Improving safety of unfractionated heparin: a retrospective, quasi-experimental, observational study of the impact of a pocket card and a computerised prescription aid tool in the University Hospitals of Geneva

Wedali E Jimaja 1, Jerome Stirnemann 1,2, Pierre Fontana 2,4, Katherine S Blondon 2,4

ABSTRACT

Background Despite the rapid rise of direct oral anticoagulants, unfractionated heparin (UFH) remains the mainstay anticoagulant in specific situations such as severe renal failure, perioperative setting or in critical care units. However, its titration is often challenging.

Objectives To investigate the effect of a pocket card and a computerised prescription aid tool (CPAT) on the quality of UFH anticoagulation.

Design Monocentric retrospective, quasi-experimental, observational study.

Setting Inpatient primary care centre between 1 January 2016 and 31 December 2019.

Participants >18 years-old treated with therapeutic UFH for more than 24 hours. There were 819 and 1169 anticoagulation episodes before and after intervention, respectively.

Intervention In October 2017, we implemented a pocket card with evidence-based recommendation for therapeutic UFH initiation, monitoring and dosing adaptation. In October 2019, we implemented a CPAT in a group subset.

Primary and secondary outcomes The primary outcome was the time needed to reach a therapeutic anti-Xa level before and after the implementation of the pocket card. The secondary outcomes included a subgroup analysis assessing the effect of the CPAT. Other secondary outcomes were the anti-Xa status (infratherapeutic, therapeutic or supratherapeutic) at 7 and 24 hours of UFH treatment.

Results We found a significant increase in the time to reach therapeutic dosing with pocket card-guided recommendations implementation (10.1 vs 14 hours, HR of 0.8, 95% CI: 0.70 to 0.93). However, the CPAT was associated with a significant decrease in the time needed to reach the therapeutic range (13.9 vs 7.1 hours, HR of 1.74, 95% CI: 1.17 to 2.60).

Conclusion Although we observed an increase in time to reach therapeutic anti-Xa with the pocket card, possibly due to a selection bias (use of activated partial thromboplastin time for monitoring before the pocket card), the implementation of CPAT significantly decreased the delay for effective therapy. Further studies are needed to confirm these findings, and to determine the optimal initial dose of UFH anticoagulation.

INTRODUCTION

The anticoagulant properties of unfractionated heparin (UFH) have been known for about a century, and UFH has been prescribed for various indications (eg, prevention and treatment of thrombosis, acute coronary syndrome, coronary bypass graft or haemodialysis). In the recent years, numerous other anticoagulant molecules have emerged, for instance low molecular weight heparin, fondaparinux and direct oral anticoagulants (DOACs), associated with several advantages including no systematic need for ongoing therapeutic drug monitoring. However, UFH remains the mainstay treatment in specific clinical situations such as in severe renal insufficiency, perioperative setting and critical care owing to its short half-life and the availability of a specific antidote.
Due to the large interindividual pharmacokinetic variability, UFH requires close laboratory monitoring and titration to keep anticoagulation within the target range. The activated partial thromboplastin time (aPTT) remains the most commonly used method for this monitoring in spite of its standardisation issues and limitations such as variable levels of clotting factors masking the UFH anticoagulant effect. The target aPTT (1.5–2.5 times basal aPTT) is based on a 1970s study. The anti-Xa chromogenic assay is a valuable alternative that does not have the limitations of aPTT and is more standardised. Its therapeutic range is set to 0.3–0.7 based normograms. Anti-Xa-based monitoring of UFH has been shown to shorten the time needed to reach the therapeutic range, to reduce the number of blood sampling and UFH dose adjustment compared with aPTT-based monitoring, but is slightly more expensive than aPTT. Moreover, the anti-Xa assay is less available worldwide and is sensitive to other anticoagulants such as DOACs.

Although an anti-Xa-based monitoring trial showed improvement in a surrogate outcome (decrease of red blood cell transfusions), trials comparing aPTT and anti-Xa-based monitoring for clinical outcomes have not found clear benefits.

