Gly74Ser mutation in protein C causes thrombosis due to a defect in protein S-dependent anticoagulant function

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Summary
Protein C is a vitamin K–dependent serine protease zymogen in plasma which upon activation by thrombin in complex with thrombomodulin (TM) down-regulates the clotting cascade by a feedback loop inhibition mechanism. Activated protein C (APC) exerts its anticoagulant function through protein S-dependent degradation of factors Va and VIIIa. We recently identified a venous thrombosis patient whose plasma level of protein C antigen is normal, but its anticoagulant activity is only 34% of the normal level. Genetic analysis revealed that the proband and her younger brother carry a novel heterozygous mutation c.346G>A, p.Gly74Ser (G74S) in PROC. Thrombin generation assay indicated that the TM-dependent anticoagulant activity of the proband’s plasma has been significantly impaired. We expressed protein C-G74S in mammalian cells and characterised its properties in established coagulation assays. We demonstrate that the protein C variant can be normally activated by the thrombin-TM complex and the resulting APC mutant also exhibits normal amidolytic and proteolytic activities toward both FVa and FVIIa. However, it was discovered that the protein S-dependent catalytic activity of APC variant toward both procoagulant cofactors has been significantly impaired. Protein S concentration-dependence of FVa degradation revealed that the capacity of APC variant to interact with the cofactor has been markedly impaired. The same results were obtained for inactivation of FVa-Leiden suggesting that the protein S-dependent activity of APC variant toward cleavage of Arg-306 site has been adversely affected. These results provide insight into the mechanism through which G74S substitution in APC causes thrombosis in the proband carrying this mutation.

Keywords
Protein C, protein S, factor Va, factor VIIIa, thrombosis

Introduction
Protein C is a vitamin K–dependent serine protease zymogen in plasma which upon activation by thrombin in complex with thrombomodulin (TM) down-regulates thrombin generation by proteolytic degradation of factors Va and VIIIa (FVa and FVIIIa) (1–3). Protein C has a multi-domain structure composed of an N-terminal γ-carboxyglutamic acid (Gla) domain (residues 1–43), two epidermal growth factor (EGF)-like domains (residues 46–136), a linking peptide (residues 137–157) between light and heavy chains, an activation peptide (residues 158–169), and a C-terminal serine protease domain (residues 170–419) which contains the trypsin-like catalytic domain (4, 5). Following removal of the activation peptide by thrombin and conversion of protein C to activated protein C (APC), the non-catalytic light chain of the protease remains covalently associated with its catalytic heavy chain by a single disulfide bond (4). The anticoagulant function of APC in degradation of both FVa and FVIIIa is stimulated by protein S bound to negatively charged membranes in the presence of calcium (6, 7). In addition to its essential role in protein S-dependent regulation of thrombin generation, APC also possesses cytoprotective and anti-inflammatory properties when it binds to endothelial protein C receptor (EPCR) to activate protease-activated receptor 1 (PAR1) (8–11). The functional significance of individual domains of APC has been extensively studied. It has been demonstrated that the Gla-domain is involved in the Ca2+-dependent in-
teraction of APC with both cofactors of the anticoagulant (protein S) and anti-inflammatory pathways (EPCR) (12, 13). The N-terminal EGF domain is also believed to be required for the protein S-dependent anticoagulant function of APC (14–16). The role of the C-terminal EGF domain in the catalytic function of APC is not well known. Nevertheless, this domain is in intimate contact with the catalytic domain, thus the two domains likely constitute a single functional unit (17). Unlike its anticoagulant function, EPCR- and PAR1-dependent cytoprotective function of APC does not require interaction with protein S. The mechanism through which protein S augments the catalytic function of APC toward the procoagulant cofactors is poorly understood (18).

