Factor V Kuwait: A Novel Mutation in the Coagulation Factor V Gene Discovered in Kuwait

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Introduction

Normal haemostasis is a balanced system which, on the one hand, prevents excessive bleeding from the site of injury while, on the other hand, it maintains blood circulation by inhibiting intravascular coagulation. Therefore, an effective haemostatic process possesses an intrinsic well-balanced regulatory system involving a number of dynamic mechanisms and chemical and physical reactions. Blood coagulation is one of these dynamic systems, in which a prominent response to an injury is recruited in the form of a series of stepwise (cascading) interactions leading to fibrin formation. This complex system involves certain proteins referred to as plasma clotting factors [1]. Among these clotting factors, factors V and VIII have enormous potential as cofactors in accelerating the activation of clotting factors X and prothrombin (factor II), respectively [2]. Factors V and VIII are high-molecular-weight glycoproteins (300–330 kDa) and are synthesized in the liver as single-chain precursors [3, 4]. Genetically, the factor V gene is present in chromosome 1q21–25, spanning more than 80,000 nucleotide bases containing 25 exons. The full nucleotide sequence of the cDNA of the factor V gene has been determined [5]. Many studies have reported cases with different point mutations at different positions in the cDNA of the factor V gene, some of which were only silent polymorphism, while a few

Key Words
Factor V Kuwait · Venous thromboembolic disorder · Restriction fragment length polymorphism

Abstract

Objective: It was the aim of this study to report a new point mutation in the clotting factor V gene in the general Arab population.

Subjects and Methods: The HR2 haplotype was tested in 288 Arabs living in Kuwait – 188 patients with venous thromboembolic disorders (VTE) and 100 healthy subjects – using polymerase chain reaction and restriction fragment length polymorphism techniques. The presence of the new mutation was verified by DNA sequencing.

Results: Two (1.06%) VTE patients had guanine instead of the wild-type adenine at nucleotide number 3935 (A3935G) of the factor V gene. This mutation caused a histidine to arginine change in amino acid number 1254 of the factor V molecule. The new mutation is termed ‘factor V Kuwait’ (His1254Arg) and was absent in the 100 healthy subjects.

Conclusion: It appears that factor V Kuwait could be a risk factor for developing VTE in Arabs. A larger study is needed to confirm this observation.
caused amino acid replacement. Factor V Leiden (G1691A; Arg506Gln) may be the most important mis-
sense point mutation in the factor V gene due to its estab-
lishment association with venous thromboembolic disor-
ders (VTE), with a high 3- to 10-fold risk in heterozygotes 
and a 30- to 140-fold risk in homozygotes [6–9]. HR2 
haplotype, which consists of 12 highly associated polymor-
phisms in the factor V gene, has been studied lately
as another possible risk factor for developing VTE [10,
11]. This haplotype has been the subject of the latest re-
search by the authors of this article, examining the preva-
ience and possible risk of HR2 haplotype in the Arab
population and patients with VTE living in Kuwait [12].
While testing for HR2 haplotype, an unexpected observa-
tion was obtained in 2 of the patients. Therefore, we in-
vestigated this observation in order to identify a new mis-
sense point mutation in the factor V gene.

**Subjects and Methods**

Two hundred and eighty-eight subjects, 188 patients with VTE 
and 100 age- and sex-matched healthy individuals, all of Arab eth-
nicity and living in Kuwait, were tested for the HR2 haplotype,
using polymerase chain reaction (PCR) and restriction fragment 
length polymorphism (RFLP) techniques according to Lunghi et al.
[10]. A full description of the above techniques has been given pre-
viously [12]. Briefly, venous blood was collected from each subject,
and DNA was extracted and purified using the conventional phe-
nol-chloroform method. For each DNA sample, PCR was per-
formed using the following two primers: forward primer 5’-C AAG 
TCC TTC CCC ACA GAT ATA-3’; reverse primer 5’-AG ATC 
TGC AAA GAG GGG CAT-3’. These primers flank a DNA frag-
ment of 703 base pairs (bp), from nucleotide number 3579 to nu-
cleotide number 4281 of the factor V gene. This fragment contains 
the R1/R2 polymorphism site (A4070G), which is one of the poly-
morphisms of HR2 haplotypes. Following the PCR procedure, the 
DNA fragments were subjected to cutting using the
RsaI restriction 
enzyme, and finally, the fragments were separated by gel elec-
trophoresis, stained with ethidium bromide, and visualized under 
the UV light.

