Coexistence of Factor V G1691A and Factor II G20210A Gene Mutations in a Thrombotic Family Is Associated with Recurrence and Early Onset of Venous Thrombosis

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Key Words
Factor V G1691A · Prothrombin G20210A · Inherited thrombophilia · Combined defects · Gene-environment interaction

Abstract
Two G-to-A mutations at positions 1691 of the factor V (FV) gene and 20210 of the prothrombin (FII) gene have been associated with an increased risk of venous thrombosis. We report a thrombosis-prone family in which one subject – the propositus who exhibited combined heterozygous FV G1691A and FII G20210A mutations – showed spontaneous and early clinical onset (at 23 years), recurrences of deep-vein thrombosis and pulmonary embolism. His asymptomatic father carried the FII G20210A substitution and his mother, characterized by an isolated thrombotic episode on occasion of surgery (at 48 years), carried the FV G1691A substitution. In the maternal lineage, one of the propositus’ uncles had thrombosis on occasion of a bone fracture (at 65 years) despite the absence of known prothrombotic defects. A sister of the propositus carried the FII G20210A and the brother the FV G1691A mutation. They have been asymptomatic until now. The propositus’ two children, 20 and 16 years old, both carry the FV G1691A substitution and have been asymptomatic until now. The plasma levels of FII were higher in carriers of the FII G20210A allele if compared with noncarriers, and the activated protein C resistance phenotype, associated with the FV Leiden mutation, showed a complete correlation with the FV G1691A mutation. Despite the very limited number of thrombotic cases involved in this survey, which does not allow statistically sound conclusions, the data obtained from this family suggest that the syn-
Introduction

Familial thrombophilia is considered a polygenic and multifactorial disorder in which gene-gene and gene-environment interactions play a pivotal role [1–3]. Point mutations in the factor V (FV) gene (FV G1691A, FV Leiden) [4, 5] and in the prothrombin (FII) gene (FII G20210A) [6] presently account for most cases of inherited thrombophilia. The FV G1691A allele is found in 90–95% of people with the activated protein C (APC) resistance phenotype [7]. This suggests that various mechanisms are responsible for the same phenotype [8]. Increased levels of plasma FII are associated with the FII G20210A allele [6, 9], but the precise mechanism by which prothrombin levels are altered has yet to be determined. Common inherited prothrombotic factors are associated with a low thrombotic risk compared with the defects that are uncommon but associated with a high risk of thrombosis [10]. Direct evidence from case-control and familial studies suggests that the coexistence of distinct prothrombotic defects increases the risk of developing thrombosis in the presence or absence of transient risk factors [11–17]. In addition, whether or not FV Leiden and FII G20210A mutations separately increase the risk of recurrent venous thrombosis is a debated question, and it has recently been reported that only carriers of both mutations have an increased risk of recurrences and are candidates for lifelong anticoagulant therapy [17]. We describe here a family with asymptomatic subjects carrying inherited prothrombotic defects (i.e. FV Leiden or FII G20210A), the propositus who carries both the FV and FII mutations with spontaneous and recurrent deep venous thrombosis (DVT) and pulmonary embolism in combination with transient risk situations, and his mother who is heterozygous for FV Leiden and has experienced an episode of thrombosis on occasion of surgery. Finally, one of the propositus’ uncles had thrombosis despite the absence of known prothrombotic defects.

Patients and Methods

Family History

Figure 1 shows the family tree of the family investigated. Subject 14, a 75-year-old woman, had an isolated episode of DVT after an appendicectomy at the age of 48. She was treated with oral anticoagulant therapy for a period of 6 months and she has since been free of thrombosis. One of her two brothers, subject 15, a 73-year-old man, had an isolated episode of DVT at the age of 65 years on occasion of a long period of bed rest because of a fracture of the leg. Subject II4 (the propositus), a 43-year-old male, had a first spontaneous episode of DVT at the age of 23 years with relapses in the following years. After a period of immobilization owing to a bone fracture, he suffered a further episode of DVT and pulmonary embolism at the age of 32. He is on lifelong oral anticoagulant therapy. All the remaining members of the family (affected and nonaffected) are currently asymptomatic.

Coagulation Studies

Anticoagulant response to APC was tested by an activated-partial-thromboplastin-time-based method (ProC®Global, Dade Behring) that was modified as previously described [18]. The results were expressed as normalized APC ratio (nAPC-r; normal values ≥0.80). Plasma prothrombin levels (normal values 70–125%) were measured using an automated one-stage functional assay (Dade Behring). All family members investigated were healthy and asymptomatic at the time of testing and did not receive any pharmacological treatment potentially interfering with coagulation assays.

