Despite the exhaustive screening of \( F7 \) gene exons and exon-intron boundaries and promoter region, a significant proportion of mutated alleles remains unidentified in patients with coagulation factor VII deficiency. Here, we applied next-generation sequencing to 13 FVII-deficient patients displaying genotype-phenotype discrepancies upon conventional sequencing, and identified six rare intronic variants. Computational analysis predicted splicing effects for three of them, which would strengthen (c.571+78G>A; c.806-329G>A) or create (c.572-392C>G) intronic 5' splice sites (5'ss). In \( F7 \) minigene assays, the c.806-329G>A was ineffective while the c.571+78G>A change led to usage of the +79 cryptic 5'ss with only trace levels of correct transcripts (3% of wild-type), in accordance with factor VII activity levels in homozygotes (1-3% of normal). The c.572-392C>G change led to pseudo-exonization and frame-shift, but also substantial levels of correct transcripts (approx. 70%). However, this variant was associated with the common \( F7 \) polymorphic haplotype, predicted to further decrease factor VII levels; this provided some kind of explanation for the 10% factor VII levels in the homozygous patient. Intriguingly, the effect of the c.571+78G>A and c.572-392C>G changes, and particularly of the former (the most severe and well-represented in our cohort), was counteracted by antisense U7snRNA variants targeting the intronic 5'ss, thus demonstrating their pathogenic role. In conclusion, the combination of next-generation sequencing of the entire \( F7 \) gene with the minigene expression studies elucidated the molecular bases of factor VII deficiency in 10 of 13 patients, thus improving diagnosis and genetic counseling. It also provided a potential therapeutic approach based on antisense molecules that has been successfully exploited in other disorders.
mon variants have been identified in the F7 gene ([http://f7-db.caahad.org/](http://f7-db.caahad.org/)). Nevertheless, in spite of an exhaustive direct sequencing of F7 exons and exon-intron junctions and of the proximal promoter region, a significant proportion of defective alleles has still not been identified. The rate of uncharacterized F7 disease alleles ranges from 2% to 8%—10 in Europe, and a similar estimate (7%) was made in India.11

In this context, subtle intronic variations outside the routinely sequenced exon-intron boundaries could have a pathological impact by impairing the splicing process. In fact, precise exon definition during RNA processing requires the interplay among several exonic and intronic splicing regulatory elements,12 which can be altered by nucleotide changes and lead to aberrant splicing.4,13-17 Various examples of “deep” intronic changes associated with miss-splicing have been reported in human disorders, including those involved in coagulation.18-20 It is worth noting that RNA splicing can be modulated for different purposes, including the development of new therapies.14-20,26 In this context, next-generation sequencing (NGS) could represent a powerful tool to characterize gene defects in patients with unknown alleles; however, only a few studies have been conducted in coagulation factor disorders.27-29

Here, we investigated 13 patients with FVII deficiency forms that have not been explained by mutations identified by conventional sequencing and used NGS to identify six rare intronic variations that could be causative. Through expression studies, we demonstrated that two of them lead to aberrant splicing, which explained the residual FVII levels in most patients and, intriguingly, that these can be rescued by an antisense-based correction approach.

### Methods

#### Patients

Since 1997, 400 FVII-deficient patients with FVII coagulant activity (FVII:C) levels <30% were referred to our laboratory for genetic analysis through conventional screening. Among them, 13 (Table 1) showed a genetic profile that, considering the identified mutated allele10,30-32 and the major F7 functional polymorphisms (c.-325_-324insCCATATCCT, A2 allele; p.Arg413Gln change, M2 allele),33-35 appeared to be incompatible with the reduced FVII levels. The local Institutional Review Board approved the study and patients provided informed consent.

#### Measurement of FVII levels

FVII:C and FVII antigen levels were determined by the one-stage method30 and Enzyme-Linked-Immunoabsorbent-Assay (Diagnostica Stago, Asnière sur Seine, France), respectively.

