/dose subcut every 12 hours are recommended for <2 months and ≥2 months
	respectively. ¹
	Safety
	A review of enoxaparin in neonates reported that minor side effects were common; major bleeding was
	recorded in 5% neonates. 5 Whether premature infants are at increased risk is unclear. No major bleeds
	were reported in a series of 10 premature neonates. ⁷
	There are no data addressing the frequency of osteoporosis, HIT, or other hypersensitivity reactions in
	children exposed to LMWH. ¹
	Enoxaparin overdose: The optimal management of LMWH overdose in the paediatric population has not
	been established. In common practice, enoxaparin overdose can be reversed by administration of
	protamine using a 1: 1 ratio to LMWH (example: 1 mg enoxaparin = 1 mg protamine). The dose of
	protamine can be given as a single dose or divided into 2-3 doses at 4 hour intervals aiming to return anti-
	Xa levels to therapeutic range. ⁴
	Dhawana kinatica
	Pharmacokinetics Enoxaparin sodium is obtained by alkaline depolymerisation of heparin benzyl ester derived from porcine
	intestinal mucosa. ³ Body weight is the most predictive covariate for clearance and central volume of
	distribution. After injection of Clexane by the subcutaneous route, the product is rapidly and completely
	absorbed. The absolute bioavailability is over 90%. It is primarily metabolised in the liver. Small amounts
	are eliminated by kidneys in an intact or slightly degraded form. ³ Elimination is not significantly modified in
	mild to moderate renal insufficiency. ³
Practice points	The dose regimen and monitoring recommendations in this formulary is based the American College of
Tarante ponito	Chest Physicians Evidence-Based Clinical Practice Guidelines 2012. These guidelines were based on studies
	for anticoagulation in neonates that have been part of larger studies reporting on children in general and
	report use of twice-daily enoxaparin targeted to an anti-Xa range (measured 4-6 h after dose) of 0.5 to 1.0
	units/mL.1 (LOE II, GOR C)
	Recommendations for dose adjustment are based on cohort studies in children.8 (LOE IV, GOR C)
References	Monagle P, Chan AK, Goldenberg NA, Ichord RN, Journeycake JM, Nowak-Göttl U, Vesely SK.
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