Continuous Infusion of Factor VIII and von Willebrand Factor in Surgery: Trials with pdFVIII LFB or pdVWF LFB in Patients with Bleeding Disorders

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Abstract

Background A plasma-derived factor VIII product (pdFVIII; Factane 100 or 200 IU/mL) and a plasma-derived von Willebrand factor product (pdVWF; Wilfactin 100 IU/mL) are approved for replacement therapy by intravenous bolus injections in hemophilia A (HA) and von Willebrand disease (VWD), respectively. However, in situations requiring intensive treatment, continuous infusion (CI) may be desirable to better control target plasma factor levels.

Aim To evaluate the perioperative hemostatic efficacy and safety of these concentrates administered by CI.

Methods Three phase III trials were conducted. Adults with HA (FVIII:C < 1%) (studies 1 and 2) or VWD (VWF:RCo < 20%) (Study 3) received a preoperative bolus followed by CI of undiluted concentrate for at least 6 days. Bolus doses and CI rates were based on individual recovery and clearance, respectively. The initial infusion rate had to be higher for 48 hours for HA and 24 hours for VWD patients to anticipate potential fluctuations of factor concentrations during major surgery. Target levels of FVIII:C in HA and VWF:RCo in VWD were 80 and 70 IU/dL, respectively. Efficacy was assessed using a global hemostatic efficacy score.

Results Studies 1, 2, and 3 included 12, 4, and 6 patients, respectively. Efficacy outcomes were excellent/good in all 22 major surgeries including 18 orthopedic procedures. Most daily measured FVIII and VWF levels (92%) were on target. No safety concerns, thrombotic events, or inhibitors were identified.

Conclusion pdFVIII and pdVWF administered by CI represent an effective and safe alternative to bolus injections in patients with severe HA or VWD undergoing surgery.
Continuous Infusion of Coagulation Factor Products  Windyga et al.  1305

Introduction

Hemophilia A (HA) and von Willebrand disease (VWD) could be associated with excessive bleeding during major surgeries especially during orthopedic surgeries. As in HA, a low factor VIII (FVIII) level is an important determinant for joint bleeding in VWD. Arthropathy caused by recurrent joint bleeding episodes therefore occurs in both diseases. Van Galen et al presented outcomes after joint bleeds in VWD and HA and showed that orthopedic surgical procedures are required in approximately one in five patients in both diseases. Perioperative prophylactic treatment with FVIII or von Willebrand factor (VWF) is required in these patients to correct the underlying coagulation abnormalities and minimize the bleeding risk. Replacement therapy of the lacking protein is classically administered by a bolus injection regimen consisting of an initial dose followed by subsequent intermittent administration to maintain adequate factor levels. This type of regimen may not be an optimal approach to manage hemostasis since it may be associated with low trough levels and therefore an increased risk of harmful bleeding. Replacement of blood coagulation factors by continuous infusion (CI) was previously reported to represent an appropriate option to prevent large variations of coagulation factor levels in the bloodstream, either low trough levels or unnecessary high peaks. Advantages are likely the improvement of the protection from excessive bleeding, and easier monitoring. Efficacy and safety of CI remain deeply investigated for management of patients with severe HA and VWD undergoing major surgery. Causes influencing CI rates are the product, its adsorption in the infusion system, its dilution, the type of surgery, and the inter-patient pharmacokinetic (PK) variability. This article aims to present the results from three phase III clinical trials assessing CI as a modality of administration within the surgery indication of Factane and Wilfactin/Willfact (LFB, Les Ulis, France) in patients with HA and VWD, respectively. Factane is a human plasma-derived FVIII (pdFVIII) product and Wilfactin/Willfact, a human plasma-derived VWF (pdVWF) almost devoid of FVIII.

Methods

pdFVIII and pdVWF Products

The therapeutic proteins are extracted from plasma using plasma fractionation techniques. Both products are purified by chromatography, treated by solvent/detergent to inactivate enveloped viruses, nanofiltered to eliminate nonenveloped viruses (filters with pore sizes of 35 and 15 nm for the FVIII concentrate and 35 nm for the VWF concentrate). A dry-heat treatment at 80°C for 72 hours is added for the VWF concentrate as a third dedicated viral removal/inactivation step. Each product administered by bolus injection was successfully used in the perioperative prevention and treatment of bleeding during surgery in adults with severe disease.

