Management of Surgical Procedures in Patients with Inherited F VII Deficiency: Six Years of Experience

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Abstract

Inherited factor VII (FVII) deficiency is the most frequently observed rare bleeding disorder. The clinical symptoms are extremely variable, and patients may be asymptomatic or may present with life-threatening bleeding disorders. FVII activity (FVII:C) values of 10–15% are considered safe for maintenance of hemostasis, and replacement therapy is recommended for some types of surgical procedures. Twenty-three adult patients with FVII:C deficiency were followed-up in the Ostrava Haemophilia Treatment Centre. Eleven patients underwent a total of fourteen invasive procedures between 2008 and 2013. In terms of replacement therapy, nine patients received plasma-derived FVII and five patients received activated recombinant FVII. None of the patients had excessive blood loss during surgery, and there were no bleeding or other complications during post-operative treatment. In addition, there were no thromboembolic events related to the use of replacement therapy.

Keywords: Inherited FVII deficiency; Surgery; Replacement therapy; Antithrombotic prophylaxis

Introduction

Inherited coagulation factor VII deficiency is the most frequently observed rare inherited coagulation disorder. The heredity of the disease is autosomal recessive, and the prevalence of a serious disorder (FVII: C < 2%) is estimated to be 1:300,000–1:500,000 inhabitants [1]. FVII: C shows variable clinical manifestation and can range from a clinically asymptomatic course in which only laboratory findings are used for the diagnosis to moderate bleeding disorders to life-threatening bleeding episodes [2,3]. In many cases, the seriousness of the bleeding does not correspond to the seriousness of the laboratory findings and most probably is not influenced by the genetic mutation either. For example, one individual with very low residual FVII activity may exhibit no bleeding problems or only a minor bleeding disorder, whereas another patient with much higher FVII activity may suffer from a serious bleeding disorder. Interestingly, patients with the exact same homozygous mutation can have different clinical phenotypes. It is thus likely that other genetic factors or environmental factors also play important roles in the disorder [3,4]. Clinical manifestations of FVII:C deficiency include predominant bleeding into tissues rich in tissue factor such as the brain, intestines, uterus, placenta, and lungs, and bleeding episode onset may be associated with a surgical procedure [2]. Frequent hemorrhages in women can result in iron-deficiency anemia. Bleeding into joints and muscles is not as typical as in hemophilia; nevertheless, this can occur in both men and women with the disorder. Upon laboratory examination, FVII:C deficiency is characterized by isolated prolongation of the prothrombin time, which is not adjusted with a corrective test performed with normal plasma. Low levels of FVII:C can also be detected by determining the functional activity of FVII with the one-stage prothrombin test with optical detection using FVII-deficient plasma. FVII: C values of 10–15% are considered a hemostatic minimum, and replacement of FVII is recommended for surgical procedures. Considering the safety risks, the replacement of choice is activated recombinant FVII (rFVIIa), or plasma-derived FVII concentrate (pdFVII); fibrin glue may also be used in certain circumstances to support local hemostasis (e.g. in stomatotomy). When these preparations are not available, concentrate of prothrombin complex (PCC) factors or plasma may also be used [1].

The aim of study is to present a management of substitution therapy in patients with different level of FVII deficiency either with pdFVII or with FVIIa in different invasive procedures performed within five years (2008 and 2013) at the University Hospital Ostrava, Czech Republic.

Patients and Methods

A total of 23 adult patients with FVII: C deficiency were treated and followed-up from 2008 to 2013 at the University Hospital Ostrava. The FVII: C level was below 1% in six patients, between 1.0 and 10.0% in ten patients, and between 10.1 and 35.0% in seven patients. The most frequently observed bleeding disorders in the female patients were frequent subcutaneous hematomas, epistaxis, and menorrhagia. The medical history of four patients included past episodes of serious bleeding and impaired wound healing following surgical procedures. We did not encounter any cases of intra-articular bleeding or bleeding into the gastrointestinal tract or the central nervous system. In eight of the patients, the diagnosis was made when preoperative coagulation tests were performed or during family screening. FVII activity was assessed with a coagulation test that used DG-FVII (Grifols) deficient plasma, the PT reagents TriniClot PT Excel S (Trinity Biotech), and the Sysmex CA 1500 analyzer.

We prepared and followed the perioperative courses of eleven patients who underwent fourteen surgical procedures between 2008 and 2013. Nine patients were treated with replacement therapy using pdFVII (Factor VII; Baxter Healthcare) (Table 1), and five were treated with rFVIIa (NovoSeven) (Table 2). Two of the surgical procedures were performed laparoscopically.

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Inherited FVII: C deficiency is a relatively rare bleeding disorder. Nevertheless, it is likely that the indicated prevalence is underestimated or is regionally dependent, which was confirmed by the number of patients followed at our center (23 patients in a 6-year period).

In two of our female patients, the disorder was diagnosed based on preoperative assessment and was specifically diagnosed based on routine coagulation tests performed prior to planned hysterectomy.

The indication for the surgical procedure was reassessed following consultation with the attending gynecologist, and these two patients were treated with conservative therapy (anti-fibrinolytic medication plus hormonal therapy).

In making a differential diagnostics, it is important to differentiate between acquired FVII:C deficiency that occurs in the absence of vitamin K versus a clinically-mute history of bleeding disorders. The former is more likely in older patients, in neglected or oligophrenic patients with nutritional disorders, in patients with nutrition absorption problems, and in patients who have received long-term antibiotic therapy. The most frequent cause of acquired FVII:C deficiency is treatment with coumarins, which is usually apparent from the patient's pharmacological history.

Low levels of FVII:C do not protect the patient from possible thrombosis. In addition to classical cases of thromboembolic disorders, atypical thrombosis localization has been described. Thromboembolic disorders can also occur in very young patients and without any relation to replacement therapy, e.g. in the portal or splenic veins, central retinal vein, or axillary and cerebral veins [6]. That is why it is always necessary to carefully evaluate antithrombotic prophylaxis, especially in patients with multiday FVII replacement therapy.

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**Conclusion**

We did not observe any excessive blood loss in the course of surgical procedures performed on patients treated with rFVIIa and pdFVII replacement therapy. None of the patients had bleeding complications during the postoperative course of treatment, and the surgical wounds healed per primam. None of the patients had thromboembolic complications. Thromboembolic prophylaxis was administered in one patient following total hip replacement for the recommended period of five weeks using LMWH without any hemorrhagic manifestations.

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References


5. SPC Factor VII Baxter 600 IU.