INTRODUCTION

Protein C is a vitamin K-dependent protein which was discovered by Stenflo in 1976. It was composed of a light and heavy chain linked by interchain sulfide bridges (Figure 1). In its activated form protein C inactivates Va and VIIIa and stimulates fibrinolysis. In this process, protein S serve as an important factor for activated protein C. Furthermore, excess protein S drives cancer cell proliferation and cell survival through oncogenic receptor Axl (Anexelektro). We determined ranges of protein C both in healthy individuals and distinct hospitalized patients.

Methods: A total of 100 patients with different diagnostic diseases and 50 healthy individuals were included in their plasma protein C determination. A rabbit antibody against human protein C was used for the quantitative estimation of plasma protein C antigen by using rocket immunossay.

Results: In healthy individuals protein C antigen(PC:Ag) ranges 0.6439-1.4752 µg/ml. The mean coefficient of variation (CV) of length of rocket was calculated to be 12.45%. PC:Ag within laboratory variation was 11.47%. Plasma protein C antigen was destroyed at 56°C for 30 minutes, whereas no significant decrease of protein C was found at 4°C refrigerator for one week.

Conclusion: The results showed that plasma protein C antigen was considerably high in 22 diabetes mellitus. On the other hand, the PC:Ag was significantly decreased in 19 liver cirrhosis (p<0.001) and was positively correlated with serum albumin levels (p<0.05). In 20 acute leukemias, on the average, there was slightly lower values in PC:Ag, and accompanied with significant decrease of PC:Ag in 5 FAB M5 subtype and in 9 hyper- leukocytes acute leukemias. However, the 3 acute promyelocytic leukemia (APL) with overt laboratory picture of DIC(disseminated intravascular coagulation) had protein C concentration no lower than the remaining 2 patients with infectious DIC, which suggested the coagulopathy in APL might be due to mechanisms different from other forms of DIC.

Keywords: Antigen, clinical determination, plasma protein C, rocket immunossay.

ABSTRACT

Objective: Protein C, a vitamin K-dependent coagulation factor, is involved in blood coagulation. Activated protein C inactivates Va and VIIIa and stimulates fibrinolysis. In this process, protein S is an important factor for activated protein C. Furthermore, excess protein S drives cancer cell proliferation and cell survival through oncogenic receptor Axl (Anexelektro). We determined ranges of protein C both in healthy individuals and distinct hospitalized patients.

Methods: A total of 100 patients with different diagnostic diseases and 50 healthy individuals were included in their plasma protein C determination. A rabbit antibody against human protein C was used for the quantitative estimation of plasma protein C antigen by using rocket immunossay.

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Keywords: Antigen, clinical determination, plasma protein C, rocket immunossay.
Griffin et al.\textsuperscript{17} reported the first family with a congenital protein C deficiency. Bertina and Broekmans et al.\textsuperscript{4,18,19} reported the first Dutch family with a congenital protein C deficiency. Using an immunologic assay for protein C\textsuperscript{20}, we carried out the measurement of protein C in patients with clinical various diseases especially bleeding diathesis.

**MATERIALS AND METHODS**

**Blood collection:**
Venous blood was collected in 1/10 volume of 0.11 M sodium citrate. Plasma was prepared by centrifugation of blood at 3000 rpm for 10 min. Platelet free plasma of 100 patients and 50 healthy volunteers were stored at -20°C for the assessment of protein C levels. Three tubes with rabbit anti-protein C serum were kindly supplied by Prof. RM Bertina, Leiden University Hospital, The Netherlands. The period of experimental procedure was completed in early 1986-89. **Procedures:**

\textbf{i. Preparation of antibody-containing agarose gels.}

A 12.5 ml of 1% indubiose A37 or commercial agarose solution is obtained by boiling in gel buffer (PH8.8 Tris-BB or Tris-BB-EDTA) during continuous stirring. The solution is allowed to cool to 55°C, and a 60 µl amount of rabbit protein C antiserum and a 0.05ml of 0.11 mol/l sodium citrate solution was added during careful mixing. A 100 x 60 x 2 mm gel is prepared on small glass plate by pouring antibody-containing agarose (50 ~ 55°C) into a coated glass plate placed on a horizontal table. After 10 minutes the solidified gel is placed in a refrigerator (4°C) in a moist atmosphere. The gel should harden before wells can be punched.

**ii. Punching out the wells.**

Wells are punched out with a gel puncher, using a template.