Protein C deficiency has an autosomal dominant pattern of inheritance and its heterozygous deficiency is associated with increased risk of venous thromboembolism (VTE) and its homozygous deficiency causes purpura fulminans, which is fatal unless treated by protein C replacement therapy (19, 20). This is in agreement with the observation that complete deficiency of protein C in knockout mice is lethal (21). More than 200 natural variants of protein C with mutations scattered at all distinct functional domains (Gla, EGF1, EGF2 and catalytic domains) have been reported in the protein C database (http://www.hgmd.cf.ac.uk/ac/gene.php?gene=PROC). In general, protein C deficiency is divided into type-I deficiency, which is characterised by equally low antigen (PC:Ag) and activity (PC:A) levels, and type-II deficiency which is characterised by only a lower activity level for APC (22). Type-II deficiency can be subdivided into type-IIa and type-IIb. Type-IIa variants show concordant reductions in PC amidolytic and anticoagulant activities because of abnormal function of the PC/APC serine protease domain, while type-IIb variants show reduced anticoagulant activity but normal amidolytic activity (23). In this study, we have identified a type-IIb protein C deficient VTE patient whose plasma PC:Ag level in ELISA and PC:A level in chromogenic assay are normal, but the PCA level in clotting assay is 34 % of that in normal plasma. By genetic analysis, we demonstrate the proband carries a heterozygous mutation (c.346G>A) in *PROC*, which leads to a Gly-74 to Ser substitution (p.Gly74Ser, G74S) on the N-terminal first EGF-like domain of protein C. We expressed this protein C variant in mammalian cells and after its characterisation discovered the variant has normal amidolytic and catalytic activity toward both FVa and FVIIa in the absence of protein S, but the anticoagulant activity of the variant was significantly impaired in the presence of the cofactor. The same results were obtained with the FVa Leiden protein, suggesting the mutation adversely affects the protein S-dependent recognition and cleavage of the Arg-306 site of FVa by APC. The results provide clinical evidence that the interaction of EGF1 of APC with protein S contributes to the anticoagulant function of the protease.

Materials and methods

**Haemostasis assays**

Routine coagulation screening assays including prothrombin time (PT), activated partial thromboplastin time (aPTT), fibrinogen (Fg), thrombin time (TT), d-dimer (DD) and fibrinogen/fibrin degradation products (FDP) were performed in all individuals under the study using an ACL-TOP automatic coagulometer (Instrumentation Laboratory, Bedford, MA, USA) according to manufacturer’s instructions. A detailed description of all haemostasis assays is presented as online Suppl. Material (available online at www.thrombosis-online.com).

**Analysis of thrombin generation in plasma**

Thrombin generation (TG) assay (Thermo Labsystems OY, Helsinki, Finland) was carried out with platelet-poor plasmas (PPP) of the proband, her affected younger and normal older brothers according to manufacturer’s instructions. The reaction was initiated with 5 pM tissue factor (TF), 4 µM phospholipids, 16.7 mM CaCl$_2$ in the absence or presence of 5 nM or 10 nM soluble thrombomodulin (sTM) (Sekisui Diagnostics, LLC, Lexington, KY, USA) as described (24). The kinetics of thrombin generation was monitored by measuring the hydrolysis of a fluorogenic thrombin substrate as described (25). The lag time (LT, min), peak height (Peak, nM), and endogenous thrombin potential (ETP, nM*min) were deduced from thrombin generation curves plotted with Thrombinscope Software, version 5.0.0.742 (Thrombinscope BV, Leiden, the Netherlands) as described (24, 25).

**Genetic analysis**

Genomic DNA was extracted from peripheral whole blood using the QIAamp DNA blood purification kit (Qiagen, Hilden, Germany) according to the manufacturer’s instructions. Detection of genetic defects of the *PROC* was carried out by directly sequencing on an ABI 3700 sequencer (Applied Biosystems, Foster City, CA, USA).

**Construction, expression, and purification of recombinant proteins**

Both wild-type (WT) and Gly-74 to Ser (G74S) substitution mutant of protein C were expressed in human embryonic kidney (HEK-293) cells as described (26). Methodologies for isolation, activation and initial characterisation of protein C derivatives and the source of plasma proteins and other reagents have been presented as Suppl. Material (available online at www.thrombosis-online.com).

**Interaction with EPCR**

The interaction of protein C derivatives with EPCR was assessed by an ELISA-based binding assay using the HPC4-tagged recombinant soluble EPCR (sEPCR) as described (27). 96-well flat bottom microtitre plates were coated with the HPC4 monoclonal antibody in TBS containing 1 mM CaCl$_2$ overnight at 4°C. Following washing and blocking of plates next day with 2 % BSA in TBS/Ca$^{2+}$, they were incubated with sEPCR (0.5 µM in TBS/Ca$^{2+}$ containing 0.1 % BSA) for 1 hour (h). The plates were rinsed and then incubated with either WT or mutant APC (7.8–500 nM) for 1 h.
After washing, a goat anti-protein C polyclonal antibody (1 µg/ml) was added and the plates were developed as described (27).