#### DNA genotyping and next-generation sequencing

Conventional Sanger technology was exploited to sequence the F7 exons, including the intronic boundaries and the 5’ untranslated region. Large rearrangements were ruled out using semi-quantitative multiplex fluorescent-polymerase chain reaction (SOF-PCR) assays.39 NGS of the F7 gene was designed to cover the intronic regions with the exception of the highly repetitive GC-rich region in intron 2 (legacy nomenclature, intron 1b). The probe-capture custom design targeted the 14893-pb F7 gene in two parts: chr13:113759000-113761350 and chr13:113764600-113775100 accounting for a total of 12,850 base-pairs with a gap of 3250 bp. DNA library generation was performed using the Custom SureSelectQXT Target Enrichment system (Agilent, Santa Clara, CA, USA) on a MiSeq platform (Illumina, San Diego, CA, USA). The sequencing data were stored in FASTQ format and analyzed using two bio-informatics

### Table 1. Features of the investigated FVII deficient patients.

| Proband | Origins                   | FVII:C (%) | FVII:Ag(%) | Conventional sequencing | Polymorphic pattern | NGS sequencing |
|---------|---------------------------|------------|------------|-------------------------|--------------------|---------------|
|         |                           |            |            |                         | c.325_324ins/     |               |
|         |                           |            |            |                         | p.Arg413Gln        |               |
|         |                           |            |            |                         |                   |               |
| #17     | Maghreb countries         | 3          | uk         |                         |                   |               |
| #19     | France                    | 2          | <5         |                         | p.Met358Ile<sup>+</sup> | het           |
|         |                           |            |            |                         | p.Met1Val*        | het           |
| #31     | France                    | <1         | 19         |                         |                   |               |
|         |                           |            |            |                         | p.Cys162Tyr<sup>*</sup> | het           |
| #113    | France                    | 1          | 66         |                         |                   |               |
| #262    | France                    | 3          | uk         |                         |                   |               |
| #341    | France                    | 16         | uk         |                         |                   |               |
| #214    | France                    | 1          | 3          |                         |                   |               |
| #28     | France                    | 13         | 39         |                         |                   |               |
| #377    | Maghreb countries         | 23         | uk         |                         |                   |               |
| #284 *  | Maghreb countries         | 20         | 15         |                         |                   |               |
| #90 *   | Lebanon                   | <1         | uk         |                         |                   |               |
| #15     | Maghreb countries         | 3          | uk         |                         |                   |               |
| #330    | France                    | 10         | uk         |                         |                   |               |

*Consanguinity NGS: next-generation sequencing; uk: unknown; A1/A2: decanucleotide insertion c.-325_-324insCCATATCCT; p.Arg413Gln polymorphism: rs5742910; M1/M2: p.Arg113Gln polymorphism: rs6046; FVII:C: FVII activity; FVII:Ag: FVII antigen; het: heterozygous; hom: homozygous.
NGS of F7 and minigene studies identify molecular bases of FVII deficiency

Results

Genotyping of FVII deficient patients

Among the 13 selected FVII-deficient patients shown in Table 1, ten had FVII:C levels below 15% and were expected to have two F7 pathogenic alleles. However, the conventional sequencing did not reveal any F7 pathogenic allele for patients #15, #90 and #330, and only one for the remaining seven patients (#17, #19, #28, #31, #113, #214, #262). On the other hand, three patients (#254, #541, #577) presented with FVII:C levels between 15% and 30% but displayed only the c.430+1G>A mutation (#341) or the A2M2 polymorphic haplotype (#254, #577), which points towards the presence of an additional F7 pathogenic allele for each patient. In this scenario, we had a total of 16 F7 uncharacterized alleles to be explored using NGS.

As far as the clinical phenotype is concerned, 3 of 13 patients were symptomatic. Patient #19 presented with bruises and frequent epistaxis, patient #90 had post-traumatic oral bleeding, spontaneous hematuria and rectal bleeding, and patient #214 suffered from provoked hematoma and severe menorrhagia resolved by replacement therapy.

