Low-molecular weight heparin infusion as anticoagulation for haemodialysis

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Abstract

Background: Low-molecular weight heparin (LMWH) is commonly used as an anticoagulant for haemodialysis by a single-bolus injection. However, its application in extended haemodialysis has been infrequently studied. In particular, for nocturnal home haemodialysis patients sleeping throughout treatment, the need for additional intradialytic bolus might render the use of LMWH impractical. To overcome this limitation, we changed traditional bolus injections to continuous infusion. We first tested our method among in-centre 4-h haemodialysis patients to establish a feasible and safe infusion regimen before utilizing it in extended dialyses at home.

Methods: Recruited patients were given nadroparin (standardized at 65 IU/kg) as an anticoagulant for haemodialysis. They were first randomized to receive nadroparin either by bolus injection or infusion. Afterwards, the patients underwent crossover to receive the alternate method of LMWH anticoagulation. The degrees of anticoagulation and bleeding complications were compared.

Results: Sixteen haemodialysis patients were recruited. After nadroparin administration, anti-Xa levels at the first hour were significantly higher by the bolus than the infusion methods (0.68 ± 0.10 versus 0.49 ± 0.10 IU/mL, P < 0.001) and were similar by the second hour (0.56 ± 0.10 versus 0.55 ± 0.11 IU/mL, P = 0.64). At the sixth hour, anti-Xa levels by the infusion method were significantly higher (0.35 ± 0.13 versus 0.25 ± 0.10 IU/mL, P < 0.001), suggesting the infusion approach required a dosage reduction. There were no bleeding events reported in either method.

Conclusions: LMWH infusion is feasible and safe. The method avoids early excessive anticoagulation caused by bolus injection and reduces the LMWH dose. Future studies should be conducted to evaluate LMWH infusion in extended haemodialysis treatment.

Key words: anticoagulation, continuous infusion, haemodialysis, low-molecular weight heparin, nadroparin

Introduction

Anticoagulation is an essential element of haemodialysis therapy for patients with end-stage renal disease (ESRD). An extracorporeal circuit is a prerequisite for the delivery of adequate dialysis. In a majority of patients with low bleeding risk, systemic heparin is utilized for anticoagulation. Given its better
bioavailability at a lower dose, longer half-life and more predictable anticoagulant responses, low-molecular weight heparin (LMWH) has been increasingly preferred over unfractionated heparin (UFH) in conventional intermittent haemodialysis [1]. Moreover, potential superior effects on lipid profiles, reduced risks for osteoporosis [2–4] and lower risks of heparin-induced thrombocytopenia (HIT) [5] favour LMWH over UFH.

LMWH is typically given as a single intravenous bolus into the arterial limb when starting dialysis. While such drug administration convenience is advantageous, the single bolus injection limits the efficacy of LMWH in haemodialysis to conventional 4-h treatments. In extended haemodialysis lasting 6–8 h, additional intradialytic boluses of LMWH are often required to ensure circuit patency [6]. However, this approach may not be practical for all patients, especially for those receiving nocturnal haemodialysis at home when they are sleeping during treatment. As a result, UFH by continuous infusion, which requires no additional intradialytic manoeuvres, is more commonly employed in this group of patients [7].

LMWH administration by continuous infusion has been less studied in the literature. Enoxaparin and nadroparin infusion have been evaluated for continuous haemofiltration in critical care settings [8, 9]. For intermittent haemodialysis, while an infusion regimen has been recommended for dalteparin [1], there is a paucity of data on LMWH infusion, let alone direct comparisons against the traditional bolus method. As a result, we performed a pilot study to first evaluate the feasibility and safety of LMWH infusion as an anticoagulant for conventional intermittent haemodialysis, to construct an appropriate infusion regimen for future use in extended haemodialysis.