**iii. Application of the samples.**

A row of wells (about 4 mm in diameter) with their centers at least 5 mm apart is punched in the gel. After appropriate dilution the samples (13 µl) are deposited in the wells. The ionic strength of the samples should preferably be approximately the same as that of the electrophoresis buffer. Subsequently, samples are applied after the glass plate has been placed on the electrophoresis apparatus and a low voltage has been applied (0.5 V/cm). Standards in suitable dilutions are deposited in the middle holes of the row.

**iv. Electrophoresis, staining and drying.**

Electrophoresis is performed at 120V (4–6 mA/plate) at room temperature 12–30°C for 20 hours overnight.

**v. Calculation of the test results.**

The length of rocket immunoprecipitate was measured by using a ruler, which is linearly correlated to the amount of antigen. Results (including normal controls and patients) are plotted on log-log paper (length of rocket in mm versus percentage antigen), based on the construction of the standard curve.

**Table 1: Plasma protein C antigen value distribution among age groups**

| Age groups | No | Protein C antigen(µ/ml) mean±SD | Range |
|------------|----|--------------------------------|-------|
| 18–25      | 9  | 1.0505±0.225                  | 0.6439–1.3588 |
| 26–30      | 11 | 1.0623±0.2066                  | 0.6870–1.4269 |
| 31–40      | 16 | 0.9767±0.1040                  | 0.8047–1.1649 |
| 41–50      | 13 | 1.1268±0.1841                  | 0.8785–1.4298 |
| 51         | 1  | 1.4752                         |       |
| Total      | 50 | 1.0579±0.1886                  | 0.6439–1.4752 |

*Data are the representation of Mean±Standard deviation without significant difference of mean for the different age groups (p>0.5).*

**RESULTS**

**Protein C antigen in healthy individuals**

Protein C antigen was measured in the plasma of 50 healthy individuals who contributed to our pooled normal plasma. A mean protein C antigen of 1.0578 µ/ml was calculated with a SD of 0.1886 µ/ml; individual protein C values ranged from 0.6439 µ/ml to 1.4752 µ/ml. The distribution of age on the protein C antigen concentration was evaluated in 50 healthy individuals. The results are summarized in Table 1, which indicated that no statistical differences in protein C antigen were found for the different age groups (p>0.5). Also, no difference was observed between men (mean 1.0613 µ/ml) and women (mean 1.0550 µ/ml). Data from those in Table 1 are the representation of Mean±Standard deviation without significant difference of mean for the different age groups (p>0.5).
The qualitative evaluation of protein C antigen determination

a) For investigation, to perform the consecutive numerous assays on mixed pooled normal plasma as standard curve, the mean coefficient of variation (CV) of length of rocket was calculated to be 12.45% (Table 2).

b) To estimate the within laboratory variation, 4 different plasma with PC:Ag level 0.9695, 0.5583, 0.2894 and 0.0581 µ/ml were measured by EIA at different gel plate, each plasma sample was measured twice. The average within laboratory precision (CV) was calculated using the following formula:

\[
CV_p = 100 \times \frac{\sum (X_i - \bar{X})^2}{nX} / 2n
\]

Where, \(X\) is the calculated PC antigen level, \(X_1 - X_2\) is the difference in protein C antigen level observed at different gel plate, and \(n\) refers to the numbers of assay. Using this formula the CVp for 4 different plasma could be calculated to be 6.69%, 18.03%, 4.2% and 15.7% respectively, the total mean CVp 11.16%. Meanwhile, another 3 plasma with low (0.1841 µ/ml), middle (0.9799 µ/ml) and high (1.3098 µ/ml) PC antigen level, and each plasma was measured 5 times by EIA at the same gel plate. The calculated mean CVs was 2.64%. The total coefficient of variation (CVl) in the calculated mean PC antigen levels within laboratory consisted of PC antigen level by EIA detection at the same gel plate variation(CV within a batch, Batch CV) and between different gel plate variation (interassay CV). Based on the formula:

\[
CV_l = \sqrt{CV_s^2 + CV_p^2}
\]

CVl in PC antigen levels within laboratory was calculated to be 11.47%.

c) Influence of PC antigen level in electrophoresis buffer with or without EDTA. To determine whether electrophoresis buffer with or without EDTA could influence the peak height of PC antigen assay, pooled normal plasma was a dilution of 1:1, 1:2, 1:4 and 1:8, no significant difference in results obtained by EIA in the presence of EDTA-laurell method or without EDTA from the electrophoresis buffer at 4°C. When normal plasma was kept at 56°C for 24, 12, 6, 2.5, 1 hour and 30 minutes respectively, no rocket-like immunoprecipitate was seen, which indicated that plasma protein C antigen could be destroyed at 56°C for 30 minutes. After plasma frozen at 4°C for 1, 3, 6, 9, 12, and 17 days, by using EIA assay, protein C antigen level could be measured at 1.0405, 0.9779, 1.0560, 0.7008, 0.8774 and 0.8950 µ/ml. The results suggested no significant decrease of protein C antigen when stored at 4°C refrigerator for one week.