Beyond the issue of the most appropriate monitoring assay, dose adaptation is often poorly standardised in hospitals, although some authors such as Schurr et al. proposed algorithms that try to organise such a standardisation. In this retrospective study, they examined the effect of nurse-driven UFH non-customisable weight-based normograms on the time elapsed to reach therapeutic anticoagulation. In the group using normograms, they observed a decrease in the time needed to reach therapeutic anticoagulation.

The aim of the present study was to investigate the impact of a pocket card exhibiting recommendations on UFH dosing according to anti-Xa result as well as a computerised prescription aid tool (CPAT) on the quality of UFH anticoagulation for therapeutic (as opposed to prophylactic) treatment in an inpatient setting.

MATERIALS AND METHODS
Setting
This was a single-centred retrospective, quasi-experimental, observational study that took place in the University Hospitals of Geneva. The data were obtained for the period of 1 January 2016 to 31 December 2019 and were extracted from the hospital’s electronic medical records (EMR).

Population
This study included hospitalised patients that were 18 years of age or older, who received a continuous intravenous infusion of UFH aiming for a therapeutic anticoagulation for at least 24 hours. This 24-hour threshold was chosen to exclude patients with acute coronary syndrome, who usually receive therapeutic dose of UFH for a short period before catheterisation. We included patients with UFH doses of ≥10 000 IU/day, which we considered as most likely prescribed for a therapeutic anticoagulation. Patients receiving doses below this threshold but with explicit mention of a therapeutic anticoagulation indication in the medical record were also included. Patients with anticoagulation episodes that were not associated with at least one monitoring test over the therapeutic threshold (anti-Xa value ≥0.5 IU/mL) were excluded from the analyses, as they were assumed to be prescribed as prophylactic purpose. Those with less than one value of anti-Xa per day were also discarded as this suggests that aPTT was used for the monitoring of the UFH therapy. Finally, patients staying in neurology, neurosurgery and haematology units were not included in the analysis as they were assumed to often not follow the usual protocol due to the very high bleeding risk associated their corresponding pathologies.

Two groups were defined: a baseline pregroup with patients included and followed up to the 1st of October 2017, when the pocket card was implemented, and another group referred to as the postgroup with patients included thereafter.

Intervention
In our institution, both aPTT and/or anti-Xa assays were used for UFH monitoring at the beginning of the period studied. In October 2017, a pocket card guideline was distributed to all the physicians to standardise the preparation, prescription and administration of therapeutic UFH (available at https://www.hug.ch/sites/interhug/files/a6_heparine_web2.pdf, or online supplemental file 1) for an English version). The card indicated the initial dose according to weight, as well as instructions to adjust the dosage according to anti-Xa values based on a protocol published in 2010 by Smith et al. The guideline recommended using anti-Xa assay 4–6 hours after introducing or modifying UFH dosing. It also proposed a platelet count 2–3 times a week during the first 2 weeks of UFH treatment. The content of the guideline was also available in the computerised UFH prescription aid tool implemented in October 2019 in the internal medicine wards. This CPAT proposes initial and titration doses of UFH according to the patient’s weight and anti-Xa results, and adds an anti-Xa assay 6 hours after each change of dose: prescribers can choose to bypass the recommended UFH doses at any time.

Hospital stays and UFH anticoagulation episodes
We considered each hospital stay as the main unit for our analysis, therefore patients hospitalised more than once during the study period have multiple hospital stays. An UFH anticoagulation episode was defined as an UFH treatment of more than 24 hours during one hospital stay, with possible interruptions of less than 48 hours. During one hospital stay, a patient could therefore have more than one UFH anticoagulation episode. Because we supposed that the therapeutic range would be reached...
faster on a second episode based on the data of the first episode, only the first anticoagulation episode was used for each hospital stay. An anti-Xa value was associated with a UFH treatment modification if the blood sample was collected more than 4 hours after initiation of treatment. The therapeutic range was defined as an anti-Xa value between 0.3 and 0.7 IU/mL.