Materials and methods

All patients ≥18 years old with ESRD receiving thrice-weekly 4-h haemodialysis at our institution were screened. Patients who had an underlying bleeding disorder, a history of intolerance to LMWH, had been receiving oral anticoagulants or other drugs that could affect heparin activity (e.g. tetracycline, digitalis, and antihistamines, etc.), or were unable to give informed consent were excluded from our study. Nadroparin (Fraxiparine, GlaxoSmithKline) was thoroughly mixed with normal saline into a 20-mL syringe before use. Such a dilution allowed more precise individual dosing of nadroparin, standardized at 65 IU/kg. For the bolus method, nadroparin was injected into the arterial limb of a haemodialysis circuit. For the infusion method, a loading dose at 35 IU/kg was given, followed by 10 IU/kg infusion per hour for 3 h (i.e. stopping 1 h before the end of dialysis). The study protocol was approved by the Ethics Committee of the hospital, and we fully adhered to the principles of Good Clinical Practice and Declaration of Helsinki. Written informed consent was obtained from all participants in the study.

Each recruited patient underwent a 4-week haemodialysis treatment at our institution. The first week was a washout period using UFH as the anticoagulant. In the second week, the patients were randomized to receive nadroparin for dialysis by either bolus or infusion method in a 1:1 manner. The third week was another washout period with UFH. In the fourth week, the patients underwent crossover to receive the alternate method of LMWH anticoagulation (Figure 1). All haemodialyses were delivered by Fresenius 4008 devices, using synthetic hollow fibre dialysers (polysulphone membrane FX80M capillary middle-flux dialysers; Fresenius Medical Care, Bad Homburg, Germany). A new dialyser was utilized for each dialysis treatment and was pre-rinsed with 1000-mL normal saline. Bicarbonate haemodialysate was used. The dialysate flow rate was maintained at 500 mL/min and the blood flow was kept between 200 and 300 mL/min depending on vascular access conditions of the patients. Ultrafiltration was performed as clinically indicated.

The evaluations of each patient, including blood sampling and thrombus assessment, were carried out during mid-week haemodialysis treatments for both methods (i.e. during the second session, in the second and fourth weeks of the study). Degrees of thrombus formation in the dialyser and arterial and venous air traps were assessed by two dialysis nurses, one of whom was blinded to the study. We employed a 5-grade scale: Grade 0, no detectable clot; Grade 1, minimal clot; Grade 2, moderate clot; Grade 3, major clot formation but dialysis still feasible; Grade 4, complete occlusion by thrombus rendering dialysis impossible. Compressions of the arterial and venous cannulation sites post-dialysis were performed sequentially by the patients themselves. Total haemostasis times required were noted, and reported as the summation of haemostasis times required for both cannulation sites. Any bleeding events during the study were also recorded.

For blood sampling, pre-dialysis blood collection was performed before nadroparin administration (Time 0), which included complete blood picture, prothrombin time (PT), activated thromboplastin time (APTT) and measurements of anti-Xa, urea and creatinine levels. Immediate post-dialysis urea and creatinine levels were also collected for the calculation of single-pool Kt/V (spKt/V) to evaluate dialysis clearance [10]. At the first, second and sixth hours after the administration of nadroparin, blood samplings for PT, APTT and anti-Xa were also performed. All blood collections during dialysis were performed in accordance with the Kidney Disease Outcomes Quality Initiative (KDOQI) guidelines [11].

Anti-Xa assay

Doubly spun (3750 rpm × 10 min) platelet-poor plasma was prepared from citrated peripheral blood. Platelet counts were verified to be <10 × 10⁹/L. Plasma samples were frozen at −70°C before testing. The plasma level of LMWH was measured by anti-Xa activity with an amidolytic method using a Stachrom Heparin Kit (Diagnostica Stago, France) on a CA-7000 coagulometer (Sysmex, Japan). Testing procedures followed the instructions of the manufacturer, except a 5-point calibration curve was used for a more accurate quantitation instead of the 3-point calibration curve recommended by the manufacturer.