d) Linear correlation and regression coefficient analysis. In this current study, protein C antigen concentration was measured according to Laurell’s rocket immunoelectrophoresis, and further Prof. Bertina’s modification. Electro-immunoassay (EIA) or rocket immunoelectrophoresis is a rapid and producible method for identification and quantitation of even minor amounts of proteins. When electrophoresis of an antigen leave the wells and enter the agarose gel containing the corresponding antibody, and eventually the antigen-antibody complex precipitates. In particular, a ‘rocket-like’ shape is seen. The majority of the antibody-antigen precipitate is indeed at the head of the rocket. More detail protocol, when 3 various concentration of PC:Ag antibodies (60 µl, 75 µl, 90 µl) were mixed with 12.5 ml of 1% indubiose A-37 agarose solution in a gel plate, consequently, pooled diluted normal plasma was applied for EIA. Three linear regression equation regarding protein C antigen could be drawn (Figure 2), and no statistical difference in results was obtained by comparative analysis of linear regression between each other, respectively (p>0.05). According to Prof. RM Bertina experience and our laboratory condition, an addition of 60 µl amounts of rabbit protein C antiserum was better available in preparation of antibody-containing agarose gels. The head of the rocket is clear, higher peak, save the amounts of antibodies and easy to be measured by a ruler. It has been found that a good correlation between the peak height and protein C antigen level. After a consecutive 18 assays, the linear regression equation:

\[
Y=1.093X + 0.5621, r=0.9887 \quad \text{(Figure 2)}
\]
Table 3: Experimental data of plasma PC:Ag determination in patients and normal healthy subjects.

| Groups                        | No | Mean±SD (µ/ml) | Range (µ/ml) | P    |
|-------------------------------|----|----------------|--------------|------|
| DM                            | 22 | 1.4238±0.2556  | 0.0834~1.8434 | <0.001 |
| PIHS                          | 15 | 1.2054±0.4930  | 0.5163~2.2525 | <0.05 |}
| Uremia                        | 10 | 1.1387±0.2512  | 0.7787~1.5507 | >0.2  |
| Thrombotic diathesis          | 9  | 1.1166±0.2699  | 0.7637~1.6893 | >0.2  |
| Acute leukemias               | 20 | 0.9349±0.3388  | 0.3971~1.9220 | >0.05 |
| DIC: APL                     | 3  | 0.8707         | 1.0019       | 1.2445 |
| Infectious diseases           | 2  | 0.2328         | 0.1646       |       |
| Liver cirrhosis               | 19 | 0.5501±0.2536  | 0.1436~1.0883 | <0.001 |
| Viral liver cirrhosis         | 14 | 0.5068±0.2514  | 0.1436~0.9291 |       |
| PC deficiency                 | 1  | <0.01          |              |       |
| Healthy individuals           | 50 | 1.0578±0.1886  | 0.6439~1.4752 |       |

The results are Mean±Standard deviation with significant difference of mean at P<0.05, and show where there is significant (P<0.05) difference in the DM and liver cirrhosis groups when compared to healthy individuals. DM: diabetes mellitus. PIHS: Pregnancy-induced hypertension syndrome. *including 1 nodular panniculitis and 1 septic shock.

Protein C antigen in distinct diseases

Diabetes Mellitus (DM)
In a group of 22 patients with diabetes mellitus (DM) an average value of protein C antigen was 1.4238 µ/ml, which is considerably higher than the average value in healthy individuals (1.0578 µ/ml). Among 8 DM, the protein C antigen level was over 1.4752 µ/ml. The increased plasma protein C antigen was in turn diabetic ketoacidosis (n=4, 1.5351 µ/ml, range 1.3516 ~ 1.8434 µ/ml), diabetic angiopathy (n=5, 1.4551 µ/ml, range 1.1888 ~ 2.0894 µ/ml), DM without vascular complication (n=9, 1.4403 µ/ml, range 1.0834 ~ 1.8926 µ/ml) and diabetic ketoacidosis (n=4, 1.2352 µ/ml, range 1.1955 ~ 1.2663 µ/ml). Plasma PC:Ag level was positively correlated with urine glucose level (r=0.9581, p<0.001).