DNA Analysis

DNA was extracted from peripheral blood utilizing standard procedures (Puregene®, Gentra Systems). For
the one-step determination of the FV G1691A and FII G20210A mutations, we utilized multiplex PCR-mediated site-directed mutagenesis, creating a neo-site for TaqI endonuclease in both the FV and FII gene-amplified fragments as recently described by Ripoll [19].

Results

Table 1 shows the coagulation findings, the DNA analysis related to the FV Leiden and FII mutations and the clinical characteristics of the family investigated. All family members underwent screening for thrombophilia as previously described [20] and none of them exhibited other known risk factors for venous thrombosis.

DNA Studies

The propositus (subject II4) was heterozygous for both the FV Leiden and the FII gene mutations. The remaining affected family members were heterozygous either for the FV Leiden (subjects I4, II3, III1 and III2) or the FII gene mutation (subjects I3 and II1). The other family members (subjects I2, I5, I6, II2 and II5) did not show either of the mutations (table 1).
Table 1. Coagulation, genotypic and clinical characteristics of the family investigated

<table>
<thead>
<tr>
<th>Subjects</th>
<th>APC-R (nAPC-r)</th>
<th>FV G1691A</th>
<th>FII activity, %</th>
<th>FII G20210A</th>
<th>Thrombosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>I2</td>
<td>0.96 GG</td>
<td>100</td>
<td>GG</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>I3</td>
<td>1.10 GG</td>
<td>135</td>
<td>GA</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>I4</td>
<td>0.65 GA</td>
<td>114</td>
<td>GG</td>
<td>48 no</td>
<td>no no</td>
</tr>
<tr>
<td>I5</td>
<td>0.90 GG</td>
<td>110</td>
<td>GG</td>
<td>65 no no</td>
<td>no no</td>
</tr>
<tr>
<td>I6</td>
<td>1.05 GG</td>
<td>95</td>
<td>GG</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>I11</td>
<td>0.99 GG</td>
<td>130</td>
<td>GA</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>I12</td>
<td>1.15 GG</td>
<td>95</td>
<td>GG</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
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<td>0.67 GA</td>
<td>100</td>
<td>GG</td>
<td>–</td>
<td>–</td>
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<td>I14</td>
<td>0.60 GA</td>
<td>150</td>
<td>GA</td>
<td>23 yes yes</td>
<td>–</td>
</tr>
<tr>
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<td>125</td>
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<tr>
<td>Normal</td>
<td>≥ 0.80 GG</td>
<td>70–125</td>
<td>GG</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

The pathologic findings are shown in italics.

1 Propositus.

Coagulation Studies

The subjects who carried the FII G20210A mutation had plasma levels of prothrombin above the upper limit of the normal range (70–125%) and of the levels observed in the subjects with a normal FII genotype (95–125%). The APC resistance phenotype associated with FV Leiden, measured as nAPC-r, showed a complete correlation with DNA analysis being below the normal cutoff value (nAPC-r = 0.80) in all the individuals with the FV Leiden mutation. It is of note that the propositus showed the highest prothrombin level and the lowest nAPC-r value of the family members investigated (table 1).

Discussion

When genetic defects are absent, thrombosis occurs in the older population, often with strong environmental risks such as surgery, bone fractures or cancer [1–3]. In contrast, inherited thrombosis is associated with younger age of onset; it is due to the presence of one or more genetic defects in which gene-gene and gene-environment interactions play a role [1–3]. FV Leiden and FII G20210A mutations account for most cases of inherited thrombophilia and they are relatively common among Caucasians [21, 22]. Therefore thrombotic patients with combined defects are not so rare.