**Endothelial cell permeability assay**

The cytoprotective signalling activity of APC derivatives was monitored by a permeability assay using transformed human umbilical vein endothelial cells (EA.hy926) as described (24, 27). The cell permeability in response to thrombin (10 nM for 10 minutes [min]) following treatment with APC derivatives (20 nM for 3 h) was quantitated by spectrophotometric measurement of the flux of Evans blue-bound albumin across functional cell monolayers using a modified two-compartment chamber model as described (24, 27). Results were expressed as mean ± SD and all experiments were repeated at least twice.

**Anticoagulant assays**

The anticoagulant activity of APC derivatives was monitored both in purified and plasma-based assay systems as described (24, 26). In the purified system, degradation of both FVa and FVIIia by APC was evaluated. In the case of FVa, the cofactor (2.5 nM) was incubated with increasing concentrations of either APC-WT or APC-G74S (0–5 nM) on 25 µM PC/PS vesicles in TBS/Ca²⁺. Following 10 min incubation at room temperature, the remaining FVa activity was determined in a prothrombinase assay as described (24). Thrombin generation was monitored by an amidolytic activity assay using S2238 (100 µM). The same assay was used to monitor the inactivation of FVa by increasing concentrations of APC in the presence of protein S (110 nM) with the exception that incubation time was decreased to 1 min. The same methods were used to measure the catalytic activity of APC derivatives toward FVa Leiden in both the absence and presence of protein S.

The inactivation of FVIIia (10 nM) by increasing concentrations of APC (0–20 nM) in the absence or presence of protein S (110 nM) ± factor V (FV, 10 nM) on PC/PS vesicles (50 µM) was monitored in TBS/Ca²⁺ as described (24). Following 3–6 or 30 min incubation at room temperature, the remaining FVIIia activity was determined by an intrinsic Tenase assay as described (24). FXa generation was measured by an amidolytic activity assay using SpFXa (200 µM).

The anticoagulant activities of APC derivatives were also evaluated in normal and protein S-deficient plasma by an aPTT assay using StART 4 fibrinometer (Diagnostica/Stago, Asnieres, France). In both cases, 0.05 ml TBS containing 0–20 nM APC was incubated with a mixture of 0.05 ml of plasma plus 0.05 ml aPTT reagent (Alexin) for 5 min before initiating clotting by the addition of 0.05 ml CaCl₂ (35 mM) at 37°C as described (24).

**Molecular modelling**

The structural model of the APC Gla-EGF1 domains was built based on the x-ray crystal structures of the Gla-domain of prothrombin and active-site inhibited Gla-domainless APC (14, 17, 28, 29). The angle between EGF1 and Gla domains was taken from the x-ray structure of factor VIIa (30). In all cases, Gly-74 is fully solvent exposed and located in a loop structure in a region of the EGF1 domain involved in Ca²⁺ binding near the last helix of the Gla-domain.

Fragment mapping approach (FTMAP) was used to predict hotspot regions on the surface of the APC and protein S Gla-EGF1 regions (31). The model structures of protein S and APC were used as input for protein-protein docking experiments (14, 17, 29). Details of molecular modelling are given in Suppl. Material (available online at www.thrombosis-online.com).

**Results**

**Clinical case**

The proband (III-3) was referred to the hematology clinic for consultation because of mesenteric and portal vein thrombosis (Figure 1A). Her younger brother (III-2), father (II-2) and aunt (II-3) had also experienced bilateral lower-limb deep-vein thrombosis (DVT), but her older brother (III-1) was normal (Figure 1A). Plasma levels of protein C obtained from the proband and her affected younger brother revealed a type-IIb protein C deficiency as evidenced by normal protein C antigen and activity levels based on ELISA and chromogenic assays, but a significantly lower activity level based on the aPTT clotting assay (Figure 1C). Results of all other routine coagulation and thrombophilia screening assays were normal (data not shown). Genetic analysis identified a heterozygous missense mutation in PROC in both the proband and her younger brother, resulting in substitution of Gly-74 of protein C in EGF1 domain (exon 5 g.9698G>A) with Ser (Figure 1B). This is a novel mutation in PROC which has not been reported before.