**Data collection**

Data were extracted from the EMR system to obtain patient demographics (age, gender, comorbidities and indication of anticoagulation), type of ward and length of stay, anti-Xa results (sampling time and result value), renal function, UFH initiation time, and dose modification.

Comorbidities were used to calculate the Multipurpose Australian Comorbidity Scoring System (MACSS) index. It is a comorbidity score integrating 102 conditions, with higher scores representing the presence of more comorbidities.15

**Statistical analysis**

**Primary outcome**

The primary outcome was the time needed to reach anti-Xa within the therapeutic anticoagulation range after UFH initiation. The Kaplan-Meier method was used to compute the cumulative incidence of anti-Xa reaching the therapeutic range for each group and the log-rank test was used to compare the two. Cox proportional hazards regression analysis was used to obtain non-adjusted and adjusted HRs.

**Secondary outcomes**

Subgroup analysis on events occurring after the 1st of October 2019 was also performed to explore the effect of a newly implemented CPAT recapitulating the pocket guide. This analysis was also performed using the Kaplan-Meier method and the log-rank test.

We investigated the difference between the two groups in terms of anti-Xa status after 7 hours and 24 hours of anticoagulation, respectively. We chose a first timepoint at 7 hours after the start of UFH infusion to decrease the risk of missing the first anti-Xa dosing. Indeed, even though the recommended time is between 4 and 6 hours, some samples may be collected beyond this interval. The anti-Xa status was defined as therapeutic, if at least one anti-Xa reached the therapeutic range during the time interval. It was deemed infratherapeutic or supratherapeutic if the last measured anti-Xa value was below or above the therapeutic range, respectively. In the case where no anti-Xa was measured during the considered period, the status was defined as missing. For the anti-Xa status after 7 hours of anticoagulation, missing values were not included in the analysis as they represented a significant amount of the overall values and because our aim was to assess adherence to pocket guide recommendations, which use anti-Xa monitoring. Within the supra-therapeutic status, the anti-Xa with values higher than 1 IU/mL were analysed separately, as they represent a significant increase in the risk of major bleeding events.16

To further explore the effect of the CPAT, we conducted the same anti-Xa status analyses comparing the CPAT subgroup with the episodes in the postgroup in internal medicine units occurring before the introduction of the CPAT. This latter group was named post-Internal Medicine (post-IM) group. Fisher’s exact test was used in this subgroup analysis.

In tables 1–3, t-test and $\chi^2$ test were used for continuous and categorical, respectively. Continuous variables were expressed as mean and SD and categorical variables as percentages. The RStudio software (V.1.2.5033, 2009–2019 RStudio) was used to perform the statistical analyses.

<table>
<thead>
<tr>
<th>Table 1 Patients characteristics</th>
</tr>
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<tbody>
<tr>
<td>Variable</td>
</tr>
<tr>
<td>Number of hospitalisation events</td>
</tr>
<tr>
<td>Number of patients</td>
</tr>
<tr>
<td>Age (years)</td>
</tr>
<tr>
<td>Weight (kg)</td>
</tr>
<tr>
<td>Length of stay (days)</td>
</tr>
<tr>
<td>Female (%)</td>
</tr>
<tr>
<td>eGFR &lt;30 mL/min/1.73 m² (%)</td>
</tr>
<tr>
<td>Anticoagulation indications*</td>
</tr>
<tr>
<td>AF</td>
</tr>
<tr>
<td>VTE</td>
</tr>
<tr>
<td>Mechanical valvular replacement</td>
</tr>
<tr>
<td>Wards:</td>
</tr>
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<td>Surgery</td>
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<td>Geriatrics</td>
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<tr>
<td>Others</td>
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<td>3+</td>
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</tbody>
</table>

*Other indications included: intracerebral venous thrombosis, inferior limb arterial thrombosis, intracardiac thrombus, splenic vein thrombosis, renal vein thrombosis and portal thrombosis, AF, atrial fibrillation or flutter; eGFR, estimated glomerular filtration rate; MACSS, Multipurpose Australian Comorbidity Scoring System (with higher score representing a more comorbid state); VTE, venous thromboembolism.