Compatibility Studies

The compatibility of products with the CI equipment (pdFVIII 100 IU/mL and pdFVIII 200 IU/mL recently manufactured with a smaller reconstitution volume, and pdVWF) was based upon assays either performed with a syringe pump or with an ambulatory infusion pump including bag and tubing sets. These compatibility studies were performed using standardized protocols by the manufacturer. In vitro stability studies were also evaluated under real conditions within hospital organization to guarantee the pharmaceutical quality of the preparations. Details of the protocols are provided in the Supplementary Material (available in the online version).

Study Designs

Three prospective, open-label, phase III studies were conducted between 2005 and 2013 in accordance with Good Clinical Practice and ethical principles outlined in the Declaration of Helsinki. The use of CI of pdFVIII concentrated at 100 IU/mL (Study 1) and at a higher concentration of 200 IU/mL (Study 2) was investigated at a single center in Poland in patients with HA. Then, to extend the knowledge of this mode of administration in a context of VWD, Study 3 was initiated with pdVWF in France and Poland. Clinical investigations for CI adhered to current guidelines in force for FVIII products or were adapted for VWF products. Ethics and national authorities’ approvals, and written informed consent were obtained prior to enrolment. Studies were registered in EudraCT (numbers 2010–023666–46 for the most recent study with pdFVIII and 2007–004116–32 for pdVWF). The studies consisted of (1) a presurgery PK evaluation, (2) a surgical period with factor replacement by CI for at least 6 days (144 hours), and (3) a safety observation period following the CI.

Patient Population

Enrolled patients were adults who had to undergo elective major surgical procedures requiring at least 6 days of treatment. In Study 1 and Study 2, patients with severe HA (baseline FVIII coagulant activity [FVIII:C] levels <1 IU/dL) could be enrolled if they had ≥150 exposure days to FVIII. In Study 3, patients with VWD should have inherited severe VWF deficiency (type 1, 2, or 3) with baseline VWF ristocetin cofactor (VWF:RCO) levels <20 IU/dL. In all cases, present or past inhibitor against FVIII or VWF was also an exclusion criterion. Patients should not be included in case of problematic venous access.

Preoperative Pharmacokinetic Study

Each patient underwent a presurgery PK evaluation to determine their individual incremental recovery and the estimate of clearance for calculations of the loading dose and the initial infusion rate, respectively. PK endpoints were assessed after a single dose of 50 IU/kg of pdFVIII in HA patients or 60 IU/kg of pdVWF in VWD patients. Blood samples were withdrawn over a period of 48 hours (Studies 1 and 2) or 9 hours (Study 3) as detailed in the Supplementary Material (available in the online version).

Continuous Infusion Protocol and Maintenance of Factors Levels

A preoperative loading dose of factor concentrate was administered by bolus injection on the day of the surgical
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procedure (D0). This bolus dose was calculated to raise a peak plasma levels between 80 and 120 IU/dl of FVIII:C for HA and at least 100 IU/dl of VWF:RCo for VWD patients according to the formula: dose (IU) = body weight (kg) x desired increment in factor level (IU/dl)/individual incremental recovery at 15 minutes (IU/dl per IU/kg). CI was initiated immediately at the end of the bolus.

In the postoperative period, the target factor levels of FVIII:C and VWF:RCo were respectively between 80 and 120 IU/dl for HA and at least 70 IU/dl for VWD patients for the first week, and thereafter between 30 and 80 IU/dl for HA and at least 50 IU/dl for VWD patients. To achieve these levels, the infusion rate was based on the formula: rate of infusion (IU/kg/h) = individual clearance (ml/h/kg) x desired factor level (IU/ml). A multiplying coefficient was applied to baseline clearance during the first 48 hours in studies 1 and 2 (2.5 for H0–H12, 2 for H12–H24, and 1.5 for H24–H48) and during the first 24 hours (from 2.0 to 1.5) in Study 3, to take into account an anticipated increased factor consumption due to postsurgical hemodynamic disturbances caused by blood loss and the nature and volume of expansion fluids infused.22 Circulating factor levels were measured at 6, 12, and 24 hours and then every 24 hours until 144 hours after the start of the CI. If needed, additional bolus doses could be also administered to reach quickly the desired level. All steps relative to the CI protocol for a specific patient are detailed in the Supplementary Material (available in the online version).