**Thrombin generation assay**

To evaluate the anticoagulant activity of protein C in the proband’s and her affected brother’s plasma, thrombin generation assay was conducted in both the absence and presence of sTM and utilising a tissue factor concentration of 5 pM to initiate clotting. The results in the absence of sTM indicated near normal thrombin generation profiles for the proband and her younger brother (Figure 2). However, in the presence of 5 nM or 10 nM sTM, plasma from the proband and her younger brother exhibited significantly higher values of Peak and ETP of thrombin generation (Figure 2). Thus, in contrast to 90–95% sTM-mediated inhibition of thrombin generation in normal plasma, the inhibition ratio was decreased to 50–70% in both the proband’s (III-3) and her younger brother’s (III-2) plasma. These results suggest that the protein C anticoagulant activity of the mutant in plasma has been markedly impaired in both subjects carrying the Gly74Ser mutation.
Expression and characterisation of recombinant protein C-Gly74Ser

Both WT and the Gly74Ser mutant of protein C were expressed in HEK-293 cells and following purification to homogeneity, activated by thrombin in TBS/ETDA buffer as described in Materials and methods. The amidolytic activity of APC-G74S toward the chromogenic substrate SpPCa was essentially identical to that observed with WT APC (Suppl. Figure 1, available online at www.thrombosis-online.com). The time course of the initial rate of activation of protein C by thrombin in both the absence and presence of sTM and calcium suggested a normal activation rate for the mutant zymogen (Figure 3). The activation of the mutant protein C zymogen by thrombin alone (in the presence of calcium) was slightly, but reproducibly improved. These results suggest that the higher incidence of thrombosis observed in the proband is not caused by the slower rate of protein C activation by thrombin.

Interaction with EPCR and cytoprotective signalling activity

The effect of G74S substitution on the affinity of APC for interaction with EPCR was evaluated by an ELISA-based binding assay. The results suggest both APC-WT and APC-G74S interact with EPCR with similar affinities thus yielding a dissociation constant of ~30 nM for both proteases (Figure 3D). A similar affinity for the interaction of APC with EPCR has been reported in other studies (12). The results further suggested that the EPCR-dependent signalling activity of APC is not adversely affected by the G74S mutation since both APC-WT and APC-G74S exhibited similar cytoprotective activity in a thrombin-mediated permeability assay (Figure 3E). The cytoprotective activity of APC was independent of protein S for both proteases (Figure 3E).

Anticoagulant activity

The anticoagulant activity of APC-WT and APC-G74S was evaluated in both the absence and presence of protein S. The APC co-
centration dependence of FVa inactivation indicated that APC-G74S has normal anticoagulant activity toward FVa in the absence of protein S and that the rate of cofactor degradation is essentially identical to that observed with APC-WT (Figure 4A). However, the same assay indicated that the anticoagulant activity of the APC mutant toward FVa has been significantly impaired in the presence of protein S (Figure 4B). Essentially identical results were obtained in reaction with FVIIIa. Thus, the APC mutant exhibited a normal anticoagulant activity toward FVIIIa in the absence of protein S (Figure 4C), but its activity toward the cofactor was markedly impaired in the presence of the cofactor (Figure 4D). Further FVa and FVIIIa inactivation studies in the presence of increasing concentrations of protein S suggested the capacity of APC-G74S to interact with protein S has been significantly (2- to 3-fold) impaired (Figure 5A, B). The PC/PS-concentration dependence of FVa inactivation in the absence (Figure 5C) and presence of protein S (Figure 5D) suggested the impaired protein S-dependent inactivation of FVa by the APC-G74S is not due to the loss affinity of the mutant APC for interaction with negatively charged membrane surfaces since both APC-WT and APC-G74S exhibited identical apparent dissociation constant of 1 µM for interaction with the PC/PS vesicles (Figure 5C, D).

Factor V is known to act as a cofactor and make additional contribution to degradation of FVIIIa by the APC/protein S complex in the intrinsic Tenase assay (32). While the addition of FV to the FVIIIa degradation assay improved the anticoagulant activity of both APC-WT and APC-G74S, nevertheless FV did not restore the defective cofactor function of protein S, suggesting that the protein S cofactor defect with the mutant protease is independent of the cofactor effect of FV (Figure 6A, B).