Pregnancy-Induced Hypertension Syndrome (PIHS)
In 15 pregnancy-induced hypertension syndrome (PIHS), 10 severe preeclampsia were observed, and a significant increase of protein C antigen (PC:Ag) value (1.4673 µ/ml, range 0.7781 ~ 2.2525 µ/ml), while in 2 moderate PIHS the PC:Ag has normal limit range (0.7787,0.9481 µ/ml respectively), another 3 mild patients with PIHS had decreased plasma PC:Ag level(0.5603 µ/ml). Intriguing, repeat PC:Ag level was recovered to normal value (1.1455 µ/ml) in one mild patient with PIHS after partum. Plasma protein C antigen was positively correlated with urine protein (r=0.9518, p<0.001).

Uremia
In a group of 10 uremia there were no significant changes of protein C antigen concentration when compared to normal controls (1.1387 vs 1.0578 µ/ml, p>0.2)

Thrombotic diathesis
To screening congenital protein C deficiency, we detected 6 patients with cerebral haemorrhagic infarction and 3 deep venous thrombosis, no PC deficiency was found in this study.

Liver cirrhosis
19 patients with liver cirrhosis included viral liver cirrhosis 14 cases and other 1 primary biliary cirrhosis, 1 schistosomal cirrhosis, 1 alcoholic liver cirrhosis and 1 primary hepatocellular carcinoma, respectively. The average value of protein C antigen was 0.5501 µ/ml (range:0.1436 ~ 0.9291 µ/ml), which was significantly lower than the average normal controls (1.0578 µ/ml). Moreover, 13(68.4%) liver cirrhosis the protein C antigen was below the lower limit of normal control (range: 0.1436 ~ 0.6188 µ/ml); 14 patients with viral liver cirrhosis the protein C antigen was 0.5068 µ/ml. Three severe liver cirrhosis had its the lowest value of protein C antigen 0.1436, 0.1846 and 0.1919 µ/ml respectively. The PC:Ag level was 1.0883 µ/ml in one liver cirrhosis complicated with diabetes mellitus. Plasma PC:Ag level was positively correlated with serum albumin value(r=0.9680, p<0.05).

Leukemias
In 20 acute leukemias, there was slightly decreased in protein C antigen level (0.9340 vs 1.0578 µ/ml), but no statistical difference was found between acute leukaemia and normal controls. When further analysis of acute leukemia with different cytological subtypes morphology according to FAB classification, it was shown that there was remarkably decrease of PC:Ag(0.8879 µ/ml, p<0.05) in 5 FAB M5 subtype. The decreased PC:Ag level(0.7833 µ/ml) was also found in 9 hyper-leukocytes acute leukemias (WBC 23.6~280 x109/l). Intriguing, there were no significant decrease in terms of protein C antigen between 3 acute promyelocytic leukaemia (APL) with DIC, whereas 2 patients with infectious DIC (1 nodular panniculitis and 1 septic shock) the average value of protein C antigen was 0.2328 and 0.1646 µ/ml respectively.

Protein C deficiency plasma
1 vial protein C deficient plasma which provided by RM. Bertina was used to measure the protein C antigen < 0.01%.

DISCUSSION

This underlines the need for each laboratory to construct its own reference values for PC antigen in the laboratory determination of different methods. We setup successfully the electroimmunoassay (EIA) of the protein C antigen with the laurel method. Based on our laboratory condition, the mean PC:Ag level in 50 healthy individuals was 1.0578±0.1886 µ/ml, which was slightly higher PC antigen levels than that of Bertina's results. To evaluate the performance and specificity of PC:Ag assays, the ICTH subcommittee
on protein C organized an international collaborative study in which 13 lymphilized plasma samples were distributed among 17 different laboratories. Three different types of methods has been used: ELISA's using polyclonal anti-protein C IgG (n=9), electro-immunoassays (n=10) and other methods (three laboratories used monoclonal antibodies against protein C, one laboratory performed a RIA with affinity purified polyclonal antibodies against protein C and one laboratory used an immunoradio- metric assay of protein C antigen). No statistically significant results were obtained with these three different methods. ELISA, RIA and IRMA methods were found to be more sensitive than the electro-immunoassay (ELIA). The mean coefficient of variation (CV) for protein C antigen in various plasma was calculated to be 22%, with CV within laboratory variation 11.7% and between laboratory variation 17.8%. In this current study, the mean CV in the construction of standard curve for protein C antigen was 12.45%, with CV within laboratory 11.47%. The results indicated that the protein C antigen measurement was specific, practice and available in clinical useful. The measurement of plasma PC:Ag has become relevant in clinical medicine since the important demonstration that hereditary protein C deficiency is linked to a history of recurrent venous thrombosis22. PC is found to be low in a number of acquired condition such as during the postoperative and neonatal period,23 DIC21,23,24, liver disease,21,23,25 and oral anticoagulant treatment (eg. warfarin)4,17,26. Elevated PC levels, on the other hand, have been detected in a number of unrelated condition, such as ischemic heart disease, diabetes, late pregnancy,27 and in women taking oral contraceptive with a high estrogen content28.

