

This Medicines Information Leaflet is produced locally to optimise the use of medicines by encouraging prescribing that is safe, clinically appropriate and cost-effective to the NHS.

Treatment of venous thromboembolism (VTE) in adults with dalteparin (Fragmin®)

Dalteparin is a low molecular weight heparin (LMWH) indicated for the treatment of deep vein thrombosis (DVT) and pulmonary embolism (PE) (collectively known as venous thromboembolism (VTE)) in adults.

The diagnosis should be confirmed radiologically (e.g. ultrasound, CTPA) since clinical diagnosis alone is inaccurate. Treatment should be started if imaging is delayed (for more than 1 hour for suspected PE or for more than 4 hours for suspected DVT) unless the risk of therapy is felt to outweigh the benefit, in which case this should be documented in the medical notes. Once the diagnosis is confirmed treatment should be continued or started immediately. When dalteparin is contraindicated, consider using unfractionated heparin (UFH) (see [MIL Vol.5 No.6 Guidelines on when to use and how to monitor unfractionated heparin in adults](#)) or fondaparinux (contact the on call haematology registrar for advice).

Advantages of dalteparin over UFH

- Dalteparin produces reliable anticoagulation rapidly, without the need for routine monitoring in most patients
- Dalteparin is convenient to administer and is as effective as UFH
- The dose of dalteparin is calculated from the patient's weight
- Dalteparin is administered subcutaneously
- The use of the subcutaneous route allows patients to be more mobile and self-caring and facilitates out-patient treatment and faster discharge

Mode of action

Dalteparin potentiates action of antithrombin; compared with UFH it results in less anti IIa (thrombin) and more anti-Xa activity.

Dose & administration

Patients starting full dose anticoagulation should have a FBC, U&Es and coagulation screen. In order to minimise the risk associated with complicated dose calculations and to reduce drug wastage, the Medicines Management & Therapeutics Committee (MMTC) has recommended that wherever possible pre-filled syringes of dalteparin are used. Various pre-filled syringes are available. Each syringe corresponds to a weight band (see Table 1). It is advisable to use the Powerplans within ePMA to aid safe prescribing. The license for dalteparin states to give 18,000 units to all patients 83kg and above. However, British and American guidelines as well

as local expert opinion recommend dosing based on actual body weight in patients who are 83kg and above.

Weight

It is imperative that the patient is weighed and that the weight is documented on the drug chart. In exceptional circumstances, when weighing the patient is not possible, the estimated weight must be documented on the drug chart. Patients who weigh less than 40kg are excluded from the product license and should be discussed with a Haematology SpR on bleep 5529 before extended treatment is initiated. Also consider discussing treatment with Haematology if the patient's weight exceeds 180 kg.

Pharmacy will not dispense dalteparin and nursing staff are at liberty to refuse to administer dalteparin from a prescription on which the weight is not documented.

Table 1: Standard dalteparin dose recommendations for treatment of VTE: Should be used in all cases except where there is risk of bleeding (see recommendations below).

Month 1

Body weight (kg)	Dose of dalteparin by subcutaneous injection using a pre-filled syringe (units)
Less than 46 Consider discussing with haem SpR if less than 40kg	7,500 once daily
46-56	10,000 once daily
57-68	12,500 once daily
69-82	15,000 once daily
83-98	18,000 once daily
99-112	10,000 twice daily
113-137	12,500 twice daily
138-165	15,000 twice daily
166 or more Consider discussing with haem SpR if greater than 180kg	18,000 twice daily

Single doses should not exceed 18,000 units

Table 2. Month 2 onwards standard dalteparin dose recommendations for treatment of VTE.

If patients are to remain on dalteparin beyond the first month as treatment for a VTE, ensure the patient is reweighed and dose reduce according to the table below:

Month 2 onwards- Ensure patient is reweighed

Body weight (kg)	Dose of dalteparin by subcutaneous injection using a pre-filled syringe
Less than 57	7,500 once daily
57-68	10,000 once daily
69-82	12,500 once daily
83-98	15,000 once daily
99-112	18,000 once daily
113-137	10,000 twice daily
138-165	12,500 twice daily
166 or more	15,000 twice daily

Patients with cancer

Randomised controlled trials have recently evaluated the use of DOACs in patients with active cancer. When compared directly with LMWH, DOACs are as effective at treating VTE as LMWH. Please see [MIL Vol 8, No 1](#) for more details.