Next-generation sequencing, besides confirming the presence of the causative variants identified by the Sanger approach, also revealed several deep intronic substitutions. Among them, only those with a coverage of 30x and observed in databases with a minor allele frequency (MAF) <0.05 were analyzed further. Six deep intronic substitutions matched these criteria (Table 2). The c.571+78G>A change, whose pathogenic effect is supported by its co-segregation with the disease phenotype in the family pedigree of patient #28 (Figure 1), was the most frequent in our series, being present in ten alleles from unrelated patients living in various areas, including France, North Africa and Lebanon (Table 1). NGS data prompted us to analyze an enlarged panel of polymorphic deep intronic variants in both c.571+78G>A homozygotes (#15, #90), who were homozygous for the major A1 and M1 polymorphic alleles. However, they differed on other intronic variants. Patient #15, of Tunisian origin, was homozygous for two variants, c.-402A (rs510317) and c.292-672G (rs12431329), that are quite rare, with a minor allele frequency (MAF) of 0.233 and 0.213, respectively. By contrast, patient #90, living in Lebanon, showed the c.-402G variant and the c.292-672A variants, and displayed two additional deep intronic polymorphic variants in the homozygous state, the c.64+196G>A (rs2774030) and the c.131-394T>C (rs1745939), with a global frequency of the c.64+196A and c.131-394C alleles of 0.851 and 0.759, respectively. Thus, two different haplotypes associated with the c.571+78G>A mutation could be defined: c.-402A, A1 c.64+196G, c.131-394T, c.292-672G, c.571+78A, M1 (Haplotype 1) and c.-402G, A1,

| Change     | Intron | Prediction | Score (wt/mutated) | Position | rs          | MAF    |
|------------|--------|------------|--------------------|----------|-------------|--------|
| c.64+305G>A | 1      | Weakening cryptic 3’ss | 0.31/0.13          | -8       | 36208414    | 0.005  |
| c.291-846C>T | 2      | Strengthening cryptic 3’ss | 0.51/0.71          | -7       | 565185989   | 0.004  |
| c.571+78G>A | 5      | Strengthening cryptic 5’ss | 0.11/0.79          | -2       | 764741909   | none   |
| c.572-392G>C | 5      | Creation of new 5’ss | -         | +1       | none       | none   |
| c.681+131G>T | 6      | Strengthening cryptic 3’ss | 0.02/0.12          | -8       | 752129277   | none   |
| c.506-329G>A | 7      | Strengthening cryptic 5’ss | 0.79/0.99          | -3       | none       | none   |

Position of the point mutation is referred to the 5’ splice site (5’ss) or 3’ss. RS: reference SNP ID number; MAF: minor allele frequency, based on 1000Genome project (http://www.internationalgenome.org). F7 gene reference sequence is NG_000262.1. NNSPLICE 0.9 software (www.fruitfly.org/seq_tools/splice.html) was used to predict and calculate the 5’ss or 3’ss of the score. Introns are indicated by legacy nomenclature wt: wild-type.
c.64+196A, c.131-394C, c.292-672A, c.571+78A, M1 (Haplotype 2). The analysis was also extended to the c.571+78G>A heterozygotes, which revealed that haplotype 1 is compatible with patients of Maghreb origin whereas haplotype 2 is compatible with patients of European origin.

The c.572-392C>G, c.64+305G>A and c.806-329G>A variants were also relatively frequent as they were found in three, two and two alleles, respectively, whereas the remaining were identified only once.

**Computational analysis of splicing regulatory elements**

We performed an *in silico* analysis of the six deep-intronic mutations to infer a pathogenic effect on splicing. In principle the nucleotide changes could affect Intronic Splicing regulatory elements such as enhancers (ISE) or Silencers (ISS) or create/strengthen 5’ or 3’ splice sites. However, regulatory elements generally reside within the first 200 bp of the intron and the main bioinformatics tools (i.e. Human Splicing Finder, www.umd.be/HSF/) have been developed to predict exonic elements, which confers prediction of the impact of the investigated changes with an unacceptable degree of speculation. Therefore, we focused the analysis on 5’ss and 3’ss (www.fruitfly.org/seq_tools/splice.html), which predicted that the c.64+305G>A, c.291+346C>T, c.681+152G>T nucleotide changes do not appreciably strengthen cryptic splice sites. Concerning the c.571+78G>A, c.572-392C>G and c.806-329G>A variants, the introduction of the nucleotide changes would result in the creation (c.572-392C>G) or remarkable strengthening (c.571+78G>A and c.806-329G>A) of a cryptic 5’ss (Table 2).