The anticoagulant activity of APC derivatives toward FVa Leiden was also evaluated in both the absence and presence of protein S. The results indicate both APC-WT and APC-G74S exhibit similar anticoagulant activities toward FVa Leiden in the absence...
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of protein S (Figure 7A). However, similar to degradation of WT FVα, the anticoagulant activity of the APC mutant toward FVα Leiden was significantly impaired in the presence of protein S (Figure 7B). In FVα Leiden, the APC recognition site, Arg-506, is mutated to Gln (33, 34). Thus, these results suggest that the cofactor function of protein S in promoting the catalytic efficiency of the APC mutant toward the FVα Arg-306 site has been impaired.

Consistent with results in the purified system, APC-G74S exhibited defective anticoagulation activity in the plasma-based aPTT assay, thus prolonging the clotting time of normal plasma with an efficiency that was significantly lower than that observed with WT APC (Figure 7C). In agreement with the hypothesis that the G74S mutation has specifically affected the protein S-dependent function of APC the mutant exhibited a normal anticlotting activity in protein S-deficient plasma (Figure 7D).

The reactivity of APC-WT and APC-G74S with plasma inhibitors was evaluated by incubating the proteases with plasma at a final concentration of 20 nM followed by monitoring their residual amidolytic activities toward the chromogenic substrate SpPCa. Time course analysis indicated both APC-WT and the variant exhibit identical reactivity with plasma inhibitors at several time points examined (data not presented).
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Discussion

We demonstrated in this study that heterozygous G74S mutation in PROC is associated with DVT in the proband and two of her family members. To determine whether a molecular defect in the anticoagulant function of the protein C mutant may be responsible for the clotting defect in the proband and her family members, we expressed the protein C mutant in mammalian cells and characterised its properties in established protein C/APC assay systems. The results indicated the protein C mutant is activated normally by the thrombin-TM complex and the resulting APC mutant has normal amidolytic and proteolytic activities in all assays examined in the absence of a cofactor. However, the results in the presence of protein S indicated that the APC mutant has lost its high-affinity interaction with its anticoagulant cofactor. Thus, the protein S concentration-dependence of FVa and FVIIIa degradation by APC in the presence of increasing concentrations of PC/PS vesicles suggested that the G74S mutation weakens the affinity of APC for protein S ~2–3-fold without adversely affecting the affinity of the Gla-domain of the APC mutant for interaction with negatively charged membrane surfaces. Further support for this hypothesis was provided by the observation that both APC-WT and APC-G74S exhibited identical anticoagulant activities in protein S-deficient plasma but APC-G74S had reduced activity in normal plasma. The G74S mutation did not adversely affect the interaction of APC with EPCR and/or its EPCR-dependent cytoprotective signalling function. Noting that the Gla-domain dependent interaction of APC with either negatively charged membrane surfaces or endothelial cell surface receptor, EPCR, is required for the anticoagulant and cytoprotective function of APC (8–16), respectively, the results suggest that G74S mutation does not adversely affect the folding and/or the conformation of the Gla-domain of APC.

The mechanism by which G74S mutation impairs the protein S-dependent anticoagulant function of APC is not known. However, it is known that APC sequentially cleaves at least two bonds after Arg-506 and Arg-306 sites to inactivate FVa (35–37). It has been demonstrated that the APC cleavage of FVa at the Arg-306 site is membrane dependent (35–37). By contrast, the APC cleavage of the Arg-506 is membrane independent, but APC cleaves this site faster than that of the Arg-306 site. The slower activity of
Figure 5: Protein S-concentration dependence of FVa and FVIIIa degradation by APC derivatives. A) FVa (2.5 nM) degradation by APC-WT (○) and APC-G74S (●) (1 nM each) in the presence of increasing concentrations of protein S (from 0 to 50 nM) was analysed on PC/PS vesicles (25 µM) in TBS/Ca\(^{2+}\) for 1 min. The remaining cofactor activity of FVa was determined by a prothrombinase assay (1 nM FXa and 1 µM prothrombin for 1 min). B) Similar to A, except that FVIIIa (10 nM) degradation by APC-WT and APC-G74S (20 nM each) in the presence of increasing concentrations of protein S (from 0 to 60 nM) was analysed on PC/PS vesicles (40 µM) in TBS/Ca\(^{2+}\) for 6 min. The remaining cofactor activity of FVIIIa was determined by an intrinsic Tenase (1 nM FIXa and 100 nM FX for 1 min) as described in Materials and methods. C-D) The same as panel A, except that the PC/PS concentration-dependence of FVa (2.5 nM) degradation by APC-WT and APC-G74S (1 nM each) in the absence (C) and presence (D) of protein S (110 nM) was monitored.