The lyophilized product was reconstituted with the provided sterile water once a day without further dilution. Products were filled under laminar air flow conditions in the pump reservoir which was either a syringe or an infusion pump. Minor amounts of unfractionated heparin previously diluted in saline (5 IU/mL) were added to pdFVIII before the transfer in the delivery system to prevent local thrombophlebitis.32 Circulating factor levels were measured at 6, 12, and 24 hours and then every 24 hours until 144 hours after the start of the CI. If needed, additional bolus doses could be also administered to reach quickly the desired level. All steps relative to the CI protocol for a specific patient are detailed in the Supplementary Material (available in the online version).

Because some VWD patients have a secondary FVIII deficiency, a bolus dose of 60 IU/kg of pdVWF was administered 12 to 24 hours before surgery to start the increase of endogenous FVIII and correct the FVIII:C level at the time of surgery. The target FVIII:C levels were ≥50 IU/dl before surgery and thereafter.

Efficacy Endpoints

The primary outcome was the global hemostatic efficacy assessment score merging intraoperative and postoperative assessments: on day 0 by the surgeon and on day 7 by the investigator. An additional assessment on the day of drain tube removal (or day 2 if no drain) was performed by the surgeon in studies 1 and 2.

Each assessment was scored on a 4-point scale (excellent, good, moderate, none). The treatment was considered as a “success” if the patient obtained an excellent or good score in the successive hemostasis assessments. Prior to each surgical procedure, the surgeon provided an estimate of the average and maximum expected blood loss during surgery for a person with a normal hemostasis under the same conditions, allowing comparison of actual to predicted blood loss. Other endpoints included the occurrence of bleeding complications, packed red blood cell units transfused, requirement of second surgical intervention, factor plasma levels, and product consumption (dosage, infusion rates, and number of additional bolus injections).

Safety Endpoints

Safety was evaluated through reports of adverse events analyzed according to their relationship with either the study drug or the method of administration. Special attention was given to possible thromboembolic events and allergic/anaphylactic-type reactions. A daily examination of infusion site for signs of redness, pain, or swelling was performed. Vital signs, physical examination, and laboratory parameters including hematology and blood chemistry were monitored at study visits. Inhibitors to FVIII:C or to VWF:RCo were measured regularly during the study in the central laboratory by assays based on the Nijmegen modification of the Bethesda assay.34

Statistical Methods

Exposure, efficacy, and safety parameters were summarized using descriptive statistics. The number and percentage together with the exact two-sided 95% Clopper–Pearson confidence interval were calculated for the primary endpoint. PK variables were reported as descriptive statistics.

Results

Compatibility Studies

Compatibility studies showed that either FVIII (Supplementary Fig. S1 [available in the online version]) or VWF (Supplementary Figs. S2 and S3 [available in the online version]) retain at least 90% of baseline potency for at least 24 hours whatever the tested conditions. Both reconstituted products were stable for 3 days in ambulatory pumps. FVIII activity was not impacted by the presence of heparin, composition of container, product potency, or low rate of infusion. There was no evolution in the multimeric composition (≥10-mers and ≥15-mers) for pdVWF product (Supplementary Fig. S4 [available in the online version]). No microbial contamination was observed.

Demographics and Baseline Characteristics

A total of 12, 4, and 6 patients were included in studies 1, 2, and 3, respectively. Baseline characteristics of the patients are summarized in Table 1. Age ranged from 28 to 75 years, and weight from 49 to 100 kg body weight. Basal FVIII:C levels in the VWD patients ranged from 2 to 18 IU/dL.

