Factor IX propeptide mutation and life threatening bleeding

Conflict of interest statement The autors reported no potential conflicts of interest.

A. Vaccarino*, M. Bazzan, O. Giachino
Rare Disease, Immunological and Haematological Department University of Turin
*Corresponding author. Tel.: +39 0112402056 fax: +39 38012402052
E-mail address: antonella.vaccarino@libero.it (A. Vaccarino)

P. Colagrande
Internal Medicine Unit Torino Nord Emergenza San Giovanni Bosco Hospital

P. Ferraresi
Biochemistry and Molecular Biology University of Ferrara, Italy

S. Stella D. Roccatello
Rare Disease, Immunological and Haematological Department University of Turin

F. Bernardi
Biochemistry and Molecular Biology University of Ferrara, Italy

Dear Editors,

Bleeding events are the most relevant complications during oral anticoagulant treatment (OAT). They have been reported to occur at a rate of about 1.35 cases each 100 patients/year [1]. In very rare instances, OAT-related bleeding in patients with INR values within the therapeutic range may be associated with a mutation of the factor IX (FIX) gene. Two propeptide mutations have been described at locus-10: Ala (GCC)-10 Val (GTC) and Ala (GCC)-10 Thr (ACC) [2,3]. The propeptide sequence at locus-10 plays an important role in the anticoagulant effect of vitamin K antagonists (VKA), since it contains the hepatic γ-carboxylase binding site [4]. In the presence of a propeptide locus-10 mutation, hepatic γ-carboxylase binding to the FIX protein is markedly reduced. This reduced binding capacity has no clinical implications (except during OAT), since the FIX plasma level is within the normal range. When patients with the mutation are on OAT, the lack of biologically active vitamin K associated with reduced binding of γ-carboxylase to the coagulation profactors, may induce a reversible but serious bleeding. This happens anytime, from days to weeks, after starting VKA treatment. The bleeding is caused by a rapid decrease in FIX activity (FIX:C), which drops to disproportionately lower values compared with the others vitamin-K dependent coagulation factors. FIX:C levels drop so low that laboratory data mimic acquired haemophilia B. Thus, patients with the mutation cannot be detected by routine monitoring of the INR. However, the sudden and disproportionately severe drop in factor IX plasma levels (b3%) induced by OAT leads to persistently abnormal prolongation of activated partial thromboplastin time (aPTT). The morbidity associated with VKA–related bleeding, and the possibility of identifying mutated patients by performing aPTT has led us to question feasibility of aPTT screening in all patients on VKA. The main argumentation against aPTT screening is the low prevalence of the mutation [5–7]. A 61 year old man was admitted to the emergency department because of an “idiopathic” proximal deep-vein thrombosis of the right leg. He was treated with LMWH, followed by VKA. Heterozygosity for Factor V Leiden mutation, and no other
thrombophilic abnormality, was found. After about two months of OAT the patient went to the emergency department of another hospital on account of severe pain and widespread ecchymosis of the right hand, which occurred the day after using a drill. Spontaneous muscle bleeding of the right thigh was also reported. Laboratory data showed an INR value of 2.15, but an abnormally prolonged aPTT ratio 4.93 (n.v. ≤ 1.2). The patient was discharged with a diagnosis of OAT related bleeding. He had never experienced any bleeding complications before starting OAT and there was no history of bleeding in his family. One month later, the patient returned to the same emergency department because of bilateral leg pain caused by deep muscle bleeding. INR was 2.1 and aPTT ratio was 5.63. The patient was again discharged with the same diagnosis. VKA dose was reduced and paracetamol was administered.

Two days later, the patient came to our hospital because of worsening of the bleeding, widespread ecchymosis on the chest, legs, hands and arms, and great difficulty in swallowing. Physical examination revealed tongue and pharynx haematoma. Laboratory data results were: haemoglobin 6.9 g/dl, INR 2.21 and aPTT ratio 5.38. VKA was discontinued and the patient underwent blood transfusion. Three days after discontinuing OAT, INR returned to 1.3 and aPTT ratio to 1.85, while after seven days both INR and aPTT ratio returned to normal values (Fig. 1). At the same time, factors II, VII and X were assayed and found within normal range, while factor IX was about 40%. Haematologic counselling was then requested. The patient’s clinical history showed that both INR and aPTT ratio were within normal ranges before starting OAT. The patient reported no previous significant bleeding episodes, and familial bleeding tendency was excluded, while inherited bleeding tendency was unlikely. Due to the normal platelet count and normal antithrombin plasma levels, we ruled out disseminated intravascular coagulation. The possible presence of antibodies against factor VIII:C or IX:C was ruled out because of the quick and spontaneous return to normal aPTT ratio values after OAT discontinuation. The only anomaly we observed in coagulation tests was a disproportionate aPTT ratio prolongation when compared to INR prolongation. A literature search was then carried out and a mutation of the factor IX propeptide was hypothesized. This is a genetic predisposition to bleeding during OAT. Blood samples were sent off for molecular evaluation. The first three exons of the factor IX gene were amplified and sequenced. Whole exon 1 was amplified by the following primers: forward 5’-ATT GAG GGA GAT GGA CAT TAT T-3’ (nt 2892–2913) and reverse 5’-AGT GAA GAA GAC AGC ATC AGA TAT-3’ (nt 3159–3136); exons 2 and 3 were amplified together using forward 5’-TTC ATG ATG TTT TCT TTT TTG CTA-3’ (nt 9237–9260) and reverse 5’-TGC AGA GAA AAA ACC CAC AT-3’ (nt 9737–9718) primers (Gene Bank K02402.1). PCR amplifications were run for 30 cycles with a starting denaturation at 95 °C for 5 min, 95 °C for 30 s, 58 °C for 30 s and final extension at 72 °C for 7 min. Direct nucleotide sequencing revealed a hemizygous -10Ala (GCC)-NThr (ACC) mutation. Only few case reports concerning the bleeding tendency in patients on OAT with factor IX propeptide mutation are present in the literature: most cases diagnosed in Switzerland and Germany were due to a founder mutation, with a clear regional focus [8–12]. To our knowledge this is the first case reported in Italy and actually it is not known if the mutation has an independent origin or results from a common founder. The bleeding tendency is usually severe, but the prevalence of the disorder in the general population is very low, although probably underdiagnosed. In our case, the patient had a very serious, life-threatening bleeding event. We highlight the fact that the significant aPTT prolongation was never considered a severe laboratory anomaly. Factor IX propeptide mutation is not well known among general practitioners or in the emergency setting. Due to the low prevalence of the defect in the general population, aPTT screening is proven not to be cost effective if evaluated in all patients on OAT. Data from literature and our experience suggest that if a male patient have significant bleeding events in the first weeks or months of OAT and INR is within the therapeutic range, the aPTT should be performed. Immediate OAT withdrawal and replacement with an alternative anticoagulant treatment would prevent a worsening of bleeding symptoms.
References


Fig. 1. International Normalized Ratio (INR-continuous line) and Activated Partial Thromboplastin Time Ratio (aPTTR-dotted line) during Oral Anticoagulant Therapy (OAT). Arrows indicate bleeding events and OAT reduction or discontinuation.