Type 2 Diabetes mellitus is characterized by vascular complication with accelerated micro and macro vascular thrombotic disease. A hypercoagulable state is appreciated in this disease. In this study, PC:Ag levels of patients especially in ketoacidosis and diabetic angiopathy were significantly higher than that of the control group (P<0.001). The results were consistent with the reports by Vignot29, Takahashi30, Saito31, and Garcia32. Moreover, evaluation of protein C and protein S levels in patients with DM receiving well-managed T2DM (type 2 DM) had significantly higher antigen levels and activity of PC(5.78 vs 4.64 µU/ml), PS and ATIII compared to those that were poorly-managed. The decreased PC antigen was observed in type 1 diabetes mellitus33,34. This may reflect the possibility that diabetic ketoacidosis (or angiopathy) and hypercoagulate may cause endothelial injury, which lead to elevated levels of thrombomodulin(TM) and compensatory increase the PC synthesis by hepatocytes. Patients with the nephrotic syndrome and uremia are at increased risk of developing renal venous and arterial thromboembolism. In the measurement of PC:Ag levels in 10 uremia, there was no significant difference compared to normal controls. Cosio35 measured the plasma concentration of protein C in 17 patients with severe proteinuria, the results was not significantly difference between the healthy subjects and 14 patients with chronic renal insufficiency (CRI). But protein C was elevated in 71% of patients with proteinuria, and was inversely correlated with serum albumin concentrations. Demaschell36 found that PC functional activity and the antigen level were normal in 30 uremic patients before dialysis. Data of current study are consistent with the previous observation. In this current study, PC:Ag levels in 19 different types of liver cirrhosis were significantly lower than those in healthy individuals (0.5501 vs 1.0578 µU/ml, p < 0.001). 68.4% of liver cirrhosis had PC:Ag level below the lower limit of normal controls. Moreover, plasma PC:Ag level was positively correlated with serum albumin values. In earlier study, Griffin37 reported that eleven of 15 patients with clinically severe liver disease (9 with positive FDP, 6 with negative FDP) has decreased plasma protein C levels. Mannucci21,22 also found that in 58 chronic liver diseases protein C levels were lower than 60 healthy subjects, the decreased degrees roughly proportional to the severity of the disease. These findings indicated that protein C deficiencies occur the dysregulation of hepatic PC synthesis, and might play a role as a predictor index in liver diseases38.

Acute leukemias varied in PC antigen level. In this current study, mean PC antigens in 20 acute leukemias were slightly lower than that of the normal individuals. Among FAB subtypes, the decreased PC:Ag concentration were found in those with M5 subtype and hyperleukocytic acute leukemias. Moreover, the PC:Ag had no lower in a limited 3 patients with APL complicated by DIC, which suggested the coagulopathy in APL might be due to mechanisms different from other forms of DIC such as infectious disease (eg. septic shock)39,40. The results was similar to previous studies by Griffin et al.,41 and Rodeghiero et al.25. In clinics, two types of hereditary protein C deficiency can be recognized42,43. In patients with type I protein C deficiency, the plasma concentration of protein C activity and/or protein C antigen will be below the lower limit of the normal range, while the ratio between protein C activity and protein C antigen is still within the normal range. A type I protein C deficiency inhibits as an autosomal dominant trait. In type II protein C deficiency, the patients protein C antigen will be within the normal range, while the protein C activity and the ratio between protein C activity and protein C antigen are below the lower limit of the normal range. In both types of protein C deficiency the plasma concentration of other vitamin K dependent coagulation factors should be within normal range. The presence of I vial PC deficiency plasma which was provided by RM Bertina was the lower than 0.01 µ/ml, and which was consisted with the criteria of type I severe protein C deficiency41. In Dutch family, more than 80 patients from 30 unrelated families have been identified with a type I protein C deficiency. Individual protein C values may vary between 0.30 and 0.76 µU/ml44,45. Up to now, only five patients with a type II protein C deficiency have been reported46. It seems likely the main prevalence of clinical type I protein C deficiency. Like protein C deficiency, familial protein S deficiency is also reported47,48. At present, some case reports and case
series suggest that protein C concentrations may improve the outcome in patients with congenital or acquired protein C deficiency. The use of protein C concentrates in adult septic shock is also testable.

