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# Low molecular weight heparin (LMWH) within North Central London (NCL)



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#### **1.0 INTRODUCTION**

Low molecular weight heparins (LMWH) are widely used for the prevention and treatment of venous thromboses as well as a number of unlicensed indications. Some, patients may be discharged from secondary care whilst on prophylaxis or treatment dose LMWH.

This guidance document aims to:

- Outline the types of patient that are considered appropriate for primary care management within NCL
- Outline those patients where management should remain within secondary care
- Outline the monitoring required for LMWHs
- Highlight essential information that should be transferred between sectors, in line with NPSA requirements 2010<sup>1</sup> (e.g. indication, dose, weight, renal function, monitoring, duration of treatment)

As there are a number of LMWHs in use across the sector, this document is not intended to be a treatment dose prescribing guide; the relevant local Trust policies and standard reference sources (e.g. BNF) should be consulted accordingly. This document is applicable to all NHS Trusts, CCGs and GPs within the NCL Joint Formulary Committee umbrella. Organisations wishing to commission services from NCL providers will be expected to abide with the recommendations within this document.

#### **Abbreviations:**

- Warfarin: where stated, this includes the other vitamin K antagonists, acenocoumarol and phenindione
- VKA: vitamin K antagonists

- HIT: Heparin induced thrombocytopenia
- AC: Anticoagulation

## 2.0 THROMBOPROPHYLACTIC LMWH – prescribing and monitoring

**2.1 Thromboprophylaxis - secondary care** supply and (where appropriate) monitor the entire defined course of LMWH (up to 6wks or longer as appropriate<sup>2</sup>). *NB: list is not exhaustive / exclusive* 

	appropriate ). NB: list is not exhaustive / exclusive				
Speciality	Indication	Usual duration (as per NICE <sup>2</sup> or local Trust			
		guidelines). Assumes pt is back to baseline mobility*			
Orthopaedics	Hip fracture surgery	28-35 days post-op			
Orthopaedics	Hip replacement	28-35 days post op			
Orthopaedics	Knee replacement	10-14 days post-op			
Orthopaedics	Major limb fracture / pelvic fracture	At least 28-35 days			
Orthopaedics	Lower limb plaster cast	For duration of plaster cast			
Orthopaedics	Other high risk major orthopaedic surgery	Up to 6 weeks or as per orthopaedic surgeon (NB:			
		primary care may be asked to prescribe if			
		thromboprophylaxis required beyond 6 weeks)			
All surgical	Major cancer surgery in the abdomen or pelvis	28 days post-op			
specialities					
All surgical	Higher VTE risk post-op (excludes patients usually As directed by the surgeon / haemat				
specialities	on warfarin)	Usually up to 28 days post-op			
Bariatric surgery	Laparoscopic sleeve gastrectomy / gastric bypass /	/ At least 5-7 days post-op			
	gastric band				
Haematology /	Higher VTE risk postnatally	Up to 6 weeks postnatally as per haematology			
obstetrics	(see section 2.2 re antenatal LMWH)	consultant			
Obstetrics	Intermediate VTE risk postnatally	At least 7 days post delivery <sup>3</sup> or as advised by			
		haematology consultant. NB prophylactic LMWH			
		doses used in pregnancy differ from the usual			
		licensed doses – see section 2.3			
Haematology	Higher VTE risk patient treated with VTE inducing	For duration of lenalidomide/thalidomide and as			
	drug such as lenalidomide/thalidomide	advised by haematology consultant			
Oncology	High risk of VTE and receiving chemotherapy (e.g.	As advised by oncologist / haematologist. Note:			
	previous PICC associated VTE)	VTE risk continues whilst ongoing active cancer.			