Preoperative Pharmacokinetic Study

All patients completed a preoperative PK study. The FVIII concentration–time profiles of pdFVIII 100 IU/mL (Study 1) and pdFVIII 200 IU/mL (Study 2) were similar (Supplementary Fig. S5 [available in the online version]). An abbreviated curve of VWF plasma decay over 9 hours after injection of 60 IU/kg of pdVWF yielded informative data in...
### Table 1 Demographics and baseline characteristics of treated patients

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Study 1</th>
<th>Study 2</th>
<th>Study 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemophilia A, N</td>
<td>12</td>
<td>4</td>
<td>–</td>
</tr>
<tr>
<td>Von Willebrand disease, N</td>
<td>–</td>
<td>–</td>
<td>6</td>
</tr>
<tr>
<td>Type 1, N</td>
<td>–</td>
<td>–</td>
<td>2</td>
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<tr>
<td>Type 2, N</td>
<td>–</td>
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<td>0</td>
</tr>
<tr>
<td>Type 3, N</td>
<td>–</td>
<td>–</td>
<td>4</td>
</tr>
<tr>
<td>VWF:RCo (IU/dL) in Type 1</td>
<td>–</td>
<td>–</td>
<td>13 (11–15)</td>
</tr>
<tr>
<td>VWF:RCo (IU/dL) in Type 3</td>
<td>–</td>
<td>–</td>
<td>Undetectable</td>
</tr>
<tr>
<td>FVIII:C (IU/dL)</td>
<td>&lt;1 (&lt;1 to &lt;1)</td>
<td>&lt;1 (&lt;1 to &lt;1)</td>
<td>7.5 (2–18)</td>
</tr>
<tr>
<td>Sex (male/female)</td>
<td>12/0</td>
<td>4/0</td>
<td>3/3</td>
</tr>
<tr>
<td>Age (y)</td>
<td>37.0 (28–67)</td>
<td>39.0 (37–43)</td>
<td>49.5 (32–75)</td>
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<tr>
<td>Weight (kg)</td>
<td>64.0 (54.0–90.0)</td>
<td>82.5 (70.0–98.0)</td>
<td>80.4 (49.0–100.0)</td>
</tr>
<tr>
<td>Body mass index (kg/m²)</td>
<td>21.6 (18.7–27.8)</td>
<td>27.4 (24.5–29.1)</td>
<td>27.7 (21.2–32.3)</td>
</tr>
</tbody>
</table>

Abbreviations: FVIII:C, factor VIII coagulant activity; N, number of patients; VWF:RCo, von Willebrand factor ristocetin cofactor.

Note: Data are given as median (min–max).

the preoperative context ([Supplementary Fig. S6](available in the online version)]. From each study, the following were found: mean incremental recovery measured 15 minutes after the end of injection was: for pdFVIII 1.9 ± 0.4 and 2.3 ± 0.4 IU/dL per IU/kg in studies 1 and 2, respectively, and for pdVWF 2.2 ± 0.5 IU/dL per IU/kg (Study 3). FVIII clearance was 2.42 ± 0.6 and 2.40 ± 1.2 mL/h/kg in studies 1 and 2, respectively. In Study 3, estimate of VWF clearance was comparable regardless of the method used, noncompartmental or compartmental analysis, with mean values of 4.7 ± 1.2 and 5.0 ± 1.1 mL/h/kg, respectively. Other PK parameters are shown in [Supplementary Table S1](pdFVIII) and [Supplementary Table S2](pdVWF) (available in the online version). A large inter-individual variability was evidenced considering the coefficient of variation of approximately 20%.

### Efficacy

A total of 22 major elective surgical procedures were performed. In line with the total absence or severe deficiency of FVIII, most (18 of 22) were orthopedic procedures, including 4 hip or knee prosthesis revisions with high risk of bleeding complications. The success rate of the treatment was 100% as all assessments were rated as excellent or good, either at the early stage of CI by the surgeon or later by the hematologist ([Supplementary Table S3](available in the online version)]. Details for each surgical procedure are presented in [Table 2]. Perioperative blood loss was within predicted ranges in all except in two patients who underwent a total knee replacement. For the first patient with HA (patient 14), the investigator reported that the joint was more destroyed and the surgery more complicated than initially expected, and thus the predicted intraoperative blood loss volume had been underestimated. For the second patient with VWD (patient 20), a tourniquet was used during the intervention and excessive bleeding occurred in the immediate postoperative period; the investigator considered that the actual volume of blood was in line with the individual surgical conditions. For these two patients, bleeding occurred at times when factor levels were within hemostatic ranges (FVIII:C at 126 IU/dL and VWF:RCo at 120 IU/dL). Both patients did not require transfusion of packed cells or reoperation to control bleeding. Ten patients experienced blood loss of >1000 mL. Half of them received blood transfusions prescribed before surgery (1) or postoperatively (4).