Risk factors for bleeding include:

- Liver disease
- Renal impairment
- History of peptic ulcer disease
- Concomitant medicines that may enhance anticoagulant effect
- Alcohol abuse
- Severe hypertension (systolic greater than 180mmHg, diastolic greater than 110mmHg)
- Congestive heart failure
- Doses based on incorrect patient weight

Note, in patients who are considered to be at an increased risk of bleeding, administration of dalteparin in divided daily doses of 100 units per kg subcutaneously twice daily may be safer.

Contra-indications (principally as for UFH)

- Known hypersensitivity to dalteparin
- Cerebral haemorrhage, acute gastric or duodenal ulceration, known haemorrhagic diathesis.
- Uncontrolled severe hypertension
- Injuries to or recent operations to the ears/eyes or central nervous system
- Infective endocarditis
- Severely disturbed liver or renal function
- Regional anaesthesia is contra-indicated in patients receiving dalteparin
- History of Heparin-Induced Thrombocytopenia (HIT)

Oral anticoagulation with warfarin

Warfarin should be started concurrently. Treatment with dalteparin should continue in parallel with oral anticoagulation for five days or until the patient's INR has been greater than or equal to 2 for at least 24 hours (two

consecutive days), **whichever is longer**. For more information, please see [MIL Vol.5 No.8](#) *Initiating Oral Anticoagulation in Adult Patients*.

Monitoring

A baseline coagulation screen and platelet count should be taken in all cases. Monitoring of the anticoagulant effect is not normally necessary for dalteparin prescriptions. However, for certain patients (see HIT, below) monitoring of platelet counts is recommended every 2-4 days from days 4 to 14 of treatment.

Inhibition of aldosterone secretion by unfractionated or low molecular weight heparin can cause result in hyperkalaemia in susceptible patients (e.g. patients with diabetes, chronic renal failure, or acidosis, or those taking potassium sparing drugs). If such patients are given dalteparin for longer than 7 days potassium should be monitored.

Plasma anti-Xa concentration can be used to monitor the anticoagulant effect of dalteparin, such as in patients with renal impairment or if abnormal coagulation parameters or bleeding should occur during therapy. Maximum plasma concentration is obtained 3-5 hours after subcutaneous injection, when samples should be taken. For patients on once daily dosing, the expected peak plasma concentration is about 1.0 anti-Xa unit per mL with a range of 0.5 – 1.5 (and 0.5-1.0 anti-Xa units per mL for twice daily dosing).

Use in renal impairment

Anticoagulation with heparins in the presence of significant renal impairment is not straightforward as both low molecular weight heparin and unfractionated heparin are likely to accumulate when the CrCl falls below 20 ml/min. In addition bleeding complications are likely to be more severe in patients with renal impairment because platelet function is impaired. For these reasons if treatment dosages of heparin are required in patients with a CrCl less than 20 ml/min there are two options:

- Use subcutaneous dalteparin with 2/3 of the normal weight adjusted dosage (see Table 3 for dose banding recommendations) and monitoring of anti-Xa plasma concentration may be considered, although the correlation between anti-Xa level and bleeding risk is very weak. The advantage of this is that the response is more predictable, but the disadvantage is that initially monitoring is still required and LMWH is not readily reversible.

OR

- Use intravenous UFH and monitor the APTT appropriately. The advantage of UFH is that it can easily be stopped if necessary and its effects wear off rapidly. It can also be easily reversed with protamine. The disadvantage is that achieving adequate anticoagulation is unpredictable and dosage adjustments based on appropriately timed APTT measurements require careful management. For more information, see [MIL Vol.5 No.6](#) *Guidelines on when to use and how to monitor Unfractionated Heparin in adults*.