\*Duration may need to be extended if not fully mobile

**2.2 Thromboprophylaxis - primary care** primary care to supply/monitor LMWH for extended duration (> 6weeks), or for long-term thromboprophylaxis. Initial supply (e.g. 4wks) from secondary care. *NB: list is not exhaustive/ exclusive* 

Speciality	Indication	Duration as per local specialist (or local Trust guidelines).
Gastroenterology / Nutrition	History of PICC associated DVT and with a need for ongoing PICC line (e.g. for parenteral nutrition)	Whilst PICC line remains in situ
Orthopaedics	Complex orthopaedic surgery	As per orthopaedic surgeon on a case-by-case basis
Oncology	High risk of VTE and not receiving	As advised by oncologist / haematologist. NB: VTE
	chemotherapy	risk continues whilst ongoing active cancer
Haematology/	Obstetric patient requiring	As advised by haemostasis/ haematology consultant
obstetrics	thromboprophylaxis during pregnancy (e.g.	(UCLH pts will be managed in secondary care). See
	for VTE / cardiac indications)	comment in section 2.3 re dose.
Haematology	Higher risk patients as per haemostasis	As advised by haemostasis / haematology consultant
	/haematology consultant	on a case-by-case basis
Haematology	Very high risk VTE pt requiring pre-flight VTE	Single doses pre-flight as advised by haemostasis /
	thromboprophylaxis – Haem advice only	haematology consultant

2.3 Dose - thromboprophylaxis	If > 100kg use higher doses as per secondary care guidelines (not included)		
LMWH	Prophylactic doses as per BNF		
Dalteparin	2500 units or 5000 units sc OD		
Enoxaparin	20mg or 40mg sc OD		
Tinzparin	3500units or 4500 units sc OD		
Dose (thromboprophylaxis) in PREGNANCY			
<ul> <li>LMWH thromboprophylactic doses used in pregnancy differ to the standard doses listed in the BNF - the haemostasis/haematology consultant will advise accordingly.</li> <li>Patients &gt; 90kg may require a higher thromboprophylactic dose as per RCOG<sup>3</sup>. local guidelines and as advised by</li> </ul>			

 Patients > 90kg may require a higher thromboprophylactic dose as per RCOG<sup>3</sup>, local guidelines and as advised by haemostasis/haematology consultant. Any potential dose changes based on weight, will be made clear in correspondence from the consultant

# 3.0 THERAPEUTIC DOSE LMWH - prescribing and monitoring

3.1 Therapeutic - secondary care

For the indications below, secondary care should

prescribe / monitor therapeutic dose LMWH and make appropriate plans for follow up. *NB: list is not* 

exhaustive / exclusive. In some cases, LMWH may be split into two divided doses as advised by haematology.						
Speciality	Indication	Usual duration				
	(or as per local Trust guidelines)					
AC clinic (hospital	C clinic (hospital Warfarin patient requiring a 'bridging' plan Peri-procedurally and unti					
based)	around surgery/procedure; patient's own AC	(NB: some patients may receive thrombo-				
	clinic to organise and manage LMWH	prophylactic rather than treatment dose)				
		in charge of the patient's clinical care, to use local				
	Trust bridging guidelines in liaison with both the	patient's own AC clinic and pre-assessment clinical				
	team, to formulate a pre-operative bridging plan.	If there is a date change to the procedure, team to				
	also communio	cate accordingly				
AC clinic	Patient in process of being newly warfarinised	Until warfarinised as per local guidelines				
(hospital based)	and LMWH being used as an interim					
Haematology	Pregnant women on therapeutic dose or higher	Antenatally and postnatally, as per haemostasis				
	intensity LMWH	/ haematology consultant				
Oncology	Patient requiring therapeutic dose LMWH and	Depends on indication for anticoagulation.				
	undergoing chemotherapy.	LMWH usually continues for duration of				
		chemotherapy and in the case of new VTE for at				
least 6 months post event (whichever i						
	longer) and then oncology to review; if ongo					
	active cancer, then AC is usually continued					
	beyond 6 months as ongoing VTE risk					
Any speciality	Patients with history of recent bleed or bleeding	Follow up as per local trust arrangement;				
	disorder (e.g. thrombocytopenia) on	haemostasis/haematology consultant to define				
	anticoagulation	when patient can be transferred to Primary care.				
Haematology	High risk VTE patient treated with VTE inducing	For duration of lenalidomide/ thalidomide and				
	drug such as lenalidomide or thalidomide as advised by haematology consultant					
3.2 Therapeutic - initial management by 2 <sup>0</sup> care, before considering transfer to 1 <sup>0</sup> care						
• The following	patient groups will initially be managed by the team	in charge of the overall care of the patient with				
specialist haemostasis/haematology advice (preferably before discharge). NB: list is not exhaustive / exclusive.						
• If it is the intention that the GP should continue the prescribing and monitoring of LMWH, then the form in appendix						
1 (or a suitably detailed letter or discharge summary outlining similar information) should be completed and faxed to						