The products were administered using individualized dosing including a safety margin dose to anticipate the factor consumption.

In HA patients, pdFVIII was given as a loading dose of mean (min–max) 65 IU/kg (48–86) and CI started with a rate of 6.8 IU/kg/h (4.5–8.9) in Study 1 while a lower loading dose (47 IU/kg [37–53]) and infusion rate (5.6 IU/kg/h [3.2–10.7]) were given in Study 2 in accordance with PK data. Mean FVIII:C levels were comparable 6 hours later: 88.6 IU/dL (71–126) and 99.8 IU/dL (88–126) in studies 1 and 2, respectively. In VWD, all patients received a bolus dose of pdVWF (mean 56 IU/kg) the day prior to the intervention. As a consequence, FVIII:C levels varied between 66 and 133 IU/dL (mean 86.7 IU/dL) at the time of the procedure, and no additional FVIII concentrate was then used. Thereafter, the mean loading dose of pdVWF was 40 IU/kg (28–56), the initial infusion rate of CI was 5.8 IU/kg/h (4.8–6.9), and the VWF:RCo level 6 hours later was 127 IU/dL (98–152).

The infusion rates administered that take into account the decreasing planned multiplying coefficient during the first 2 days and the calculated clearance is shown in [Table 3]. The clearance of FVIII:C or VWF:RCo decreased overtime ([Fig. 1]). It was evidenced that the factor consumption decreased over the course of 3 days in HA and 2 days in VWD patients. After 48 hours, the subsequent infusion rates were between 2.2 and 2.7 IU/kg/h either for pdFVIII or pdVWF products. Median (min–max) consumption per surgery was 608 IU/kg (376–1074) for pdFVIII and 579 IU/kg (411–825) for pdVWF.

In HA patients ([Fig. 2A, B]), mean FVIII:C levels reached the desired level of 80 IU/dL throughout the 6 days of CI. In
### Table 2: Characteristics of surgeries and details of factor administration by continuous infusion and blood loss

<table>
<thead>
<tr>
<th>Patient</th>
<th>Disorder</th>
<th>Type of surgery</th>
<th>Surgery duration (h:min)</th>
<th>Individual incremental recovery (IU/dL)/ (IU/kg)</th>
<th>Loading bolus dose (IU/kg)</th>
<th>Continuous infusion period</th>
<th>Blood loss volume</th>
<th>RBC transfusion and time</th>
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<tr>
<td>1</td>
<td>HA</td>
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<td>1:55</td>
<td>2.0</td>
<td>64.4</td>
<td>2.74</td>
<td>8.1</td>
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<tr>
<td>2</td>
<td>HA</td>
<td>Inguinal hernia repair</td>
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<td>3</td>
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<td>1.63</td>
<td>5.4</td>
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<td>HA</td>
<td>Socket of hip arthroplasty revision</td>
<td>1:15</td>
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<td>2.96</td>
<td>8.9</td>
<td>3.3</td>
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<td>HA</td>
<td>Total knee arthroplasty revisionb</td>
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<td>HA</td>
<td>Bilateral total knee replacementb</td>
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<td>2.5</td>
<td>54.4</td>
<td>1.63</td>
<td>4.5</td>
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<td>HA</td>
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<td>HA</td>
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<td>Total knee replacementb</td>
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<td>73.4</td>
<td>2.83</td>
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<td>3.1</td>
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<td>1.7</td>
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<td>3.20</td>
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<td>10.7</td>
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<td>Total knee arthroplasty revisionb</td>
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<td>1.9</td>
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<td>2.0</td>
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<td>2.2</td>
<td>43.3</td>
<td>1.9</td>
<td>4.4</td>
<td>2.2</td>
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<td>Type 3 VWD</td>
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<td>56.0</td>
<td>4.7^e</td>
<td>6.6</td>
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<td>36.8</td>
<td>3.5^e</td>
<td>4.9</td>
<td>2.5</td>
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<td>Total knee replacementb</td>
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<td>2.2</td>
<td>45.4</td>
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<td>6.9</td>
<td>2.5</td>
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<td>Type 1 VWD</td>
<td>Hemicolectomy</td>
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<td>5.4</td>
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<td>22</td>
<td>Type 1 VWD</td>
<td>Hysterosalpingo-oophorectomy</td>
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<td>38.8</td>
<td>3.6</td>
<td>4.9</td>
<td>2.5</td>
</tr>
</tbody>
</table>