**In vitro characterization of the splicing variants**

Based on the bioinformatics prediction of the impact of these variants on splicing, on the number of affected alleles, on the MAF, and on identification in homozygous conditions, we selected the c.571+78G>A, c.572-392C>G and c.806-329G>A changes for further characterization. Due to the impossibility of investigating F7 mRNA processing in patients’ hepatocytes, the physiological site of F7 synthesis, or to the unavailability of fresh leukocytes as ectopic source of F7 mRNA, we exploited the expression of F7 minigenes (Figure 2A). The transfection of the pIVS6-wt minigene and splicing pattern analysis revealed correct splicing (Figure 2C, transcript 2) but also, albeit to a lesser extent, exon 6 skipping (transcript 1) and usage of the weak cryptic intronic 5’ss at position +79 that leads to partial intron retention (transcript 3G).

The splicing pattern analysis of cells transfected with the pIVS6+78A minigene showed an aberrant transcript (Figure 2C, transcript 3A) that, upon sequencing, indicated the usage of the strengthened intronic 5’ss at position +79 (Figure 2B, transcript 3A). This leads to partial intron retention resulting in a deleted and frame-shifted mRNA harboring a premature nonsense triplet at position p.201, not expected to produce a functional FVII protein. To evaluate the presence of residual FVII levels, the RT-PCR was fluorescently labeled and the amplicons evaluated by denaturing capillary electrophoresis, which ensures high sensitivity. This approach led us to identify very low levels of correct transcripts, which roughly accounted for approximately 3% of the overall transcripts (Figure 3B).

While the splicing analysis of c.806-329G>A construct in pCDNA3 did not reveal any alteration by transient transfection in HEK293T (Online Supplementary Appendix and Online Supplementary Figure S1) or in Baby Hamster Kidney cells (data not shown), the assessment of splicing pattern of cells transfected with the FVII minigene harboring the c.572-392C>G change revealed splicing abnormalities (Figure 2B). In addition to the correctly spliced mRNA, we identified transcripts arising from skipping of
exon 6, from usage of the weak cryptic 5’ss at position +79 and, most importantly, from a pseudo-exonization event (transcripts 1, 3G, and 4, respectively). More precisely, this new pseudo-exon 5b originates from the usage of a cryptic 5’ss and of a 282 bp upstream cryptic 3’ss. This finding was further strengthened by a PCR using a primer in the pseudo-exon 5b (F7PsExR), which gave rise to amplified fragments only in cells expressing the pIVS6-392G construct (Figure 2B, lower panel). The inclusion of the pseudo-exon leads to a frame-shifted mRNA with a premature nonsense codon, predicted to encode a dysfunctional FVII protein. 

Semi-quantitative evaluation of transcripts by fluorescent labeling of amplicons and denaturing capillary electrophoresis revealed that the correct and the aberrant forms are present in the relative proportion of 74% and 26%, respectively (Figure 3C), compared to 90% and 10% in the pIVS6-wt context (Figure 3A). Overall, these data demonstrate that both mutations exert their detrimental effect by impairing FVII splicing, strengthening the usage of cryptic 5’ss.

**Investigation by antisense U7snRNA**

Since the observed aberrant splicing is caused by the usage of new 5’ss, we hypothesized that masking them would weaken or abolish their detrimental role. To this purpose, we exploited variants of the U7 small nuclear RNA (U7smOPT) as potent antisense molecules to target the alternative splice sites (Figure 4A).