Patient and public involvement
Due to the retrospective design of the study, patients were not involved in its design and the results have not been disseminated to the participants.

RESULTS
From the 7757 hospital stays of patients treated with UFH during the study period, 1988 met the inclusion criteria, the main reason for exclusion being an anticoagulation duration inferior to 24 hours. Out of these, 819 stays from 777 patients were part of the pregroup, while 1169 stays of 1054 patients were part of the postgroup.

The mean ages were 71.3 (±14.8) and 73.5 (±14.8) years old for the pre and postgroups, respectively. There was a predominance of male gender, with female gender representing 40.2% and 40.9% in pre and postgroups, respectively (p=0.78). The most prevalent identified indications for anticoagulation were atrial fibrillation/flutter and VTE in both groups, and most patients were in either surgery or general internal medicine wards (table 1).

Primary outcome
Heparin anticoagulation quality and efficiency
A total of 819 and 1169 anticoagulation episodes were extracted in the pre and postgroups, respectively. After removing the episodes without anti-Xa values within the 7 hours after the beginning of anticoagulation, 310 episodes remained in the pregroup and 535 in the postgroup. For the pregroup, the median time to reach an anti-Xa within therapeutic range was 10.1 hours (95% CI: 7 to 13 hours) and for the postgroup, it was 14 hours (95% CI: 13 to 15 hours), with a statistically significant difference (p<0.01). The HR of reaching a therapeutic anti-Xa is 0.8 for the postgroup (95% CI: 0.70 to 0.93, p<0.01) when comparing both groups. When adjusted for age, wards, gender and weight, the statistical significance remains, with a HR of 0.8 of the postgroup to reach therapeutic anticoagulation (95% CI: 0.69 to 0.92, p<0.01) (figure 1).
Secondary outcomes

Time to reach therapeutic anticoagulation in subgroup analysis

In the subgroup analysis, the postgroup was divided into two groups: one with hospital stays occurring after the 1st of October 2019 in the internal medicine ward that used the CPAT, and those occurring before, that did not. The analysis compared only hospital stays in the internal medicine wards; 424 episodes were found in the post-IM group, and 28 in the CPAT group.

The median time to reach the therapeutic range was 13.9 hours (95% CI: 11.9 to 15 hours) in the post-IM group, whereas it was 7.1 hours (95% CI: 6.8 to 15.8 hours) in the CPAT group, with a p<0.01 (figure 2).

When compared with the post-IM group, the CPAT group exhibited a HR of 1.74 (95% CI: 1.17 to 2.60, p<0.01).

After performing an adjusted analysis, the CPAT group had a HR of 1.77 (95% CI: 1.18 to 2.65, p<0.01) of having a therapeutic anti-Xa value compared with the post-IM group.

Anti-Xa status in the 7 first hours of anticoagulation

The anti-Xa status were first checked after 7 hours of UFH infusion. In the pregroup, 509 (62%) episodes had no anti-Xa measured, compared with 634 (54%) in the postgroup, with a statistically significant difference (p<0.01).

Initial UFH dosing was 276 IU/kg (IQR=111 IU/kg) in the pregroup and 336 IU/kg (IQR=139 IU/kg) in the postgroup, with a p<0.01. After 7 hours of UFH infusion, the doses were 278 IU/kg (IQR=111 IU/kg) and 339 IU/kg (IQR 140 IU/kg) in the pre and postgroups, respectively, p<0.01.

The percentage of episodes reaching the therapeutic range was higher in the pregroup than in the postgroup with a statistically significant difference (151 (49%) vs 214 (40 %) episodes, p=0.02).