Abbreviations: D, day, H, hour (after loading bolus dose); HA, hemophilia A; VWD, von Willebrand disease; RBCs, red blood cells.

*The patient received an additional bolus dose of 16 IU/kg.

bUse of tourniquet.

cThe patient received an additional bolus dose of 13 IU/kg.

dThe patient received an additional bolus dose of 14 IU/kg.

^eCompartmental analysis.

gTransfusion prescribed before surgery (hemoglobin 9.8 and 8.1 g/dL in patient 17 and 21, respectively) and administered during the surgery.

^hArthroscopy of the left knee, shaving and micro fractures of chondral defects, synovectomy and arthroscopy and arthrolysis of the right knee, arthrolysis, removal of adhesions.
the first 72 hours after surgery, four FVIII:C levels were below
the target level in four patients but without being lower than
71 IU/dL. As per protocol, three of them received an addi-
tional bolus injection—with pdFVIII (between 13 and 16
IU/kg) to correct levels. In VWD patients (►Fig. 2C), median
VWF:RCo levels were maintained over 70 IU/dL and FVIII:C
levels over 100 IU/dL along 6 days, indicating an effective
FVIII binding to VWF infused. ►Table 4 shows individual data
in both diseases. A total of 14/174 (8%) FVIII:C or VWF:RCo
levels were lower than expected during the perioperative
period, hence daily 92% were on target. The six surgeries in
these patients with severe VWD were performed without
adding FVIII concentrate.
No unfractionated heparin nor low-molecular-weight hepa-
rin or antifibrinolytics were administered in HA patients.
Prophylactic enoxaparin was administered in two patients
with VWD: the first undergoing a total knee replacement and
the second a hemicolectomy, both starting from the operative
day. The first patient also received tranexamic acid the day of
the surgery.

**Safety**
The CI period ranged from 7 to 13 days with a median factor
consumption per surgery (after the loading dose) of 69 IU/kg
of FVIII per day in Study 1, 65 IU/kg of FVIII per day in Study 2,
and 65 IU/kg of VWF:RCo per day in Study 3. No side effects
were recorded. Across all studies, a total of three nonserious
adverse events related to treatment were reported in two
patients: an infusion site reaction followed by an increase in
diastolic blood pressure during the bolus administration for
the preoperative PK resolved within 10 minutes without
medication (Study 3); a venepuncture site inflammation
occurring after 2 days of CI, which led to the interruption
of the study drug for few minutes to change the infusion site
and resolved without other action (Study 3).