**AUTHOR’S CONTRIBUTION**

All authors have worked equally in this work.

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**CONFLICT OF INTEREST**

No conflict of interest associated with this work.

**REFERENCES**

1. Stenflo J. A new vitamin K-dependent protein. Purification from bovine plasma and preliminary characterization. J Biol Chem 1976; 251:355-363. PMID: 1245477
2. Kisiel W. Human plasma protein C. Isolation, characterization and mechanism of activation by a-thrombin. J Clin Invest 1979; 64:761-769. https://doi.org/10.1172/jci109521
3. Broekmans AW and Bertina RM. Protein C In: L. Poller, ed., Recent advances in blood coagulation, number four. Churchill Livingstone, Edinburgh 1985; 117-137. https://doi.org/10.1016/S0165-5656(88)50003-7
4. Bertina RM, Broekmans AW, van der Linden LK, et al. Protein C deficiency in a Dutch family with thrombotic disease. Thrombosis and Haemostasis 1982;48:1-5. PMID: 6897135
5. Bertina RM. An international collaborative study on the performance of protein C antigen assays. Report of the ICTH subcommittee on protein C. Thromb Haemost 1987;57(1):112-117. PMID: 3590075
6. O'Bryan JP, Frye RA, Cogwell PC, et al. Axl, a transforming gene isolated from primary human myelob leukemia cells, encodes a novel receptor tyrosine kinase. Mol Cell Biol 1991; 11(10):5016-31. https://doi.org/10.1128/mcb.11.10.5016
7. Nuzzo S, Catuogno S, Capuzzo M, et al. Axl-targeted delivery of the oncosuppressor miR-137 in non-small-cell lung cancer. Molecular Therapy: Nucleic Acids 2019;17:256-263. https://doi.org/10.1016/j.omtn.2019.06.002
8. Lingeve RMA, Keating AK, Earp HS, et al. TAM receptor tyrosine kinases: biologic functions, signaling, and potential therapeutic targeting in human cancer. Adv Cancer Res 2008;100:35-83. https://doi.org/10.1016/S0065-230x(08)00002-x
9. Burrstyn-Cohen T, Heeb MJ, Lemke G. Lack of protein S in mice causes embryonic lethal coagulopathy and vascular dysgenesis. J Clin Invest 2009;119: 2942. https://doi.org/10.1172/jci39325
10. Wimmel A, Rohner I, Ramaswamy A, et al. Synthesis and secretion of the anti-coagulant protein S and co-expression of the Tyro3 receptor in human lung carcinoma cells. Cancer 1999; 86:439. https://doi.org/10.1002/(SICI)1097-0424(19990701)86:1%3C43::AID-CNC2%3E3.0.CO;2-d
11. Abboud-Jarrous G, Priya S, Mairon A, et al. Protein S drives oral squamous cell carcinoma tumorigenicity through regulation of Axl. Oncotarget 2017; 8(8):13986-14002. https://doi.org/10.18632/oncotarget.14753
12. Brand TM, Iida M, Stein AP, et al. Axl is a logical molecular target in head and neck squamous cell carcinoma. Clin Cancer Res 2015;21(11):2601-12. https://doi.org/10.1158/1078-0432.CCR-14-2648
13. Lee N, Jang WJ, Seo JH, et al. Deoxy-D-glucose-induced metabolic alteration in human oral squamous SCC15 cells: involvement of N-glycosylation of Axl and Met. Metabolites 2019:9:188. https://doi.org/10.3390/metabo9090188
14. Wilmes S, Hafer M, Vuorio J, et al. Mechanism of homodimeric cytokine receptor activation and dysregulation by oncogenic mutations. Science 2020; 367:643-652. https://doi.org/10.1126/science.aaw3242
15. Zhu G, Saboor-Yaraghi AA, Yarden Y, Santos J, Neil JC. Downregulating oncogenic receptor: from bench to clinic. Hematol Med Oncol 2016;1(1):30 – 40. https://doi.org/10.15761/HMO.1000106
16. Song XZ, Akasaka H, Jang HW, et al. Hematopoietic progenitor kinase 1 down-regulates the oncogenic receptor tyrosine kinase Axl in pancreatic cancer. J Biol Chem; 2020; 29. https://doi.org/10.1074/jbc.RA119.012186
17. Griffin JH, Evatt B, Zimmerman TS, et al. Deficiency of protein C in congenital thrombotic disease. J Clin Invest 1981; 68:1370-1373. https://doi.org/10.1172/JCI110385
18. Broekmans AW, Veitkamp JJ and Bertina RM. Congenital protein C deficiency and venous thromboembolism: A study in three Dutch families. N Eng J Med 1983;309: 340-344. https://doi.org/10.1056/NEJM198308133090604