**Results**

The results of PCR/RFLP in 186 VTE patients and 
100 normal subjects revealed the amplification of 703-bp 
DNA fragments, which were cut by the RsaI restriction 
enzyme into 492- and 211-bp fragments. Only 2 VTE 
cases gave unexpected results. The 703-bp fragments in 
these 2 cases were cut into two fragments with an esti-
mated size of 350 bp (fig. 1). Accordingly, this suggested 
a possible presence of a newly created restriction site for 
the RsaI enzyme, somewhere in the middle of the 703-bp 
fragment, thus cutting it into two halves. A review of the 
DNA sequence of the factor V gene revealed a potential restriction site for the RsaI enzyme starting at nucleotide 
3935, with the sequence ATAC. The restriction sequence 
for the RsaI enzyme is GTAC, and thus a possible point 
mutation at nucleotide 3935 (adenine to guanine) could 
create a restriction site for the RsaI enzyme. This was 
confirmed by DNA sequencing for the 2 suspected cases 
using the dideoxy-mediated chain termination technique, 
and both cases were found to have the expected mutation.
Consequently, a new restriction site was created for the 
RsaI enzyme, which cut the 703-bp fragment into 358- 
and 345-bp fragments. These two bands were very close 
to each other and were shown as one band on agarose gel 
electrophoresis (fig. 1). This new missense point mutation 
caused a histidine to arginine change in the amino acid 
code number 1254 in the factor V molecule (His1254Arg) 
which we have termed ‘factor V Kuwait’.

**Fig. 1.** Agarose gel electrophoresis showing the 703-bp PCR ampli-
scons before (b) and after (a) cutting with RsaI restriction enzyme 
in 5 cases (1–5). Cases 2 and 5 had 358- and 345-bp fragments after 
cutting with RsaI enzyme, which indicated the presence of the new 
factor V Kuwait mutation (FVK). Cases 1, 3 and 4 did not show 
the above cutting pattern, and thus did not have the new mutation.
M1 and M2 are DNA markers.
Discussion

Blood coagulation is a very efficient haemostatic process involving many factors and cofactors, which function in a very organized manner under full control to prevent bleeding from sites of injury [1]. Many genetic and acquired abnormalities can affect the clotting factors, which would result in bleeding tendencies or the development of VTE, as exemplified by factor V Leiden (G1691A; Arg506Gln), which is believed to cause VTE in carriers [6–9]. Another important example is the HR2 haplotype, which consists of a group of associated polymorphisms in the factor V gene and has been reported to be associated with VTE, although this is still uncertain [10, 11]. Other rare mutations in the factor V gene were also reported, such as factor V Liverpool (Ile359Thr) [13, 14], factor V Cambridge (Arg306Thr) [15], and factor V Hong Kong (Arg306Gly) [16], which have an additional effect in causing VTE.

In this study, a new missense point mutation in the clotting factor V gene was identified, where an adenine nucleotide was replaced by a guanine nucleotide at nucleotide position 3935 of the factor V gene sequence (A3935G; factor V Kuwait mutation). Factor V Kuwait mutation caused a change from histidine to arginine in the amino acid number 1254 of the translated factor V molecule (His1254Arg). Since changes in the amino acid sequence of proteins can cause changes in their functions, this newly discovered mutation may cause an adverse impact on the function of factor V, or it may simply have no effect. It is still not possible to estimate the consequences of this mutation in the absence of a proper knowledge of the possible effects of the above changes in amino acids.

The factor V Kuwait mutation was found to be present in 2 out of 188 VTE patients, and equally important, it was absent in the 100 normal controls. This may suggest a possible risk factor for developing VTE in carriers of this mutation. Moreover, in both cases, 1 patient had antithrombin deficiency, while the other was heterozygous for the factor V Leiden mutation. Both defects are known risk factors for VTE. It is therefore tempting to postulate that factor V Kuwait might have added to the risk of developing VTE in these 2 patients, especially since it is widely accepted that VTE disorders usually need coexistence of more than one risk factor to clinically manifest themselves. However, a larger prospective study is required to confirm any relationship between factor V Kuwait and VTE.

This new mutation has been submitted to the GenBank and registered with the accession numberAY881018.

Conclusion

The factor V Kuwait mutation, found in Arabs with VTE, is a new entity in the pool of genetic abnormalities of the clotting factor V.

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References


