

# Who can administer

May be administered by registered competent doctor or nurse/midwife

# Important information

- Ordered on advice of consultant haematologist only
- Unlicensed preparation in Ireland
- IMPORTANT: The ACT is NOT the same as the aPTT. Results are not interchangeable.
- See under Dose for adjustments required in renal or hepatic impairment

## Available preparations

Exembol 250mg per 2.5ml vial (contains ethanol 1g / 2.5ml vial)

## Reconstitution

- Already in solution
- Dilute further prior to administration

## Infusion fluids

- Sodium chloride 0.9% or Glucose 5%
- Mix by repeated inversion of the infusion bag for 1 minute

### Methods of intravenous administration

#### Continuous intravenous infusion (administer using an electronically controlled infusion device)

- Add 250mg (2.5ml) vial to 250ml infusion fluid (no need to remove 2.5ml from bag)
- Using a 1mg per ml solution (250mg in 250ml) set up the continuous infusion
- Adjust dose as per 'Dose'

# Slow intravenous injection (Percutaneous intervention only, see dose under Further Information)

• Given over 3 to 5 minutes

## Dose in adults

#### Heparin induced thrombocytopenia (HIT)

- All patients with HIT must be managed in conjunction with haematology
- If baseline aPTT elevated- discuss target aPTT with Haematology
- **APTTr;** the patient's activated partial thromboplastin time divided by either the laboratory's normal value or the patient's own baseline value
- Prior to commencement **stop heparin** and take baseline aPTT, PT and fibrinogen (CLAUSS) (ref 1)
- Argatroban is a direct thrombin inhibitor. Its anticoagulant effect is measured by the APTTr and INR, which increase in a dose dependent manner. Dose adjustments are based on the APTTr as shown in the

table below.

- Daily monitoring is sufficient after 2 consecutive aPTT within target range if no dose adjustments were made (ref 1)
- Argatroban interferes with the Clauss Fibrinogen assay resulting in falsely low results. The derived fibrinogen assay may be more reliable but both will be abnormal at higher argatroban concentrations<sup>(ref 1)</sup>
- The lab MUST be informed that patient is receiving argatroban<sup>(ref 1)</sup>

 Table 1: Recommended starting doses and monitoring intervals (see table 3 below for flow rates)

|                                  | Argatroban dose<br>(mcg/kg/minute) | Interval to check aPTT after<br>initial dose and each dose<br>change thereafter |
|----------------------------------|------------------------------------|---|
| Standard dose                    | 2                                  | 2 hours   |
| Critically ill patients*         | 0.5                                | 4 hours   |
| Moderate hepatic<br>impairment** | 0.5                                | 4 hours   |

\* Multi-system organ failure, ICU patients, heart failure, post cardiac surgery, anasarca

\*\* Moderate hepatic impairment (Child Pugh B)

| Table 2: Dose modifications and monitoring intervals (see Table 3 below for flow rates) |   |   |   |   |  |  |
|---|---|---|---|---|--|--|
| Standard dosing schedule  |   | Critically ill/Hepatically impaired patients  |   |   |  |  |
| Initial infusion rate 2mcg/kg/minute  |   | Initial infusion rate 0.5mcg/kg/min   |   |   |  |  |
| aPTT  | Infusion rate change  | Next aPTT   | Infusion rate change  | Next aPTT   |  |  |
| <1.5<br>patient's<br>baseline   | Increase by<br><b>0.5</b> mcg/kg/min  | 2 hours   | Increase by<br><b>0.1</b> mcg/kg/min  | 4 hours   |  |  |
| 1.5 to 3<br>times<br>patient's<br>baseline<br>(and less<br>than 100<br>seconds)         | No change<br>Ideal range: 1.5 to 2.5<br>patient's baseline aPTT<br>as per Dr Gilmore  | 2 hours<br>After 2<br>consecutive<br>aPTT within<br>target range,<br>check at least<br>24 hours | No change   | 4 hours<br>After 2<br>consecutive<br>aPTT within<br>target range,<br>check at least<br>every 24 hours |  |  |
| > 3 times<br>patient's<br>baseline or if<br>over 100<br>seconds                         | Discontinue infusion<br>until APTT is within<br>desired range (typically<br>within 2 hours of<br>discontinuation) - restart<br>at 50% of the previous<br>rate | 2 hours   | Discontinue infusion<br>until APTT is within<br>desired range (typically<br>within 2 hours of<br>discontinuation) -<br>restart at 50% of the<br>previous rate | 4 hours   |  |  |