**Table 3 Infusion rates and evolution of clearance during the first 2 days by study**

<table>
<thead>
<tr>
<th></th>
<th>H0–H12</th>
<th>H12–H24</th>
<th>H24–H48</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Study 1 (HA), N = 12</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Recommended multiplying coefficient</strong></td>
<td>× 2.5</td>
<td>× 2</td>
<td>× 1.5</td>
</tr>
<tr>
<td><strong>Mean (SD)</strong></td>
<td>6.8 (1.4)</td>
<td>6.9 (1.4)</td>
<td>5.3 (1.0)</td>
</tr>
<tr>
<td><strong>Median (min–max)</strong></td>
<td>6.8 (4.5–8.9)</td>
<td>7.4 (4.9–8.7)</td>
<td>5.3 (3.6–6.8)</td>
</tr>
<tr>
<td><strong>Study 2 (HA), N = 4</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Recommended multiplying coefficient</strong></td>
<td>× 2.5</td>
<td>× 2</td>
<td>× 1.5</td>
</tr>
<tr>
<td><strong>Mean (SD)</strong></td>
<td>5.6 (3.4)</td>
<td>5.4 (2.5)</td>
<td>5.0 (2.5)</td>
</tr>
<tr>
<td><strong>Median (min–max)</strong></td>
<td>4.3 (3.2–10.7)</td>
<td>4.7 (3.2–9.0)</td>
<td>4.4 (2.6–8.5)</td>
</tr>
<tr>
<td><strong>Study 3 (VWD), N = 6</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Recommended multiplying coefficient</strong></td>
<td>× 2.0a</td>
<td>× 1.5a</td>
<td>No corrective factor</td>
</tr>
<tr>
<td><strong>Mean (SD)</strong></td>
<td>5.8 (1.0)</td>
<td>4.0 (0.7)</td>
<td>4.6 (1.3)</td>
</tr>
<tr>
<td><strong>Median (min–max)</strong></td>
<td>5.7 (4.8–6.9)</td>
<td>4.0 (2.8–4.7)</td>
<td>4.3 (3.3–6.9)</td>
</tr>
</tbody>
</table>

Abbreviations: H, hour (after loading bolus dose); HA, hemophilia A; N, number of patients; SD, standard deviation; VWD, von Willebrand disease.

Note: The recommended multiplying coefficient was applied in the infusion rate formula as follows: rate of infusion (IU/kg/h) = multiplying coefficient × individual clearance (mL/h/kg) × desired factor level (IU/mL).

aAs per protocol, the recommended multiplying coefficients (2.0 for the first 12 hours and 1.5 for the following 12 hours) was applied for the 4 first enrolled patients. After a protocol amendment, the multiplying coefficients remained at the investigator’s discretion.
bVWF:RCo level missing in one patient.

![Fig. 1](image-url)  
**Fig. 1** Evolution of the mean clearance of FVIII:C and VWF:RCo during the course of continuous infusion with pdFVIII (100 IU/mL and 200 IU/mL) and pdVWF (100 IU/mL). Clearances were calculated from factor infusion rate and factor measurements at the steady state. pdFVIII, plasma-derived factor VIII; pdVWF, plasma-derived von Willebrand factor; VWF:RCo, von Willebrand factor ristocetin cofactor.
There was no sign of local thrombophlebitis or infection during daily inspection of the catheter site. None of the patients developed inhibitors during the study periods.

Discussion

The data from the three clinical studies demonstrated the clinical feasibility and relevance of replacement therapy with pdFVIII and pdVWF administered for at least 6 days by CI in patients undergoing major surgeries. No treatment-related serious adverse events, development of inhibitory antibodies, or thrombotic events were reported.

As a prerequisite for application via CI, pharmaceutical stability of products was demonstrated: no changes in the FVIII and VWF concentrate characteristics were detected at least within 72 hours when stored in ambulatory pumps. Strengths up to 200 IU/mL of the pdFVIII concentrates appear favorable for CI as no dilution was required after reconstitution avoiding a possible denaturation.

For all 22 surgeries, hemostatic efficacy assessment scores were classified as excellent except for one surgery classified as good. Importantly, results were comparable with those reported in pivotal studies with the same product but administered by bolus.24,25 The treatment recommendations called for levels of FVIII:C above 80 IU/dL in HA, and levels of VWF:RCo above 70 IU/dL in VWD during the post-operative period, and these objectives were met. In patients with VWD, mean FVIII:C remained stable over time after the second day of VWF treatment (Fig. 2C). These findings are in line with those from an earlier study reported by Borel-Derlon et al, which also provide identical FVIII profile when patients received intermittent boluses.25 Most patients with HA or VWD achieved plasma levels (94% for FVIII:C and 100% for VWF:RCo, respectively) above the lowest predefined target range levels during the first 24 hours after surgery, a well-known crucial period. This result was in contrast to published data by Hazendonk et al in a retrospective study evaluating a large population of 119 HA patients undergoing a total of 198 surgical procedures including 115 surgeries performed by CI. The authors found that 44.5% (137/308) of FVIII levels were below the same lowest predefined target range.25 Although the sample sizes were different between studies to allow for a strict comparison, our results were in favor of mitigating the risk of drop under the targeted FVIII level by increasing the initial infusion rate.36,37 The initial infusion rate was based on the baseline clearance corrected by a multiplying coefficient to anticipate coagulation factor consumption. While a decreasing multiplying coefficient from 2.5 to 1.5 over 48 hours provides an effective guide to managing initial infusion rates in HA, its use from 2.0 to 1.5 over a shorter period (24 hours) in VWD seemed not clearly optimal since some levels of VWF:RCo >200 IU/dL were observed. Two of four patients with type 3 VWD had VWF: RCo levels at 211 and 294 IU/dL 24 hours after the CI initiation, suggesting that the multiplying coefficient could be reduced. In both diseases, the inter-patient variability with respect to incremental recovery and estimated clearance observed at baseline stressed the importance of a presurgical PK evaluation and individualized approach. The decline in clearances of infused FVIII observed in our patients over the treatment period remained comparable to other studies.33,38