APC toward the Arg-306 site is compensated by the cofactor function of protein S which has been shown to preferentially improve the cleavage rate of Arg-306 site to a greater extent than that of the Arg-506 site (36, 37). To evaluate whether or not the defective protein S binding property of APC-G74S affects the cleavage rate of Arg-306 in the absence of protein S, but the activity of APC-G74S in the presence of protein S was impaired to a similar extent as with WT FVa suggests that the cofactor function of protein S in promoting the catalytic efficiency of the APC mutant toward cleavage of Arg-306 site has been impaired. It should be noted that these results do not exclude the possibility that the cofactor activity of protein S toward recognition of Arg-506 has also been impaired in the APC mutant.

The structural basis for the weaker affinity of APC-G74S for protein S is not known. We investigated this question using several computational approaches and structural analyses of both x-ray structures and homology models. Structural data shows that Gly-74 is located near Ca\(^{2+}\)-binding residues of EGF1 of APC (5, 17, 29). The binding of Ca\(^{2+}\) to this site of APC-EGF1, immediately outside of the Gla-domain is required for the normal anticoagulant function of APC and its interaction with protein S (38). Thus, it is possible that the G74S mutation alters the Ca\(^{2+}\)-dependent affinity of APC for protein S. This site has a much higher affinity for Ca\(^{2+}\) than the Gla-domain of APC, thus the evaluation of the effect of mutagenesis on the affinity of EGF1 for Ca\(^{2+}\) was not feasible by functional assays. However, molecular modelling of Gla and EGF1 domains of APC (Figure 8A) predicts that the Gly to Ser substitution at this position should be structurally tolerated. Moreover, based on simulation data, the Ser side chain in the APC variant is expected to point away from the Ca\(^{2+}\)-binding site without affecting the interaction of EGF1 with the metal ion. Short simulations indeed suggested that a Ser at this position could be inserted without causing steric clashes or folding problems. A Gly at this site is found in APC sequences from different species but this residue is not conserved in the same position of other vitamin K–dependent
coagulation proteins such as FIX and FVII which also possess a similar high-affinity Ca$^{2+}$-binding site on EGF1 domain (5). Further, this residue is Ser in FIX and Lys in FVII, yet the EGF1 domains of both coagulation factors harbour a high-affinity Ca$^{2+}$-binding site in the same position as in APC (5). The prediction of the stability change (ΔΔG value) upon the amino acid substitution of Gly to Ser, as computed with two different software programs (39, 40), indicated the mutation is neither stabilising nor destabilising and, as such, it should be tolerated structurally. Thus, the molecular modelling data does not predict a disruptive role for Ser at this position for the ability of APC-G74S to interact with Ca$^{2+}$.

**Prediction of protein-protein interaction sites**

Mutations in the protein-protein interaction sites are known to be enriched in disease-causing missense mutations compared to other protein surface regions (41). Since Gly-74 of APC is in a fully exposed region and its substitution to Ser is predicted to be structurally tolerated, we hypothesise that this residue should be within a protein-protein interaction site area, and according to the experimental data, possibly in a region interacting with protein S. Numerous computational approaches can be used to predict protein-protein interaction sites and hotspot regions (42, 43). Here we used the fragment mapping approach, FTMAP (31), as a tool to identify hotspot regions potentially present in the interface regions between the Gla-EGF1 domains of APC and the Gla thrombin-sensitive region (TSR)-EGF1 domains of protein S (14). Two main binding hotspot regions were identified on the surface of human protein S: hotspot 1 involves a region of the Gla-domain and several residues of the TSR (Arg-49 and/or Gln-52) that have been hypothesised to be involved in APC binding (44–46); hotspot 2 involves a region at the interface of EGF1 domain (i.e. Asp-95) that has been shown to play a role in binding to APC (Figure 8B) (47). Of interest, these predicted hotspot regions are supported by results of previously reported mutagenesis studies (44–47). Using the same orientation as in Figure 8A, the two main computed APC hotspot regions are shown in Figure 8B, hotspot 1 is at the Gla-EGF1 interface near Gly-74 and hotspot 2 is located in the EGF1 domain. The protein-protein docking experiments employing the pyDock computational modelling package predicts that Gly-74 is located in the vicinity of several residues of the Gla-TSR-EGF1 regions of protein S and expected to be in the interface of the APC-protein S interaction sites (Figure 8C). This structural model is still a “low resolution” structure at this time, thus it is not possible to predict whether the Gly to Ser substitution would directly clash against some nearby protein S residues or if the substitution induces some local dynamic structural changes in this region of APC that impedes the proper interaction of APC with this region of protein S (e.g. perturb stabilising hydrogen bond networks including water molecules commonly found at protein-protein interfaces).