We describe a family in which the propositus (II4) was characterized by the combined presence of heterozygous FV Leiden and FII G20210A mutations. The severity of symptoms, the recurrences and the early onset of a spontaneous thrombosis together with a positive family history recommended a long-term anticoagulant treatment. These features could be due to the coexistence of two inherited prothrombotic defects in conjunction with important triggering factors such as bone frac-
ture and immobilization. Most likely, these were the only prothrombotic conditions in the propositus, since all family members underwent screening for thrombophilia, which did not reveal any other genetic defects. The propositus’ mother (I4) and one of her two brothers (I5), were the other symptomatic members of the family. The mother was heterozygous for the FV Leiden mutation and had an isolated thrombosis on occasion of surgery at age 48. After anticoagulant treatment was stopped she remained free of thrombosis until today. De Stefano et al. [17] reported that in patients with only an FV Leiden mutation, the risk of recurrences was nearly identical to that in patients with no mutation, regardless of the type of recurrence (any, spontaneous, spontaneous after either a first idiopathic event or in conjunction with defined risk factors). The risk of spontaneous recurrences in the propositus’ mother should fall within the latter subgroup, since she had her first episode on occasion of surgery. It is of note that she had had four pregnancies with no complications in the period from 23 to 32 years. Transient risk situations such as pregnancy, childbirth, surgery and trauma are known risk factors for thrombosis, especially in genetically affected subjects in whom they could act as precipitating factors. However, not all subjects carrying an FV Leiden mutation experience thrombosis [23–25]. The increased risk of thrombosis is lifelong and increases with age. In a study of 50 Swedish families with inherited APC resistance, the thrombosis-free survival curves showed that at 33 years of age only 20% of heterozygotes and 40% of homozygotes had manifestations of venous thrombosis [7]. The propositus’ mother had pregnancies when she was relatively young (23–32 years), and established prothrombotic risk factors did not cause any clinical accident in her, while at older age thrombosis occurred on occasion of surgery. The modulatory role of age in combination with circumstantial risk situations at the onset of thrombosis in genetically predisposed patients is clearly shown in this case. Finally, in the maternal lineage, subject I5 had thrombosis in the absence of known prothrombotic defects, following a long period of immobilization due to a bone fracture. The importance of coinherited genetic risk factors and environmental risk conditions in the occurrence of thrombosis is well demonstrated by the present study, although in this patient (subject I5) no congenital defects were found. It is of note that in this subject thrombosis occurred at an older age and in combination with strong environmental risk conditions. However, it could be speculated that some inherited predisposition was present in him, since thrombosis occurred in his sister too.

To summarize, in this family thrombosis occurred in the propositus who exhibited combined FV and FII mutations, with a first idiopathic episode (at 23 years) and further recurrences in conjunction with acquired risk conditions, his mother who had an FV mutation suffered an isolated episode on occasion of surgery (at 48 years) and an uncle of the propositus also experienced an isolated episode despite the absence of prothrombotic defects (at 65 years). This means that the maternal lineage, where other unidentified prothrombotic influences could be present, can be identified as the thrombosis-prone lineage in which most likely common prothrombotic polymorphisms can cause familial thrombosis. On the other hand, all family members, affected and nonaffected ones, have a finite risk of developing thrombosis. In a recent large study on family members with FV and/or FII gene mutations, Martinelli et al. [26] reported that the absolute annual risk of a first episode of venous thrombosis was higher in carriers of both mutations (0.42%) and lower in cases with either mutation (0.19%.)
and 0.13% for FV and FII gene mutation, respectively) compared with 0.066% in relatives with neither mutation. In particular, a recent report on the risk assessment of single and combined prothrombotic polymorphisms in patients with idiopathic venous thromboembolism ascribed the highest risk to the coexistence of the FV Leiden and the FII gene mutations [17]. Moreover, in thrombophilic families, the FII G20210A allele was recently found to be associated more frequently with the FV G1691A mutation than with the protein S defect [27].

In the family we studied, the combination of asymptomatic and genetically affected subjects and a thrombotic relative who is genetically unaffected is in accordance with the multifactorial nature of thrombosis and raises the problem whether the presence or absence of defined prothrombotic defects (i.e. FV Leiden or FII gene mutations) should influence the use of primary prophylaxis or the duration of treatment after an episode of thrombosis [26]. The decision should take into account that the risk of thrombosis has to be weighed against the risk of bleeding associated with oral anticoagulation ranging from 1.1 to 3.8% per patient-year [28, 29].

Notwithstanding the limitations of our survey due to the small size of the population studied and the fact that up till today it is not certain whether FV Leiden or FII gene mutations are by themselves independent risk factors for venous thrombosis, the presented family could be a good model which allows to highlight and to discuss the multifactorial and polygenic nature of thrombosis in which inherited and acquired conditions act in synergism on the onset and progression of venous thromboembolism.

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References


Defect G20210A: A Combined Thrombotic Factor V G1691A and Factor II 16