19. Wintzen AR, Broekmans AW, Bertina RM, et al. Cerebral hemorrhagic infarction in young patients with hereditary protein C deficiency. British Med J 1985; 290:350 – 352. https://doi.org/10.1136/bmj.290.6465.350
20. Zhu YJ, Li JX. Plasma concentration of the natural anti-coagulants protein C and antithrombin III in leukemia (abstract). Thromb Haemost 1989;62(suppl, XII ISTH meeting, Tokyo, Japan):391. PMID: 3160800
21. Mannucci PM, Simerly PC, Tripodi A, et al. Multicenter comparison of five functional and two immunological assays for protein C. Thromb Haemost 1987; 57(1):44-48. PMID: 3590079
22. Broekmans AW. Hereditary protein C deficiency. Haemostasis1985; 15:233. https://doi.org/10.1111/j.1432-1076.2000.tb01514.x
23. Mannucci PM, et al. Deficiencies of protein C an inhibitor of blood coagulation. Lancet, 1982;2:463-7. https://doi.org/10.1016/S0140-6736(82)90494-9
24. Griffin JH, Mosher DF, Zimmerman TS, et al. Protein C, an antithrombotic protein, is reduced in hospitalized patients deficiencies of protein C, an inhibitor of blood coagulation. Blood 1982;60(1):261-264. https://doi.org/10.1182/blood.V60.1.261
25. Rodeghiero E, et al. Liver dysfunction rather than intravascular coagulation as the main cause of low protein C and antithrombin III in acute leukemia. Blood 1984; 63:965. PMID: 6584188
26. Vigano S, Mannucci PM, Solinas S, et al. Increase in protein C antigen and formation of an abnormal protein soon after starting oral anticoagulant therapy. Br J Hematol 1984;57:213-20.PMID: 6547348
27. Vigano S, Mannucci PM, D'Angelo A, et al. Protein C antigen is not an acute phase reactant and is often high in ischemic heart disease and diabetes. Thromb Haemost 1984;52:263-6. PMID: 6549416
28. Mannucci PM, Vigano S, Bottasso B, et al. Protein C antigen during pregnancy, delivery and puerperium. Thromb Hemost 1984;52:217.PMID: 6549233
29. Meade TW, Stirling Y, Wilkes H, Mannucci PM. Effects of oral contraceptives and obesity on protein C antigen. Thromb Hemostas1985; 53:198-9. PMID: 3839605
30. Takahashi H, Tatewaki W, Shihata A, et al. Plasma protein S in disseminated intravascular coagulation, liver disease, collagen disease, diabetes mellitus, and under oral anticoagulant therapy. Clinica Chimica Acta1989; 182(2):195-208. https://doi.org/10.1016/0009-8981(89)90078-8
31. Saito M, Kumabashivi, Jokaji H, et al. The levels of protein C and protein S in plasma in patients with type II diabetes mellitus. Thromb Res 1988;52:479-486. https://doi.org/10.1016/0049-3848(88)90031-X
32. Aslan B, Eren N, Cigerli S, et al. Evaluation of plasma protein C antigen, protein C activity and thrombomodulin levels in type 2 DM. Turkish J Med Sci 2005;15:305-310.
33. Ochere Addai-Mensah, Max Efui Annani-akoffor, Linda Ahenkorah Fondjo, et al. Effect of poor glycaemic control on plasma levels and activity of protein C, protein S, and antithrombin III in type 2 diabetes mellitus. Plos one 2019;14(9):e0223171. https://doi.org/10.1371%2Fjournal.pone.0223171
34. Ceriello A, Quatraro A, Dello Russo P, et al. Protein C deficiency in insulin dependent diabetes: A hyperglycemia-related phenomenon. Thromb Haemost 1990;64:104-7. PMID: 2274914
35. Koert M, Nowak-Gottil U, Kreuz W, et al. 15 parameters of coagulation and fibrino-lysis in children with type I diabetes mellitus. Klin Padiatr 1991; 203:429-32. https://doi.org/10.1055/s-2007-1025468
36. Cosio FG, et al. Plasma concentrations of the natural anti-coagulants protein C and protein S in patients with proteinuria. J Lab Clin Invest 1985; 106:218. PMID: 3160809
37. Demichell M, Contino L, Aldo Ortenisia, et al. Protein C and protein S levels in uremic patients before and after dialysis. Thromb Res 1993; 68(6):451-7. https://doi.org/10.1016/0049-3848(92)90057-h