Co-expression of the c.571+78G>A change with antisense U7smOPT variants resulted in an appreciable rescue of splicing, as evaluated by densitometric analysis of bands upon semi-quantitative PCR. In particular, the proportion of correct transcripts, barely appreciable in untreated conditions, remarkably increased to approximately 9% or to approximately 20% of total transcripts upon co-expression of the pU7+78Ash or pU7+78A, respectively (Figure 4B, bottom).

Concerning the c.572-392G variant, co-expression of pU7-392G, designed on the cryptic 5’ss, resulted in a 3-fold reduction in aberrantly spliced mRNA containing the pseudo-exon 5b and conversely favored (1.3-fold increased) the synthesis of correctly spliced transcripts that rose from approximately 70% to approximately 90% of all forms (Figure 4B, bottom), resembling the proportion observed in the wild-type context.

Overall, these data further demonstrate the causative role of the mutations that create/strengthen cryptic intronic 5’ss.

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**Figure 2. Alternative splicing patterns associated with the c.571+78G>A and c.572-392C>G mutations.** (A) Schematic representation of the pIVS6 minigene. Mutations (blue) are reported on top. The presence of cryptic splice sites (5’ss in light blue, 3’ss in green), with related scores are indicated by arrows. Polymerase chain reaction (PCR) oligonucleotides are shown in red. (B) The schematic representation of splicing patterns is reported together with relative sequencing chromatograms of the amplicons, obtained by T7bisF7ex7R PCR (see panel A). PCR fragments were cloned before sequencing. (C) Splicing pattern analysis in HEK293T cells transiently transfected with pIVS6 wild type (wt) or with pIVS6 variants c.571+78A (+78A) and c.572-392G (-392G); the PCR with T7bisFF7ex7R oligonucleotides is reported in the upper panel: PCR with T7bisF7psExR oligonucleotides specifically designed to amplify transcripts with the pseudo-exon 5b, is reported in the lower panel. M:100 bp ladder; hd: heteroduplex.
Discussion

Uncharacterized F7 pathogenic alleles are mentioned in all patient databases, and NGS would represent a powerful tool to tackle this; however, so far, this has not been fully explored. Here, we applied this approach in 13 FVII deficient patients who were only partially characterized through conventional sequencing, and identified a panel of deep intronic substitutions as candidates to explain the reduced FVII:C levels in patients. However, as for the numerous deep intronic nucleotide changes identified by NGS and associated with inherited diseases, their pathogenic role requires experimental support.

The evidence for the pathogenicity of deep intronic variations relies on several clinical and molecular observations. The first aspect to consider is their virtual absence in databases. This led us to exclude from our selection the c.64+305G>A and c.291+846C>T variants, with a minor allelic frequency of 0.005 and 0.004, respectively. The same applies to the c.681+132G>T change, reported in dbSNP databases as rs752129277 but with an estimated frequency of <0.0004. Another aspect that could suggest the pathogenicity of the nucleotide changes is their distribution among affected relatives or non-related individuals. This was the case of the c.571+78G>A, c.572-392C>G and c.806-329G>A changes that occurred in eight, two and two unrelated FVII-deficient patients, respectively. Altogether these elements prompted us to explore the impact of these three variants on the splicing process through the expression of minigenes in eukaryotic cells, a well-proven approach used to dissect splicing abnormalities.4,13,14,42,45

Regarding the c.571+78G>A variant, the in vitro characterization demonstrated an aberrant splicing profile that was consistent with the FVII:C levels reported in patients. Interestingly, the amount of correctly spliced transcripts (approx. 3%) reflects the FVII:C levels (3% and <1%) in the two c.571+78A homozygotes (patients #15 and #90) (Table 1), which explains the asymptomatic or moderate clinical phenotypes. This finding is also consistent with the observation that the mutation co-segregated with the disease phenotype through the pedigree of patient #28, and the 50% FVII:C levels detected in the heterozygous mother.

