Intraventricular Hemorrhage in a Term Neonate: Manifestation of Protein S Deficiency- A Case Report

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Abstract
Protein S (PS) is an antithrombotic plasma protein that plays essential roles in limiting thrombus formation in the anticoagulant system. Protein S deficiency is related with recurrent thrombosis. Here, the authors report a case of a term neonate with severe PS deficiency in year 2015, Imam Hospital, Tehran, Iran, that had seizures and intraventricular hemorrhage (IVH) since the age of 3 days. Nine-month follow-up did not show any developmental problems and MRI showed no hemorrhage.

Keywords: Intraventricular hemorrhage, Protein S deficiency

Introduction
Protein S (PS) is an antithrombotic plasma protein that acts mainly as a cofactor of activated protein C (APC) anticoagulant activity in the degradation of factor Va and activated factor VIII(1). PS deficiency is a hereditary autosomal dominant disorder, described in 1984 in several cases with low level of protein S and A outstanding history of recurrent thrombosis (2). Findings show in healthy individuals, approximately 60% of PS is bound in a complex with C4b-binding protein (C4bBP) that is a regulatory factor of the complement system. Only the remaining 40% of PS is capable to be a cofactor for APC (3). PS deficiency is an infrequent but harsh hereditary thrombophilic disorder with a prevalence of between 0.03% and 0.13% in the general population (2, 4). Main symptoms of PS deficiency are superficial thrombophlebitis, deep venous thrombosis and pulmonary emboli (5). In the present study the authors describe a case of a term neonate with severe PS deficiency that had seizures and intraventricular hemorrhage (IVH) since the age of 3 days.

Case Report
A 31-yr-old woman has delivered a term female infant with 40wk of gestational age. The newborn Apgar score in ten min after delivery was 7, the newborn weight was 4000 g. Because of progression of jaundice at the first day of life baby was admitted to the Imam Hospital, Tehran, Iran, in yr 2015. Parents filled an informed consent to be involved in this study. All information remained secret and no clear picture of newborn was published.
The primary physical examination showed no abnormalities. On the 3rd day of life, newborn was transferred to the Neonatal Intensive Care Unit of Imam Hospital, because of seizures. Despite phenobarbital administration, the seizures persisted. No sign of infections and metabolic disorders were found in the test results and the spleen and liver were not palpable. Brain MRI showed diffuse periventricular and intraventricular hemorrhage on both sides. The internal cerebral vein was dilated and a small number of irregular vessels in the mid sagittal region were observed. Acute ischemic foci (limited on DWT) were observed in both genu of the corpus callosum and in the frontal lobes as well. Ventricular dilatation was acknowledged too. Brain MR-angiography (MRA) revealed that main intracranial vascular structure was normal (Fig.1). A few days later, Grade 1 intraventricular hemorrhage was extended to Grade 2 and 3 hemorrhages; sonography of the brain showed Grade 2 intraventricular hemorrhage in the right and Grade 3 in the left side. Head circumference was not obviously increased from the initial measurement. Moreover, mobile masses in the inferior vena cava (IVC), protruding into the right atrium (RA) and multiple thrombosis of the portal vein were reported in neonatal echocardiography (Fig. 2). Metabolic profile (including homocysteine, ammonia, and lactate) and plasma amino acid level were normal. The blood concentration of antithrombin III, fibrinogen, protein C and factor V Leiden were within normal ranges. The study revealed the quick decrease of protein S in the infant's blood while the parent’s samples were normal. Thrombocytopenia was shown in the history of pregnancy, and the mother had been treated with dexamethasone. She was the first offspring of a consanguineous marriage. Fresh frozen plasma was infused at that time. Aspirin therapy was continued, leading to the improvement of the infant condition. During the nine-month follow-up, the infant did not show any developmental problems and MRI revealed no hemorrhage.
Discussion

Seizures happen more frequently during the neonatal period compared with any other period of the human lifetime. Population based studies recommend that the incidence of seizures in term newborns is 1 to 3 per 1000 live births (6). Generally, clinical diagnosis of neonatal seizures is difficult, since seizures are usually not accompanied by any particular clinical symptoms (7, 8).

On the other hand, the relationship between hereditary thrombotic disease and deficiencies of protein C and protein S is distinguished. We described a case of a term neonate with seizures, mobile masses in the IVC, IVH, and multiple thrombosis of portal vein due to decline in PS level. Because of the normal level of protein S in the parents, and in the infant as well, in the period of the follow-up, it appears that the protein S deficiency in this infant was temporary and as a result of A-V malformation.

To our knowledge, just a concise similar case-report has been published. Comp et al. (2, 9) described two patients that had a double heterozygous or homozygous deficiency of protein S. Fischer et al. (10) reported a newborn with homozygous qualitative PS deficiency who had a PS plasma activity <10% and intracerebral massive bleeding. Moreover, hemorrhagic ischemic stroke with protein S deficiency and thrombosis were reported in young patients (11-13).

In preterm neonates, IVH is observed mainly in the context of germinal matrix hemorrhage. On the contrary, in the term of neonates, IVH initially occurs as a result of hemorrhage in the choroid plexus or thalamus (14, 15). The long-term prognosis of PS deficiency after IVH in term newborns is ambiguous (16). Long-term follow-up is necessary to find out the predictors of neurological sequelae after IVH in term newborns.

Ethical considerations

Ethical issues (Including plagiarism, informed consent, misconduct, data fabrication and/or falsification, double publication and/or submission, redundancy, etc.) have been completely observed by the authors.

Acknowledgement

The authors declare that there is no conflict of interests.
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