1 (or a suitably detailed letter or discharge summary outlining similar information) should be completed and faxed to the GP. This should be scanned onto the hospital electronic patient records, as proof of communication

<sup>•</sup> Clear instructions must be provided to the GP regarding when an adjustment to the LMWH dose would be required (e.g. weight change, change in renal function) – see section 4.0)

•	In some	e cases, L	MWH m	ay be split into two	divided doses as a	dvised by haematology

In some cases, Linver may be spirit into two divided doses as advised by naematology				
Speciality	Patient groups		Comments	
Any speciality	Renal impairment*:		Dose reductions required for renal impairment as	
	CrCL < 30mL/min: enoxapari	n or dalteparin;	advised by haemostasis / haematology consultant	
	CrCL < 20mL/min: tinzaparin			
Any speciality	Significant hepatic impairmer	nt		
Any speciality	High body weight e.g.	Or low body weight		
	> 110 kg for dalteparin,	e.g. < 40kg		
	> 105 kg for tinzaparin,			
	> 100kg for enoxaparin			
Haematology	Ongoing LMWH monitoring a	nd patient review by	These patients may have e.g. 3-6 monthly /	
	a haemostasis/haematology of	consultant, but	annual follow-up with haematology and it is	
			usually inappropriate to prescribe for these	
			durations. 1 <sup>0</sup> care may be asked to prescribe.	
Any speciality	lity Patient requiring higher intensity anticoagulation		Higher intensity for a period of time (patient	
(e.g. higher than standard BNF VTE dose) as		specific) as defined by haemostasis / haematology		
	advised by haemostasis/haematology consultant		consultant (follow up as per local trust	
(example: dalteparin 120 units/kg sc BD)		arrangement)		
Oncology	ncology Whilst on chemotherapy, LMWH should be		Duration depends on indication and cancer status.	
	managed by secondary care.	When chemotherapy	First VTE event in setting of cancer: LMWH usually	
has stopped, transfer to primary care could be		for at least 6 months then review by oncology; if		
	considered.		ongoing active cancer, then AC is usually	
			continued beyond 6months although choice/dose	
			can be reviewed. Patient may require longterm	
		LMWH		

\*could use eGFR in place of CrCL for <u>most adult patients of average build and height as per BNF</u>. **But**, for extremes of body weight, use CrCL (see comment in section 4.3 and appendix 2)

**3.3 Therapeutic - primary care** For the following indications, it is considered appropriate for primary care to supply and monitor **therapeutic dose LMWH**. Initial supply is usually from secondary care. *NB: list is not exhaustive / exclusive. GPs can prescribe and monitor for other patient groups if it is in the patient's best interest to do so.* In some cases, LMWH may be split into two divided doses as advised by haematology