In summary, we have demonstrated that EGF1 Gly-74 of APC plays a key role in the protein S-dependent anticoagulant function of APC. The heterozygous Ser substitution of this residue was determined to be responsible for VTE in the proband and her affected family members. The results suggest that the G74S mutation would be most harmful under conditions where protein S levels are low (i.e. pregnancy, oral contraceptive use, etc.) (48, 49). Molecular modelling of the APC-protein S complex predicts the lack of a side chain at position 74 of the APC EGF1 domain facilitates its optimal interaction with a region in the TSR-EGF1 domain of protein S on the membrane surface. This interaction appears to be important for maintaining the active-site of APC in a topographical orientation that is optimal for efficient recognition and degradation of its procoagulant substrates, FVAs and FVIIIa, on the membrane surface.

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Author contributions
C. C., L. Y. designed experiments and performed research; B. O. V. performed molecular modelling; X. W. and Q. D. designed experiments, supervised studies conducted in Ruijin Hospital with the subjects’ plasma and contributed to writing of the manuscript; and A. R. R. designed experiments, analyzed data, wrote the manuscript and supervised the project. All authors approved the final version of this manuscript.

Conflict of interest
None declared.

What is known about this topic?
- Heterozygous protein C deficiency is associated with increased risk of venous thrombosis.
- Activated protein C (APC) functions as an anticoagulant by degrading factors Va and VIIIa.
- The cofactor function of protein S is required for the anticoagulant function of APC.

What does this paper add?
- This paper demonstrates that Gly-74 to Ser mutation on EGF1 domain of protein C causes venous thrombosis.
- The Gly74Ser mutant of APC has defect in its protein S-dependent anticoagulant function.
- This paper provides in vivo evidence that interaction of EGF1 of APC with protein S is required for recognition of procoagulant substrates on negatively charged membrane surfaces.

Figure 7: Purified FVa Leiden degradation and clotting activity of APC derivatives in protein S-deficient plasma. A) The degradation of FVa Leiden (2.5 nM) by increasing concentrations of APC-WT (○) and APC-G74S (●) was carried out on PC/PS vesicles (25 µM) in TBS/Ca²⁺ in a 96-well assay plate. Following 10 min incubation at room temperature, the remaining cofactor activity of FVa Leiden was determined by a prothrombinase assay (0.5 nM FXa and 500 nM prothrombin for 1 min) as described in Materials and methods. B) The same as A, except that the APC concentration dependence of FVa Leiden degradation was carried out in the presence of protein S (110 nM) for 1 min. C) The anti-clotting activities of APC-WT (○) and APC-G74S (●) were determined in normal plasma by an aPTT assay as a function of increasing concentrations of APC at 37°C as described in Materials and methods. D) The same as panel C except that the anticoagulant activities of the proteases were evaluated in protein S-deficient plasma.
Figure 8: Structural models of the APC and protein S Gla-EGF1 domains. A) The model of APC Gla-EGF1 domains was built based on the x-ray structures of prothrombin, Gla-domainless APC and FVIIa as described in Materials and methods. The side chains of several residues near the Ca$^{2+}$-binding residue, β-hydroxylated Asp-71, in the vicinity of Gly-74 are displayed. Some polar and negatively charged residues are shown in red while some surrounding hydrophobic and/or aromatic residues are shown in yellow. B) The fragment mapping approach, FTMAP, was used to predict protein-protein interaction sites and hotspot regions as described in Materials and methods. Two main hotspots in these regions of both proteins were identified. C) The pyDock protein-protein docking algorithm was used to propose about 100 different models of the APC-protein S complex as described in Materials and methods. Structural analysis and integration of previously reported mutagenesis data, predicted hotspot data and results of the present analysis of the G74S variant led to the shown structural model. The omega loops (in orange) of the two proteins are shown pointing toward a membrane surface.
References


