Thrombosis Associated with Protein S Deficiency: A Case Report and Literature Review

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Abstract
We have reported a rare case of deep vein thrombosis (DVT) of both legs due to protein S (PS) deficiency in a 28-year-old female. She was admitted to intensive care unit (ICU) on the basis of lower limb swelling due to thrombosis as confirmed by Doppler ultrasonography. Immunoassay by ELISA demonstrated markedly reduced total PS levels in her serum, which is likely due to a defect in the PROS1 gene. The patient was successfully treated with heparin and warfarin. This case adds to the growing evidence that PS deficiency is one of the rare cause of DVT, and also raises awareness about prophylactic treatment especially in those with PROS1 gene abnormalities or other risk factors.

Keywords: Thrombosis, protein S deficiency, PROS1 gene, deep vein thrombosis.

Introduction
Protein S (PS) is a vitamin K-dependent serum protein, of 75kDa, that has a pivotal role in the anticoagulant system[1]. Its major function is as a cofactor to facilitate the action of activated protein C (APC) on its substrates, activated factors V (FVa) and (FVIIIa)[2]. Deficiency of PS is rare [3], but has been associated with deep vein thrombosis (DVT) [4], thrombophlebitis [5], and pulmonary embolism [6]. We described a case of young female presented with deep venous thrombosis (DVT) in both legs due to PS deficiency.

Case Report
A 28-year-old normotensive and normoglycemic female presented in the emergency department (ER) of the Capital Development Authority (CDA) Hospital, Islamabad, Pakistan, with sudden onset painful swelling of both legs for last seven days. Her past medical and surgical history were unremarkable, and she belongs to low socioeconomic status. Her menstrual cycle was regular, and her two children were delivered by caesarean section without any bleeding and thrombotic complications during or after delivery. She used oral contraceptive pills (OCP) intermittently, and her family history was unremarkable.
The patient physical examination showed 80kg (176lbs) and 4ft 7in (140cm) weight and height, respectively. There were no features of jaundice, pallor, leukonychia, clubbing, edema, lymphadenopathy. Her vital signs were within normal limited.

Both legs were swollen with right and left calf circumferences of 48.5 and 47.5 cm, respectively, and the overlying skin was congested as shown in Figure 1. The respiratory, gastrointestinal, cardiovascular, and central nervous system examinations were all normal. Thrombosis was detected by Doppler ultrasound exam on both lower limbs, right femoral vein and left popliteal vein.

Investigations revealed normal complete blood count (CBC). Complement levels, antinuclear antibody (ANA), anti-smooth muscle antibody (SMA), anti-double stranded DNA (dsDNA), anti-cardiolipin antibodies (ACA), anti-mitochondrial antibodies (AMA) and rheumatoid factor (RF) were found to be undetectable which rules out the possibility of autoimmune diseases. Bleeding time, clotting time, activated partial thromboplastin time (APTT), and prothrombin time (PT), serum protein C were also within normal limits, while serum PS test was on lower limits 24 U/dl (normal > 63 U/dl for females).

The diagnosis of PS deficiency was made on the basis of low PS levels. The patient was admitted, and started on low molecular weight heparin (LMWH) along with warfarin. Prothrombin time (PT) and INR (International Normalized Ratio) were monitored. The treatment goal was to maintain INR range from 2-3, and prevent potential complications.

Patient was stable and discharged on the tenth day of hospital stay, on warfarin 10mg once a day with INR 2.6; and advised to follow-up monthly to monitor INR, CBC and serum PS level.

**Discussion and Literature Review**

The present case adds to the growing evidence that PS deficiency is associated with thrombosis [3,4]. This is perhaps not surprising since PS has a crucial role in the control of normal blood clotting via inhibition of coagulation [1,2]. It does this through a feedback mechanism in which it activates activated protein C (APC) eventually leading to proteolysis and breakdown of coagulants and products [10]. In addition, PS has anticoagulant properties even in the absence of APC since it is able to inhibit thrombin production [11], and a role in the tissue factor pathway inhibitor (TFPI) system has been proposed [1,12]. The deficiency may be either acquired [13], or hereditary [3,4]. It is detected in about 1:700 of the general population although its frequency is markedly increased to 3-6% in subjects with thrombophilic disorders, further indicating its pivotal role in thrombosis [4,7]. Acquired forms can occur as a result of a number of factors such as low vitamin-K levels, liver disease, HIV infection, sickle-cell anemia, and in warfarin therapy [13-15]. Inheritable forms are rare, with an incidence of about 0.03% and usually due to large deletions in the PS gene, PROS1 [16], although other defects such a frameshift [17], and nonsense mutations have been described [18]. Since PS deficiency is autosomal-dominant (AD), even one allele may...
still be wild type, heterozygous individuals have an increased risk of thrombophilic events and their PS levels are diminished [16]. Without molecular analysis of the relevant gene, it is difficult to decipher whether any observed PS deficiency is acquired or inheritable[19]. Unfortunately, due to the lack of facilities in the hospital, it was not possible to assess the PROS1 gene in the present case. Nevertheless, there are some differences in serum biochemistry and phenotypic presentation between the acquired and inheritable forms [16] and the latter is usually associated with DVT rather than arterial thrombosis [20]. In our case patient had DVT in both legs revealed by doppler ultrasound examination. In addition, deficiency of PS predisposes to recurrent venous thromboembolism (VTE) [4,21]. Thus, although there was no family history of thrombosis, the patient's history of the previous DVT is consistent with PS deficiency due to a possibly inheritable gene defect. In support of this, no factor associated with acquired PS deficiency was found.

The diagnosis of PS deficiency can be achieved by either assessing biological activity or immunoassay [19]. Immunoassays can measure both free and bound forms. Careful evaluation is required to interpret the results correctly since many factors can influence serum PS levels including other diseases, pregnancy, use of oral contraceptives, insufficient vitamin-K intake, and medication with vitamin-K antagonists activity [3,13,15].

Three broad types have been described in the literature [22,23]. Type-I – is characterized by a decrease in both free and total PS. Type-II – represents functional deficiency, since total free and bound PS are normal. Type-III – occurs when the free is low but total PS is normal. An ELISA for total PS was used to monitor the present case and concluded that the patient suffered from the type-I form of PS deficiency.

Because of the associated increased risk of thrombophilic events, PS deficiency should be clinically addressed [24]. This could include the use of prothrombin complex concentrates [25]. Oral contraceptives are contraindicated because of their lowering effects on PS and association with thrombosis [9]. In the event of presenting thrombophilic events, the initial treatment consists of unfractionated heparin or low molecular weight heparin (LWHM) in conjunction with warfarin until an INR of 2.0–3.0 is reached on two consecutive days [21,26]. Warfarin should be administered for at least 5 days to prevent skin necrosis, a rare adverse effect that occurs during early warfarin treatment in patients with PS and protein C deficiency[15]. Warfarin treatment should be considered for up to 2 years, and in some cases for life-long [21].

Conclusion
We reported a rare cause of a deep vein thrombosis of both legs due to protein S deficiency in a 28-year-old female. Clinicians should have a high index of suspicion of such rare causes of thrombophilia, when evaluating DVT to prevent life-threatening complications.

Author Contributions
AB Bhatti clinically examined the patients and diagnosed protein S deficiency. F Ali supervised the laboratory biochemical studies. SA Satti helped confirm the clinical and laboratory diagnosis.

References