It is interesting to note that the c.571+78G>A mutation, albeit absent from databases, occurred in eight apparently unrelated patients in our cohort. The polymorphic analysis in the homozygous patients led us to identify two different haplotypes that suggested two distinct mutational
events leading to two subsequent founder effects. In particular, one haplotype was found in patients of Maghreb origin whereas the second one was compatible with genotypes of patients of European and Lebanese origins.

Concerning the c.572-392C>G change, the aberrant splicing profile displayed approximately 70% of correctly spliced transcripts, which is not apparently consistent with the FVII:C levels (10%) observed in homozygous patient #380. However, the mutation was associated with the less frequent A2 and M2 polymorphic alleles, which have been demonstrated to halve the FVII expression when present in the homozygous state.33-37

The c.571+78G>A and c.572-392C>G mutations were also found in patients with moderately reduced FVII levels (#28, #377, #284). In patients #377 and #284, this can be explained by the fact that: i) the mutations are present in heterozygous condition; and ii) by the additional contribution of the functional polymorphisms A2 and M2. For patient #28, heterozygous for the c.571+78G>A mutation, the residual expression could arise from the allele bearing the p.Arg59Trp change, which, however, has never been characterized.

In contrast to the c.571+78G>A and c.572-392C>G variants, the in vitro splicing analysis did not reveal detrimental effects on splicing for the c.806-329G>A change. Therefore, we were unable to explain the genotype-phenotype relationship for three patients (#31, #262 and #341) who were not carriers of the c.571+78G>A or c.572-392C>G mutations. Since we have not identified other candidate pathogenic variants besides the mutations previously identified by conventional sequencing, it is tempting to speculate that the genetic defect could be in the unexplored highly repetitive rich GC region of intron 2 or in the 5’ or 3’ regulatory regions of the F7 gene.

Knowledge of the alternative splicing patterns and of the mechanisms involved offers the opportunity to design correction strategies that could have therapeutic implications.20-22 Here, we exploited variants of the U7 small nuclear RNA, the RNA component of the U7 small nuclear ribonucleoprotein that is biologically involved in histone RNA 3’ end processing.40 By changing the Sm consensus sequence of the endogenous U7snRNA, it is possible to express U7snRNA variants (U7smOPT) that no longer modulate histone processing, but can bind to RNA targets though base-pair interaction and efficiently accumulate into nucleus as snRNP.41 Therefore, by changing the 5’ tail of the U7smOPT, it is possible to target a desired RNA sequence and avoid its recognition by splic-

Figure 4. Investigation of aberrant splicing mechanisms by using antisense U7smOPT variants. (A) Schematic representation of engineered U7smOPT exploited in this study. The sequence of intronic mRNA and of the engineered 5’tail of U7smOPT with relative base-pairing is reported. The nucleotide changes identified by next-generation sequencing are indicated in bold and red, as well as the corresponding base in U7smOPT antisense sequence. (B) Splicing pattern analysis in HEK293T cells transiently transfected with plVS6 wild type (wt) and variants (c.571+78A and c.572-392G) alone or in combination with engineered U7smOPT (pU7). PCR with T7bisF-F7ex7R oligonucleotides performed at 32 cycles (top) or, for semi-quantitative evaluation, at 25 cycles (bottom). M: 100 bp ladder.
ing factors. This opportunity has been exploited to induce both exon skipping for therapeutic purposes and to dissect splicing regulatory elements by masking them. In our study, the observation that antisense U7smOPT variants masking the cryptic 5′ss were able to rescue the splicing pattern further demonstrated the pathogenic role of the nucleotide changes and provided a correction approach that has potential therapeutic implications for the c.571+78A mutation. In fact, if translated into patients, the correct transcripts rescued for this mutation (a well-represented change in our patient cohort and associated with severe forms) would account for FVII levels well beyond the therapeutic threshold.

In conclusion, the combination of NGS of the entire F7 gene with the expression of minigenes elucidated the molecular bases of FVII deficiency in ten out of thirteen FVII deficient patients, thus improving diagnosis and genetic counselling, and provided insight into a potential therapeutic approach based on antisense technology, successfully exploited in other disorders.

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