Table 3: Dalteparin dose recommendations for treatment of venous thromboembolism (VTE) in patients with significant renal impairment (CrCl less than 20mLs/min) in Month 1.

Body weight (kg)	Dose of dalteparin by subcutaneous injection using a pre-filled syringe
Under 46	5,000 units daily
46-56	6,500 units daily*
57-68	8,500 units daily*
69-82	10,000 units daily
83-98	12,500 units daily
99-112	15,000 units daily
113-137	18,000 units daily
138-165	10,000 units twice daily
166 or more	12,500 units twice daily

* Graduated syringes containing 10,000 units in 1mL of dalteparin are available to administer these doses

Adverse effects

- Commonly reported adverse effects include subcutaneous haematomas at the site of injection
- Systemic bleeding is a rare complication of treatment with dalteparin.
- Heparin-induced thrombocytopenia (HIT) has been reported in association with low molecular weight heparins and is an indication for immediate cessation of treatment.

Heparin products can cause hypoadosteronism which may result in an increase in plasma potassium. Rarely, clinically significant hyperkalaemia may occur, particularly in patients with chronic renal failure and diabetes mellitus.

Patients with platelet counts which are less than 50 or expected to fall to less than 50

Therapeutic anticoagulation is not generally used in patients who have platelet counts less than 50 due to the risk of bleeding. If patients have an acute VTE then anticoagulation should be carefully considered. Please consider discussing with the Haemostasis SpR Bleep 5529 or Consultant.

Pregnancy and lactation

Dalteparin has been assessed in pregnant women and no harmful effects are known with respect to the course of pregnancy and the health of the unborn and neonate. Dosing in pregnant patients may vary from those quoted above due to differences in the volume of distribution of dalteparin in pregnancy. Separate guidance is available from the Women's Services Directorate.

Overdose / reversal

In an emergency the anticoagulant effect of dalteparin can be partially reversed by protamine sulphate. One mg of protamine sulphate inhibits the effect of 100 units (anti-Xa) of dalteparin. The usual maximum dose is 50 mg given by slow IV injection (rate not exceeding 5 mg per minute). The Haemophilia and Thrombosis Centre can be contacted for further advice. (Out of hours, contact the on call haematology registrar via switchboard)

Dalteparin and intramuscular injections

Intramuscular injections should be avoided in patients receiving anticoagulants, except for adrenaline in severe anaphylaxis.

Dalteparin and surgery

Separate guidelines are available for the peri-operative management of anticoagulation (see [MIL Vol.10 No.5](#) for full guidance). If necessary, contact the on call haematology registrar for advice.

Heparin-induced thrombocytopenia (HIT)

Clinically important HIT is rare with LMWH except in patients receiving the drug in some post-operative settings. Evidence suggests the risk of developing HIT with LMWH is greatest in patients who have undergone cardiac surgery, and that other patients do not require monitoring. The more common type of HIT is immune-mediated and does not normally develop until 5-10 days after starting unless the patient has been exposed to heparins in the previous 100 days. All patients who are to receive dalteparin should have a platelet count on the day of starting therapy. For patients receiving the drug after cardiac surgery, check the platelet count every 2-4 days between days 4 to 14 (or from day 1 if the patient has been exposed to heparins in the previous 100 days). All other patients do not require platelet count monitoring unless they show signs of HIT such as thrombosis or skin allergy. If HIT is strongly suspected or confirmed, dalteparin should be stopped and an alternative anticoagulant, such as danaparoid, lepirudin, bivalirudin, argatroban or fondaparinux should be given (see [MIL Vol. 5 No. 10 Alternative Anticoagulants for use in Heparin-Induced Thrombocytopenia in Adults](#) or contact the on call haematology registrar for advice).

Safe medication practice

- Dalteparin should always be prescribed with "UNITS" written in full.
- When using pre-filled single dose syringes, to ensure delivery of the full dose, do not expel the air bubble from the pre-filled syringe before injection
- Dalteparin doses are weight-based. Ensure that the patient's weight is documented on **ALL** prescriptions for dalteparin.

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