Speciality	Indication	Usual duration* (or as per local Trust guidelines)	
Any speciality	<ul> <li>Unsuitable for warfarin or alternatives e.g.</li> <li>Poor compliance (i.e. likely or proven non- clinic attendance e.g. IVDU, homeless etc)</li> <li>Unable to attend AC clinic (housebound, bedbound) and with poor peripheral venous access (i.e. safety issue regarding monitoring)</li> <li>Anticipated poor INR control or failure to achieve therapeutic anticoagulation on VKA</li> <li>Intolerance to VKA or contraindication</li> <li>Excess alcohol, binge drinking</li> </ul>	For VTE, usually up to 6 months but may be longer as advised by haemostasis / haematology consultant	
Any speciality	Defined course of anticoagulation (e.g. superficial thrombophlebitis, provoked calf or proximal DVT, line associated thrombosis etc) and where oral anticoagulation with VKA is not suitable for clinical reasons	The duration of anticoagulation needs to be clearly defined by secondary care	
Any speciality	Warfarin patient requiring a 'bridging' plan around surgery or procedure and where the GP practice provides an AC service	This applies if the GP practice AC service includes 'bridging' anticoagulation, which should be in accordance with the relevant hospital's bridging guidelines. GP can refer to relevant anticoagulant clinic if needed.	
	The pre assessment clinic may request that the GP prescribes 3-5 days of LMWH proof the bridging plan. In such instances, the GP will be forwarded a patient specific prelevant guidelines as appropriate. Note: it is the duty of the surgical/medical team in charge of the patient's clinical constraints bridging guidelines in liaison with the patient's own AC clinic and pre-ass clinical team, to formulate a pre-operative bridging plan. If there is a date change to procedure, team to also communicate accordingly		
Oncology	Patient does not require, or has completed chemotherapy and is no longer under the direct care of the oncologist , but requires ongoing LMWH	Duration depends on indication and cancer status. First VTE event in setting of cancer: LMWH usually for at least 6 months then review by oncology; if ongoing active cancer, then AC is usually continued beyond 6mths although choice and dose can be reviewed at that point. Patient may require longterm LMWH	

### **3.4 Dose (therapeutic)**

- The relevant local Trust policies and standard reference sources (e.g. BNF) should be consulted
- For patients of **high body weight** (e.g. > 110kg for dalteparin, > 105kg for tinzaparin or > 100kg for enoxaparin) or **low body weight** (e.g. < 40kg), seek advice from the local haemostasis SpR
- In some cases, LMWH may be split into two divided doses as advised by haematology
- Antenatal / postnatal: therapeutic / higher intensity dose LMWH will be as per haemostasis consultant
- **Oncology**: On occasions, the dose may be higher or lower than the standard weight based BNF dose. This may depend on whether the patient is at a higher risk of bleeding or requires high intensity LMWH. The haemostasis/haematology cons. will advise accordingly. If this occurs, then it should be clearly documented by secondary care.

### 4.0 MONITORING OF LMWH

In general and unless otherwise specified by secondary care, the following is advised:

4.1 MONITORING - th	erapeutic dose LMWH	
Parameter	Comment	
FBC, U&E and LFTs every 4 to 6 weeks (or as clinically indicated; maximum interval 3 months)	<ul> <li>Enoxaparin and dalteparin: review dose if estimated CrCL* falls to &lt; 30mL/min.</li> <li>Tinzaparin: review treatment dose if CrCL* falls to &lt; 20mL/min</li> <li>Discuss with haemostasis SpR if there is any significant change in renal function or if CrCL falls below the values above</li> <li>Monitoring for HIT is not routinely required unless the risk is &gt; 1% (see section 5.0)</li> </ul>	
Weight every 4-6 weeks or as clinically indicated (maximum interval 3 months)	<ul> <li>Dalteparin: adjust dose in line with the BNF weight bandings (or as advised by secondary care)</li> <li>Tinzaparin / enoxaparin: review dose if weight changes by more than 5kg (or as advised by secondary care)</li> </ul>	

\* could use eGFR in place of CrCL for *most adult patients of average build and height as per BNF*. **But** for extremes of body weight, use CrCL with caution (see section 4.3 and appendix 2)

4.2 MONITORING – thromboprophylactic dose LMWH			
Parameter	Comment		
Clinical monitoring is not required	Patient should be advised (with this advice documented), to report bleeding/bruising symptoms to his/her doctor		
HIT monitoring is not routinely required unless the risk is > 1%	See section 5.0		

### 4.3 COMMENT RE RENAL FUNCTION - eGFR vs CrCL

The BNF states the following:

- Renal function is measured either in terms of estimated glomerular filtration rate (eGFR) normalised to body surface area (BSA) 1.73m<sup>2</sup> or it can be expressed as creatinine clearance, often calculated from the Cockcroft & Gault (C&G) formula.
- In adults, renal function is increasingly being reported on the basis of eGFR (mL/min/1.73m<sup>2</sup>). However, published information on the effects of renal impairment on drug elimination and manufacturer's dosing recommendations, are usually stated in terms of CrCL (C&G).
- Although eGFR and CrCL are not interchangeable, in practice and for *most adult patients of average build and height*, eGFR can be used to determine dose adjustments in place of CrCL.
- For patients at extremes of body weight (BMI < 18.5kg/m<sup>2</sup> or > 30kg/m<sup>2</sup>) the *absolute* glomerular filtration rate\* or CrCL (from C&G formula\*) should be used to adjust doses. Note that there are inherent inaccuracies when estimating renal function in certain patient populations, such as the very elderly, reduced muscle mass, poor nutritional status or at extremes of body weight. It is advised that estimates are used with caution and dosing advice sought as appropriate from the local anticoagulation teams

\*See appendix 2 for relevant calculations.

# 5.0 MONITORING FOR HIT (heparin induced thrombocytopenia)<sup>5,6</sup>

- The monitoring of HIT is only occasionally required and will affect relatively few patients
- From a GP perspective, HIT monitoring (if indicated), amounts to one platelet check between days 4-7 post starting LMWH and once again between days 10-14, (assuming the patient is still on LMWH)
- If HIT monitoring is required, secondary care should clearly document this on the discharge summary. Baseline platelet counts (i.e. pre LMWH) should be documented, as should any subsequent in-patient results. The discharge summary should be faxed to the GP as soon as possible pre-discharge with the monitoring requirement highlighted. The patient should be informed.
- Suspect HIT if platelets fall by 50% or more of pre-treatment baseline (even if the platelet count nadir remains above the lower limit of the normal range).
- In addition, consider whether HIT is a possible diagnosis if patient develops venous/arterial thrombosis or skin lesions at heparin injection sites.
- If HIT is suspected, stop LMWH and urgently contact the local haemostasis SpR for advice.

Recommendations for platelet	<b>Recommendations for platelet monitoring</b> (based on ACCP 2012 <sup>5</sup> and BCSH 2012 <sup>6</sup> recommendations)			
Secondary care should use this table to identify those patients requiring HIT monitoring. If this is required on				
discharge, then the secondary care	team should ensure that the GP is notified accordingly			
Patient type	Platelet monitoring for HIT			
LMWH only (prophylactic or therapeutic) and where: 1. the risk of HIT is < 1% (see incidence table below) AND	<ul> <li>Baseline platelet count</li> <li>Subsequent monitoring not required</li> <li>i.e. HIT monitoring is not required for all medical, obstetric and surgical patients (including orthopaedic). <i>Exception:</i> cardiothoracic surgery (where the incidence)</li> </ul>			
2. pt does not fall into the other heparin categories below	of HIT is 1-3%) and cancer patients undergoing surgery (where the risk of HIT is unclear but likely to be at least 1%).			
LMWH and HIT incidence > 1% (see incidence table below)	<ul> <li>Baseline platelet count</li> <li>Once between days 4-7 post starting LMWH</li> <li>Once again between days 10-14 whilst on LMWH</li> </ul>			
UFH (unfractionated heparin) during the current in-patient episode and now on LMWH	<ul> <li>Baseline platelet count</li> <li>Once between days 4-7 post starting UFH and</li> <li>Once again between days 10-14 whilst on LMWH</li> </ul>			
ANY type of heparin within the previous 100 days	<ul> <li>Baseline platelet count</li> <li>Check at 24 hours</li> <li>Thereafter as per other categories as appropriate</li> </ul>			
UFH (unfractionated heparin) infusion	<ul> <li>Baseline platelet count</li> <li>Check at 24 hours if UFH/LMWH has been administered within the previous 100 days</li> <li>Every 2-3 days from days 4-14 or until UFH is stopped (whichever occurs first)</li> </ul>			