Statistical methods

Statistical evaluations were performed using the SPSS version 13.0 software package (SPSS, Chicago, IL, USA). Continuous variables were expressed as mean ± standard deviation or median (interquartile range), whereas categorical variables were expressed as proportions. Comparisons of data were made by paired Student’s t-test, Wilcoxon signed-rank test or Fisher’s exact test as appropriate. Analysis of variance (ANOVA) with Bonferroni post hoc test was used if three or more groups of data were involved for a comparison. Two-sided P-values were reported, with P-values <0.05 considered significant.

Results

A total of 16 patients were recruited and all completed our study. Their baseline characteristics are shown in Table 1. The mean age was 58.9 years, with a median haemodialysis vintage of 3.6 years.
Most patients underwent dialysis via native arteriovenous fistulae. Individual nadroparin dosages were prescribed in accordance with body weight at 3928.5 ± 422.2 IU. All patients underwent thrice-weekly 4-h haemodialysis treatments uneventfully during our study. During LMWH administration via the bolus method, two patients terminated dialysis 16 and 8 min early, whereas one patient terminated 5 min early during administration via infusion. These early termination cases were due to significant muscle cramps. None of the patients encountered premature termination of dialysis due to clotting of the extracorporeal circuit.

The baseline anti-Xa levels of recruited patients upon study entry were confirmed to be undetectable. The comparisons of various parameters between the bolus and infusion methods are shown in Table 2. At Time 0, PT was slightly prolonged during the infusion method, likely with little clinical relevance (10.68 ± 0.52 versus 10.45 ± 0.50, P = 0.005). Other parameters at Time 0 were similar between the two methods, including negligible anti-Xa levels. After nadroparin administration at the start of dialysis, a typical early peak was avoided with the infusion method, as shown by the lower anti-Xa level at the first hour (0.49 ± 0.10 versus 0.68 ± 0.10 IU/mL, P < 0.001). At the second hour, the anti-Xa level in the infusion method was similar to that in the bolus approach (0.55 ± 0.11 versus 0.56 ± 0.10 IU/mL).

Table 1. Baseline demographics of patients (n = 16)

| Parameter                                | Value                  |
|------------------------------------------|------------------------|
| Age, years                               | 58.9 ± 7.6             |
| Male gender, n (%)                       | 8 (50)                 |
| Body mass index, kg/m²                   | 24.2 ± 2.0             |
| Aetiology of ESRD, n (%)                 | 2 (12.5)               |
| Hypertension                             | 3 (18.8)               |
| Glomerulonephritis                       | 2 (12.5)               |
| Polycystic kidney disease                | 4 (25)                 |
| Others                                   | 5 (31.2)               |
| Comorbidities, n (%)                     | 15 (93.8)              |
| Hypertension                             | 4 (25)                 |
| Diabetes mellitus                        | 3 (18.8)               |
| Ischemic heart disease                   | 1 (6.3)                |
| Cerebrovascular disease                  | 5 (31.3)               |
| History of malignancy                    | 3.6 (2.6–7.3)          |
| Hemodialysis vintage, years              |                        |
| Dialysis access, n (%)                   | 14 (87.5)              |
| Synthetic graft                           | 2 (12.5)               |

Continuous variables are expressed as mean ± standard deviation or median (interquartile range). ESRD, end-stage renal disease.

Fig. 1. Study flowchart. UFH, unfractionated heparin.
Our study provides the first clinical data on the safety and feasibility of LMWH infusions compared with bolus injections for haemodialysis. Conventionally, LMWH action has been assessed based on anti-Xa activity, for which 0.5 IU/mL has been recommended as the target level [12]. An ideal anticoagulant should be safe to use, without causing excessive anticoagulation leading to potential haemorrhage. For LMWH, one drawback of the traditional method is an unavoidable overanticoagulation effect early after injection. Within the initial 2 h, high anti-Xa levels of >1.0 IU/mL were reported after the administration of enoxaparin, nadroparin or tinzaparin [13–15]. These results have been summarized in a meta-analysis of randomized trials on LMWH [16]. In our study, similar early peaks of the anticoagulant effect were observed among patients treated with the bolus method, with a mean anti-Xa level of ~0.7 IU/mL during the first hour. For the infusion method, sudden spikes in LMWH action were avoided, and the anti-Xa levels were stably maintained at 0.5 IU/mL during the first and second hours. This highly suggested that our proposed infusion approach could be a safer option within the first 2 h of dialysis. Given that overanticoagulation is uncommon beyond 2 h of LMWH administration in previous trials [16], bleeding risks related to our infusion approach are probably of less concern despite not recording anti-Xa levels at the third and fourth hours. Additionally, the uneventful completion of dialysis treatment and the similar dialytic clearances and thrombus scores observed for the infusion method compared with those in the bolus method supported the feasibility of our proposed infusion regimen to maintain extracorporeal circuit patency during haemodialysis.