Concerning the episodes with an infratherapeutic status, 118 (38%) were found in the pregroup and 205 (38%) in the postgroup with no statistical significance for these results (p=1). For the supratherapeutic episodes, however, there were more episodes in the postgroup than in the pregroup (116 (22%) vs 41 (13%), with a p value<0.01). Among these supratherapeutic episodes, 15 (5%) had anti-Xa above 1 IU/mL in the pregroup and 48 (9%) in the postgroup, with a statistically significant difference (p=0.04) (table 2).

In the subgroup analysis, we compared the CPAT group with the post-IM group. No significant difference was observed between the two groups for therapeutic anti-Xa (59% vs 43%, OR 1.94, 95% CI: 0.73 to 5.39, p=0.18). While there were no infratherapeutic episodes (0%) in the CPAT group, there were 71 infratherapeutic episodes (34%) in the post-IM group. There were 9 (41%) supratherapeutic episodes in the CPAT group and 49 (22%) in the post-IM group. Episodes with an anti-Xa >1 IU/mL were 1 (5%) in the CPAT group and 23 (11%) in the post-IM group. Six episodes had no anti-Xa values within the first 7 hours in the CPAT group, and there were 215 episodes without anti-Xa values in the post-IM group. These episodes were not included in the analysis.

Anti-Xa status in the 24 first hours of anticoagulation

Within the first 24 hours of anticoagulation, 550 (67%) episodes reached the therapeutic range in the pregroup compared with 778 (67%) in the postgroup, with no statistically significant difference (p=0.82). There were 181 (22%) episodes with the infratherapeutic status in the pregroup and 241 (21%) in the postgroup (p=0.46). The 59 (7%) supratherapeutic episodes in the pregroup was significantly lower...
than the 136 (12%) in the postgroup (p<0.01). In particular, the 64 (5%) episodes with anti-Xa over 1 IU/mL in the postgroup was significantly higher than the 22 (3%) episodes in the pregroup (p<0.01). Finally, there were more episodes missing anti-Xa in the 24 first hours of anticoagulation in the pregroup than in the postgroup (4% vs 1%) (table 3).

In the subgroup analysis, episodes reaching therapeutic range were 21 (75%) in the CPAT group compared with 308 (73%) in the post-IM group, with no statistical difference (OR 1.13, 95% CI: 0.45 to 3.23, p=1). For infratherapeutic episodes, 1 (4%) were found in the CPAT group and 52 (12%) in the post-IM group. Episodes with supratherapeutic status were 6 (21%) in the CPAT group and 60 (14%) in the post-IM group. Episodes with anti-Xa over 1 IU/mL were 2 (7%) in the CPAT group compared with 31 (7%) in the post-IM group. Finally, the episodes without any anti-Xa in the first 24 hours were 0 (0%) in the CPAT group and 4 (1%) in the post-IM group.

DISCUSSION

This study showed that after the dissemination of a pocket card of recommendations for heparin treatment initiation, monitoring, and dosage adaptation, a significant increase in the time to reach the first anti-Xa was found. Several factors may contribute to these results. First, our definition of therapeutic versus prophylactic anticoagulation was based on heparin dose, since it is not specified in the medical chart. Second, we only considered anticoagulation episodes with anti-Xa monitoring for this analysis, with a large proportion of missing data for anti-Xa values: at the time of the study, access to anti-Xa tests, especially during weekends and nights was limited, and heparin was still often monitored with aPTT rather than the anti-Xa assay. It is therefore possible that anti-Xa monitoring was preferentially used in the pregroup by doctors that were also already using the new recommendation and nomogram for heparin doses. The postgroup included these ‘good prescribers’ as well as other prescribers who may have continued prescribing without the recommended doses and titration. To test this hypothesis, we conducted the same analysis within the first 7 hours with aPTT instead of anti-Xa tests and found a decrease of the missing values (18% in the pregroup and 15% in the postgroup); the number of episodes with aPTT in the therapeutic range was not significantly different between the two groups (12% in the pregroup vs 15% in the POST, p=0.13). A selection bias due to missing data is therefore highly possible. Future studies with defined therapeutic goals for anticoagulation and more reliable anti-Xa values are needed to verify our findings.