Incidence of HIT according to patient population and type of heparin exposure (ACCP 2012 <sup>5</sup> )					
Patient population (min. of 4days exposure) Incidence Patient population (min. of 4days exposure) of HIT			Incidence of HIT %		
Post-operative patients		Medical			
Heparin prophylactic dose	1-5%	Cancer	1%		
Heparin therapeutic dose	1-5%	Heparin prophylactic of therapeutic dose	0.1-%1		
Heparin flushes	0.1-1%	LMWH prophylactic or therapeutic dose	0.6%		
LMWH prophylactic of therapeutic dose	0.1-1%	ITU patients	0.4%		
Cardiac surgery patients	1-3%	Heparin flushes	< 0.1%		
		Obstetric patients	< 0.1%		

### **6.0 IN-PATIENT DISCHARGES FROM SECONDARY CARE**

- The discharging team must ensure that the patient receives uninterrupted anticoagulation therapy (usual minimum of 2 weeks supply) until the patient can be reviewed by the GP or hospital clinic, whichever has been agreed. Note that some patient groups who will be managed by primary care (e.g. homeless, IV drug misusers), may not be safe for a 2 week supply. In these cases, dialogue with the GP pre discharge is required and early GP follow-up organised
- If a finite period of *thromboprophylaxis* is required and it is clinically appropriate to do, then the entire quantity of LMWH can be supplied on discharge (e.g. up to 6 weeks)
- The discharging team must also ensure that the patient or a relative is able to administer LMWH, otherwise referral to a district nurse should be made. For out of area patients, if self administration or district nurses are not viable options, then consideration should be given to Homecare services.
- A yellow sharps bin should also be provided by secondary care
- Clear documentation regarding anticoagulation management, MUST be included on discharge paperwork from secondary care (see section 7.0)
- The GP may occasionally be required to undertake HIT monitoring (section 5.0). In these cases secondary care should clearly document this on the discharge summary. Baseline platelet counts (i.e. pre LMWH) should be documented, as should any subsequent in-patient results. The discharge summary should be faxed to the GP as soon as possible pre-discharge with the monitoring requirement highlighted. The patient should be informed.
- If it is the intention that the GP should continue the prescribing and monitoring of LMWH, then the form in appendix 1 (or a suitably detailed letter or discharge summary outlining similar information) should be completed and faxed to the GP

# 7.0 INFORMATION PROVISION - between secondary and primary care (and vice versa)

For all transfers of care between primary/secondary care and *vice versa*, it is expected that the following information should be included on relevant paperwork. This is in line with good practice as per the NPSA<sup>1</sup>.

- Indication for anticoagulation
- Dose prescribed including intentional dose adjustments and rationale where appropriate
- Approximate start date and expected duration of therapy
- Who is responsible for clinical review and when
- Most recent weight in kg (with date) and an 'alert' weight (where LMWH dose adjustment needed)
- Baseline and recent blood results (e.g. Hb, platelets, serum creatinine)
- Renal function (section 4.3)
- Requirements for ongoing monitoring (section 4.0), including platelet monitoring for HIT if appropriate (section 5.0)
- Who is responsible for prescribing and monitoring
- Any additional relevant information (e.g. district nursing services required, patient selfadministering)

#### References

1. NPSA, National Patient Safety Agency Rapid Response Report (RRR 014): Reducing treatment dose errors with low molecular weight heparins. July 2010

2. NICE clinical guideline 92 Venous thromboembolism: reducing the risk

http://www.nice.org.uk/nicemedia/live/12695/47197/47197.pdf

3. Royal College of Obstetricians and Gynaecologists (RCOG) Green top guideline 37a: Reducing the risk of thrombosis and embolism during pregnancy and the puerperium. Nov 2009. <u>http://www.rcog.org.uk/files/rcog-corp/GTG37aReducingRiskThrombosis.pdf</u>