38. Abo-Eleinein AM, Mabrouk MM, Abou-Saif S, et al. Role of both protein C and antithrombin III as predictors of stage of liver disease in chronic viral hepatitis B or C infected patients. Endocrine, Metabolic Immune Disorders- Drug Targets 2020; 20(1):112 – 117. https://doi.org/10.2174/187153031966119052902555
39. Leclerc F, Hazelzet J, Jude B, et al. Protein C and S deficiency in severe infectious purpura of children: a collaborative study of 40 cases. Intensive Care Med 1992; 18(4):202-5. https://doi.org/10.1007/BF01709832
40. Fourrier F, Chopin C, Goudemand J, et al. Septic shock, multiple organ failure, and disseminated intravascular coagulation. Compared patterns of antithrombin III, protein C, and protein S deficiencies. Chest 1992;101(3):816-23. https://doi.org/10.1378/chest.101.3.816
41. Seligsohn U, Berger A, Abend M, et al. Homozygous protein C deficiency manifested by massive venous thrombosis in the newborn. N Engl J Med 1984; 310:559-562. https://doi.org/10.1056/NEJM19840313100904
42. Estelles A, Garcia-Plaza I, Dasi A, et al. Severe inherited "homozygous" protein C deficiency in a newborn infant. Throm Haemost 1984; 52:53-56.
43. Sills RH, Marlar RA, Montgomery RR, et al. Severe homozygous protein C deficiency. J Pediat1984; 105:409-413. https://doi.org/10.1007/978-94-011-4722-4_4
44. Marciniak E, Wilson HD, Marlar RA. Neonatal purpura fulminans: a genetic disorder related to the absence of protein C in blood. Blood1985; 65:15-20. https://doi.org/10.1182/blood.65.1.15.15
45. Dreyfus M, Maguy JP, Briley F, Schwarz HP. Treatment of homozygous protein C deficiency and neonatal purpura fulminans with a purified protein C concentrate. N Engl J Med 1991; 325:1565-1568. https://doi.org/10.1056/nejm1991112823252207
46. Comp PC, Nixon RR, Cooper MR, Esmon CT. Familial protein S deficiency is associated with recurrent thrombosis. J Clin Invest1984; 74:1082-88. https://doi.org/10.1172/JCI1632
47. Schwarz HP, Fischer M, Griffin JH, et al. Plasma protein S deficiency in familial thrombotic disease. Blood 1984; 64:1297-1300. PMID: 6238642
48. Broekmans AW, Bertina RM, Reinalda-Poot J, et al. Hereditary protein S deficiency and venous thromboembolism. A study in three Dutch families. Thromb Haemost 1985; 273-277. https://doi.org/10.4066%2FJAM1.2012.1309
49. Shen MC, Lin JS, Tsay W. Protein C and protein S deficiencies are the most important risk factors associated with thrombosis in Chinese venous thromboembolic patients in Taiwan. Thromb Res 2000; 99(5):447-52. https://doi.org/10.1016/s0049-3848(00)00265-6
50. Kroiss S, Alibetti M. Use of human protein C concentrates in the treatment of patients with severe congenital protein C deficiency. Biologic Targets Therapy 2010;4:51-60. https://doi.org/10.2147%2Fttt.s3014
51. Sanz-Rodriguez C, Gil-Fernandez JJ, Zapater P, et al. Long-term management of homozygous protein C deficiency: replacement therapy with subcutaneous purified protein C concentrate. Thromb Haemost 1999; 81(6):887-90. PMID: 10404762
52. de Kort EHM, Vranken SLAG, Van Heijst AFJ, et al. Long-term subcutaneous protein C replacement in neonatal severe protein C deficiency. Pediatrics 2011; 127(5):e1338-42. https://doi.org/10.1542/peds.2009-2913
53. Crivellari M, Marino G, Belloni S, et al. Administration of human protein C concentrates in patients with double organ failure and severe sepsis after cardiac surgery. Minerva Anestesiollogica 2008; 74:48-49. https://doi.org/10.1007/s00134-009-1584-3
54. Landoni G, Monti G, Facchini A, et al. Human protein C concentrate in pediatric septic patients. Signa Vitae 2010;5(1):13-19. https://doi.org/10.22534/SV51.042010.2
55. Veldman A, Bridey F, Schwarz HP. Treatment of homozygous protein C deficiency and neonatal purpura fulminans with a purified protein C concentrate in neonatal severe purpura fulminans: a genetic disorder related to the absence of protein C in blood. N Engl J Med 1984; 310:559-562. https://doi.org/10.1186%2Fcc9226