The CPAT in the subgroup analysis significantly decreased the interval between the initiation of UHF and the first anti-Xa in therapeutic range (7.1 hours [95% CI: 6.8 to 7.5 hours] in the CPAT subgroup vs 14 hours [95% CI: 13 to 15 hours] for the postgroup) and is also lower than in the pregroup (10.1 hours, 95% CI: 7 to 13 hours). This result could be a consequence of the regulating aspects of the integration of the recommendation in the prescription tool.

Since prescriptions are computerised, the CPAT provides the right assistance at the right time for each prescription of heparin. This significant finding although based on a small sample size in the CPAT, is consistent with a previous study by Kershaw et al.17

Furthermore, use of a computerised prescription tool was more effective, with a decrease of the number of infratherapeutic episodes of anticoagulation, both within the first seven or 24 hours of therapy. The initial heparin dose is 400 IU/kg in the CPAT and pocket guide, and is only dependent on the patient’s weight. A possible collateral effect of this initial dose and higher efficacy may be seen in the increased number of supratherapeutic episodes with CPAT, with the subsequent risk of bleeding with higher anti-Xa values.18 A further study with a larger data set of CPAT use is being conducted to verify these findings, and to determine if adjustments are needed for the initial heparin dose.

Regarding the anti-Xa status, the analysis in the first 7 hours used was as a surrogate to assess the efficiency of the anticoagulation initiation: again, the missing data need to be considered to interpret the lower proportion of results in the therapeutic range in the postgroup compared with the pregroup. The infratherapeutic status were not different between the two groups, but the postgroup had a higher number of supratherapeutic anti-Xa and especially anti-Xa above 1 IU/mL. This may be due to several reasons. The missing values of anti-Xa (where perhaps aPTT testing was used instead) could also be a source of bias. We did not explore aPTT values in this analysis because of its lower reliability of heparin effect.18

The identification of anticoagulation episodes with a therapeutic aim may have been suboptimal: a bias may have been induced when we removed certain specialties (neurology, haematology, etc.), where dosing does not follow recommendations because of a higher risk of bleeding complications. The higher supratherapeutic values in the postgroup suggest may also be due to the initial recommended dose, which may be higher than the doses used by clinicians for certain populations (eg, elderly patients).

When looking at the anti-Xa status in the 24 first hours of anticoagulation, there is no difference between groups for anti-Xa in the therapeutic range and an increased proportion of supratherapeutic anti-Xa in the postgroup.

The main limitations of this study are its retrospective and quasi-experimental design, and the possible inherent bias. Other limitations arise from the clinical documentation: the time of UFH administration and of blood samples were approximated from times registered in the EMR, thus less accurate. When blood samples were analysed outside of the hospital (eg, for baseline data), they could not be integrated to the analysis. Information concerning the occurrence of bleeding could not be obtained and were thus lacking in the analysis. Indications were inferred from ICD codes entered in the system after the hospital stay and thus could not be assessed with certainty. Moreover, their acute or chronic nature were not known. There was also a bias induced by the exclusion of the neurology, neurosurgery and haematology wards, because they represent a significant portion of the therapeutically anticoagulated patients.
Finally, the therapeutic intent of UFH treatment is not clearly identified in the EMR, and our identification method may have not excluded all the prophylactic events in the analysis.

One of the strengths of this study is the size of the sample. Moreover, the quasi-experiment nature of this study also has the upside of offering real-life results of the intervention. Indeed, there is no one controlling that physician strictly observe the pocket card recommendation usually, and a more controlled design would have more accurately extracted the efficiency of this pocket card use but might have overestimated its impact.