Higher levels of anti-Xa activity in the infusion group were observed at the sixth hour (i.e. 2 h post-dialysis) when identical dosages of nadroparin were utilized for both methods. This may indicate higher post-dialytic haemorrhagic risk by the infusion method due to a more residual anticoagulant effect. Such a drawback could be easily resolved by reducing the LMWH dosage, such as a reduction in the hourly infusion rate or an earlier cessation of infusion, to alleviate post-dialysis anticoagulant activity. Further clinical study is necessary to validate our postulates.

Discussion

Our study provides the first clinical data on the safety and feasibility of LMWH infusions compared with bolus injections for
and to define optimal infusion regimens. Blood sampling at mid-week haemodialysis in our study also allowed us to identify any potential LMWH accumulation with the infusion method. Anti-Xa levels at Time 0 (i.e. 48 h after nadroparin administration in preceding haemodialysis treatment) were negligible, implying no significant LMWH accumulation with the infusion method for conventional thrice-weekly haemodialysis.

As mentioned earlier, one major aim of our proposed LMWH infusion method was for application in dialysis beyond conventional 4-h treatments, especially for nocturnal home haemodialysis patients. Our favourable results show that the infusion approach could be a safe and feasible option for LMWH administration in the literature, studies on anticoagulation in extended haemodialysis therapy have been relatively scarce. LMWH administration has been limited to the bolus method and has principally been applied to nocturnal dialysis delivered in-centre only [5, 17]. The need for intradialytic boluses of LMWH prohibits its use for patients on nocturnal haemodialysis at home. The potential superior effects on lipid and bone metabolism and lower risk of HIT may make LMWH a preferred choice over UFH. The latter is particularly relevant because the development of HIT would render nocturnal home haemodialysis problematic, if not impossible [18]. Certainly, an extrapolation of our results into extended haemodialysis treatment requires additional clinical trials that apply infusion approaches of LMWH to this particular group of patients.

There were several limitations in our study. The sample size was relatively small, and we lacked more data on intradialytic anti-Xa levels. Therefore, our study was inevitably underpowered to report potential significant differences between the two anticoagulation methods. Additionally, the extrapolation of our results to other ethnic groups may not be necessarily applicable because all recruited subjects were Chinese. Furthermore, the markers of coagulation activation, such as thrombin-antithrombin complex and prothrombin fragments 1 and 2, were not monitored. This restricted our circuit patency evaluation only to any visible clots. However, with dialysis prescriptions kept separate from the anticoagulation method, the evaluation only to any visible clots. However, with dialysis pre-2, were not monitored. This restricted our circuit patency

of HIT may make LMWH a preferred choice over UFH. The latter is relatively small, and we lacked more data on intradialytic anti-Xa levels. Therefore, our study was inevitably underpowered to report potential significant differences between the two anticoagulation methods. Additionally, the extrapolation of our results into extended haemodialysis treatment requires additional clinical trials that apply infusion approaches of LMWH to this particular group of patients.

To conclude, we have modified the administration method of LMWH for anticoagulation in intermittent haemodialysis. Compared with the traditional single-bolus approach, our infusion regimen was found to be equally safe and feasible, with additional merits of avoiding excessive anticoagulation early after bolus injection and possible LMWH dose reduction. Further clinical trials are needed to determine whether the LMWH infusion method could be applied in extended haemodialysis treatment.

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Conflict of interest statement
There are no conflicts of interest to declare for all authors.

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