The CI treatment regimen successfully stabilized FVIII or VWF levels, mitigating the risk of thrombosis since constitutively high plasma FVIII levels have been reported to be associated with increased risk of venous thromboembolism.39,40 Other safety issues in the context of replacement therapy with CI are potential risk of infection or phlebitis at the infusion site, but no sign of local thrombophlebitis or infection during daily inspection of catheter sites was observed during the observation period of the three studies. No adverse events such as thromboses, wound infection, or inhibitor development were observed.
In conclusion, taken together, the results reported in the present article show that products used by CI are effective and well tolerated when used to cover major surgical procedures.

**What is known about this topic?**

- Replacement therapy by continuous infusion aims to provide patients with a safe and constant level of the deficient coagulation factor.
- Continuous infusion is known to be an effective alternative to bolus injections in clinical management of surgical procedures in patients suffering from bleeding disorders.
- Causes influencing continuous infusion rates are the product, its adsorption in the infusion system, its dilution, the type and course of surgery, and the inter-patient pharmacokinetic variability.

**What does this paper add?**

- Continuous infusion of pdFVIII or pdVWF, i.e., Factane and Wilfactin respectively, is effective to control bleeding, and represents a safe alternative to bolus injections to patients undergoing major surgery.
- Continuous infusion of undiluted pdFVIII or pdVWF maintains factor levels within the hemostatic target range during surgery and in the postoperative period.
- Considering inter-patient variability of FVIII and VWF pharmacokinetic parameters, a presurgical pharmacokinetic evaluation is helpful to individualize treatment.

**Author Contributions**

J.W., C.N., and V.C. contributed in designing the studies, and analyzed and interpreted the data. B.G., L.R., A.F., and
E.S.-W. contributed to patient enrolment and collected the data. S.P. and C.H. contributed to the analysis and wrote the manuscript. F.B. designed the studies, interpreted the data, and wrote the manuscript. All authors critically revised the manuscript and approved the final version.

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Conflict of Interest
J.W. received grant support from Alnylam Pharmaceuticals, Baxalta, LFB, Novo Nordisk, Octapharma, Rigel Pharmaceuticals, Roche, Shire/Takeda, and Sobi; also sponsored lectures from Alexion, Baxalta, CSL Behring, Ferrin Pharmaceuticals, Novo Nordisk, Octapharma, Roche, Sanofi/Genzyme, Shire/Takeda, Siemens, Sobi, and Werfen. B.G. has received grant support from CSL Behring, Octapharma and Sobi; also sponsored lectures from CSL Behring, Novo Nordisk, Roche, and Shire/Takeda. L.R. has no conflicts of interest to declare. E.S.-W. has no conflicts of interest to declare. V.C. has received honoraria or consultation fees from Roche, Novo Nordisk, and Sobi, also support for attending scientific meetings from LFB and Sobi. S.P. is an employee of LFB; C.H. is an employee of LFB. F.B. is a former employee of LFB. C.N. received research grant or honoraria or participated in clinical trials from Alnylam/Sanofi, Bayer, BioMarin, Bioverativ/Sobi, Bayer, CSL Behring, Novo Nordisk, Octapharma, Pfizer, Roche/Spark, and Shire/Takeda.

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References