CONCLUSION

In conclusion, the results of our study showed that dissemi- nating a pocket card of recommendations did not reduce the time to reach therapeutic anti-Xa range, nor did it decrease the number of infratherapeutic events. This result should be verified in a separate, more specific study with less missing anti-Xa data, as it might change the initial dosage of UFH used in our protocol. The implementation of a computerised UFH prescription tool showed potential in bolstering the efficiency of the pocket card recommendation, with a significant decrease in the time to therapeutic range. This study reinforces the concept of providing the right tool (computerised tool) at the right time (prescription entry) to optimise the impact of the intervention. A further study with a larger sample of CPAT is needed to confirm the findings of this study.

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Contributors KSB (lead investigator and guarantor) conceived the study, obtained funding, developed the analysis plan and supervised the study analysis. PF and JS contributed to the study design and analysis plan. WEJ contributed to the study design, did the statistical analysis and wrote the first draft of the manuscript. All authors critically revised and approved the final manuscript. The corresponding author attests that all listed authors meet authorship criteria and that no others meeting the criteria have been omitted.

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REFERENCES

This UFH pocket card was translated solely as a reference for this publication (original in French)

**Monitoring**

Check anti-Xa HNF activity:
- 6h (at least 4h) after initial dose
- 6h after each change of dose, or at least 1x/day
- Check platelet levels before starting UFH, then 2-3x/week for 2 weeks

<table>
<thead>
<tr>
<th>If anti-Xa level &gt;1.0 IU/ml or &lt; 0.20 IU/ml, check:</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Where blood was drawn: same side as UFH? On the UFH catheter?</td>
</tr>
<tr>
<td>• Any other IV meds on the same catheter as UFH?</td>
</tr>
<tr>
<td>• Is the syringe pump working? Tubing unobstructed?</td>
</tr>
</tbody>
</table>

**Therapeutic anticoagulation with IV UFH**

**Overview**

- Prescribe in IU/24h, administer with a syringe pump
- Only use heparin vials (20’000 IU/48 ml)
- CBC, Quick and a PTT should be drawn before starting UFH perfusion

**Administering UFH**

- Initial dose: 400 IU/kg/24h in syringe pump (max 40’000 IU/24h)
- In the case of an acute thrombotic event, consider an initial bolus of 50 IU/kg (max 5000 IU)

**No bolus if:**
- Recent surgery (< 48-72hrs)
- Thrombocytopenia < 50 G/l
- Thrombolysis
- Other causes for increased bleeding
# Therapeutic anticoagulation with IV UFH

## Titration dose protocol (Smith ML et al, Am J Health Syst Pharm 2010)

<table>
<thead>
<tr>
<th>Anti-Xa HNF activity (UI/ml)</th>
<th>Bolus if prescribed</th>
<th>Other weights</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>If 55-64 kg</td>
<td>If 65-74 kg:</td>
</tr>
<tr>
<td>&lt; 0.20</td>
<td>26 UI/kg*</td>
<td>🔺 by 96 IU/kg/24hr</td>
</tr>
<tr>
<td>0.20-0.29</td>
<td>Non</td>
<td>🔺 by 48 IU/kg/24hr</td>
</tr>
<tr>
<td>0.30-0.70</td>
<td>Non</td>
<td>-</td>
</tr>
<tr>
<td>0.71-0.80</td>
<td>Non</td>
<td>🔺 by 24 IU/kg/24hr</td>
</tr>
<tr>
<td>0.81-0.99</td>
<td>Non</td>
<td>🔺 by 48 IU/kg/24hr</td>
</tr>
<tr>
<td>≥1.00</td>
<td>Non</td>
<td>STOP</td>
</tr>
</tbody>
</table>

*Except if major risk of bleeding **Check how the blood was drawn

If you have any questions, call the hemostasis consultation (XX XXXXX)
Check anti-Xa HNF activity 6hrs after any change in dose.