• The maximum recommended dose is 10 microgram/kg/minute

• The maximum recommended duration of treatment is 14 days, although there is limited experience with administration for longer periods

| Table 3: Infusion rate in ml/HOUR of a 1mg/1ml infusion |                      |                              |                            |                            |  |  |
|---|----------------------|------------------------------|----------------------------|----------------------------|--|--|
| Dose/Weight<br>(kg)                                     | 0.1micrograms/kg/min | 0.5micrograms/kg/ <b>min</b> | 1micrograms/kg/ <b>min</b> | 2micrograms/kg/ <b>min</b> |  |  |
| 50  | 0.3                  | 1.5                          | 3                          | 6                          |  |  |
| 60  | 0.36                 | 1.8                          | 3.6                        | 7.2                        |  |  |
| 70  | 0.42                 | 2.1                          | 4.2                        | 8.4                        |  |  |
| 80  | 0.48                 | 2.4                          | 4.8                        | 9.6                        |  |  |
| 90  | 0.54                 | 2.7                          | 5.4                        | 10.8                       |  |  |
| 100   | 0.6                  | 3                            | 6                          | 12                         |  |  |
| 110   | 0.66                 | 3.3                          | 6.6                        | 13.2                       |  |  |
| 120   | 0.72                 | 3.6                          | 7.2                        | 14.4                       |  |  |
| 130   | 0.78                 | 3.9                          | 7.8                        | 15.6                       |  |  |
| 140   | 0.84                 | 4.2                          | 8.4                        | 16.8                       |  |  |
| 150   | 0.9                  | 4.5                          | 9                          | 18                         |  |  |

#### Patients with HIT Type II undergoing percutaneous coronary intervention (PCI)

- Limited data is available from the use of argatroban in patients with HIT Type II undergoing percutaneous coronary intervention
- Based on the data, when there is no alternative, therapy could be initiated with a bolus dose of 350 microgram/kg over 3 to 5 minutes
- This is followed by an infusion dose of 25 microgram/kg/min
- ACT should be checked 5 to 10 minutes after the bolus dose is completed
- The procedure may proceed if the ACT is greater than 300 seconds
- If the ACT is below 300 seconds, an additional bolus dose of 150 microgram/kg should be administered, the infusion rate be increased to 30 microgram/kg/min, and the ACT should be checked 5 to 10 minutes later
- If the ACT is higher than 450 seconds the infusion rate should be decreased to 15 microgram/kg/min and ACT values be checked 5 to 10 minutes later
- Once a therapeutic ACT between 300 to 450 seconds has been achieved, the infusion dose should be continued for the duration of the procedure
- ACT measurements were recorded using both Haemotec and Haemochrom devices
- The efficacy and safety of argatroban use in combination with GPIIb/IIIa inhibitors has not been established.

#### **Renal impairment**

- No **initial** dosage adjustment necessary in mild to severe renal impairment. Monitor aPTT closely.
- Haemodialysis: limited data. Consult haematologist and see specialist texts

#### Liver impairment

- Use with extreme caution in hepatic impairment
- See Tables 1 and 2 above for doses
- Reversal of the anticoagulant effects of argatroban may take longer in this setting (more than four hours)
- Argatroban is **contraindicated** in patients with severe hepatic impairment

#### Changing to oral anticoagulation (ref 1)

- A Haematology Consultant (ideally with Special Interest in Coagulation) will decide on the timing, duration and choice of oral anticoagulant
- Patients with HIT require a minimum of three months of oral anticoagulation
- If starting DOAC
  - $\circ~$  stop argatroban and start DOAC ~
  - $\circ\,$  Rivaroxaban, apixaban and dabigatran have all been used in this situation
- If starting warfarin therapy for patient on argatroban
  - Warfarin should only be commenced when there is resolution of thrombocytopenia (to avoid coumarin-associated microvascular thrombosis and venous limb gangrene)
  - $\circ~$  To avoid prothrombotic effects and to ensure continuous anticoagulation, argatroban must be continued during the initiation of warfarin therapy
  - $^\circ$  A minimum overlap of argatroban and warfarin of at least 5 days is advised
  - $\circ\,$  Argatroban has a significant effect on the INR
  - NO loading doses of warfarin to be given
  - $\circ\,$  Start with the intended maintenance dose of warfarin (no greater than 5mg daily)
  - Discontinue argatroban when INR reaches up to 4 for at least two days, on COMBINED therapy (INR should be 2 greater than desired target range- eg in patients with target INR of 2 to 3, INR on combined argatroban and warfarin should be 4 to 5) <sup>(ref 3)</sup>
  - Repeat INR 4 to 6 hours after stopping argatroban, to ensure INR is therapeutic prior to permanent discontinuation of argatroban<sup>(ref 3)</sup>
  - If INR is below the desired therapeutic range recommence argatroban and repeat the procedure above (steps 2 to 6)
  - Take INR at least daily

# Monitoring

• See above

## Storage

- Store below 25<sup>°</sup>C
- Do not refrigerate or freeze
- The diluted solution is stable for 24 hours if kept at  $25^{\circ}$ C or less (do not expose the diluted solution to direct sunlight)

## References

UK SPC (Exembol) 31/10/2024

1. Dr Ruth Gilmore, Consultant Haematologist, 26/03/2025

2. Heparin induced thrombocytopenia - a comprehensive clinical review, Salter et al. JACC Vol 67, No 21, 2016

# Therapeutic classification

Parenteral anticoagulants

BNF Blood clots