4. British national Formulary No 66. BMJ Group and the Pharmaceutical Press; London

5. Linkins et al. American College of Chest Physicians. Treatment and prevention of heparin induced thrombocytopenia: antithrombotic therapy and prevention of thrombosis 9th edition. ACCP clinical practice guidelines. *Chest* 2012; 141;e495S-e530S

6. Watson et al. BCSH Guidelines on the diagnosis and management of heparin induced thrombocytopenia: second edition. 2012; <u>http://www.bcshguidelines.com/4\_HAEMATOLOGY\_GUIDELINES.html</u>

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### Appendix 1: <u>LMWH</u> transfer to primary care within NCL

Secondary care: To complete this form OR to ensure that all information is included in a discharge summary or an out-patient letter to the GP

Consultant:					Hospital A	Hospital Address		
Speciality:								
Clinic date:								
Patient name:					Patient's a	Patient's address		
DOB:								
Hospital No:								
Dear Doctor					GP details			
The above patient requires ongoing subcutaneous treatment with								
injection, for the indication as outlined below. We have supplied								
and would be grateful if you could please continue to								
supply and monit	tor thereafter.							
Indication	Indication		Wt(Kg)+ date		Height + date	Dose & frequency	y Std BNF dose? Y / N	
Details of individualised dosing, if standard weight based BNF dosing not applicable:								
Start date:	Hb (date)	Plts (date)		Serum Cr (date)		CrCL (mL/min)	eGFR (mL/min/1.73m <sup>2</sup> )	
Proposed duration				Da	ate for review			
Responsibility for reviewing AC								
Other relevant conditions								
			N					
<ul> <li>Standard LMWH monitoring (unless otherwise advised):</li> <li>FBC, U&amp;E and LFTs every 4 to 6 weeks (or as clinically indicated, max. interval 3mths). Seek advice from haem SpR if renal function significantly deteriorates or if CrCL/eGFR &lt;30mL/min with enoxaparin /dalteparin or CrCL/eGFR &lt;</li> <li>20mL/min with tigagaparin (see section 4.3 of NCL LMMHL document regarding aCFR/CrCL)</li> </ul>								
<ul> <li>20mL/min with tinzaparin (see section 4.3 of NCL LMWH document regarding eGFR/CrCL).</li> <li>Check weight every 4-6 wks or as clinically indicated, max 3mthly. Tinzaparin/enoxaparin: review dose if</li> </ul>								
> 5kg weight change. Dalteparin: review dose in line with BNF weight bandings. Additional specific monitoring for this patient:								
Additional comments (including administration details if applicable):								
Thank-you for fo	llowing up If you	wish to	discuss fu	rthe	er nlease cont	act using the deta	ils helow	
Thank-you for following up. If you wish to discuss further, please contact using the details below.								
Yours sincerely:Print:Print:								
GradeDateDate								

Confidentiality info

# Appendix 2: Cockcroft-Gault and Absolute GFR

Cockcroft & Gault (C&G) Formula					
Estimated CrCL (mL/min)	= <u>(140-age) x weight x constant</u> Serum Cr				
Age in years					
• Weight in Kg (ideal be	odyweight but see below)				
• Serum Creatinine in r	nicromole/L				
• Constant = 1.23 (ma	ile) or 1.04 (female)				
Estimated ideal body weig O Males: O Females:	ght (IBW) kg IBW = 50 kg + 2.3 kg for each inch over 5 feet IBW = 45.5 kg + 2.3 kg for each inch over 5 feet				
, , ,	TBW in Kg) is ≤ 120% IBW, then use TBW in C&G formula TBW in kg) is > 120% IBW, then calculate and use adjusted body weight (ABW in Kg) TBW-IBW)]				

Absolute Glomerular Filtration Rate				
Actual GFR	= eGFR x (patient's BSA /1.73)			
BSA = Body surface area (formulas available via Internet)				

### Bibliography

Cockcroft DW and Gault H. Prediction of creatinine clearance from serum creatinine. Nephron 1976;16: 31-41. Basic Clinical Pharmacokinetics 4th edition; 2004. Michael Winter. Editor: DB Troy. Lippincott Williams& Wilkins, Philadelphia