Sturge-Weber Syndrome - Clinical Case Study

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Abstract: Sturge-Weber syndrome (SWS) or encephalotrigeminal angiomatosis is a syndrome, characterized by local brain atrophy and calcifications in the cerebral cortex combined with ipsilateral “Port Wine” facial nevus in the area innervated by the nervus trigeminus. SWS is a rare syndrome having incidence of 1 in 20,000-50,000 people. It affects men and women equally. SWS is a sporadically occurring congenital malformation and the fetal cortical veins (during 4-8 gestational week) develop abnormality, i.e. they remain “fetal”. There is no evidence of hereditary transmission. As associated gene mutation was found in nucleotide transition in GNAQ on chromosome 9q21. The venous drainage from the superficial brain part is carried out through sinus sagitalis superior, and the deep veins – through sinus cavernous and straight sinus. Cortical veins serve as a liaison between the two vein systems that flow in an internal jugular vein

Keywords: Sturge-Weber syndrome, epilepsy

1. Introduction

Sturge-Weber syndrome (SWS) or encephalotrigeminal angiomatosis is a syndrome, characterized by local brain atrophy and calcifications in the cerebral cortex combined with ipsilateral “Port Wine” facial nevus in the area innervated by the nervus trigeminus.

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The venous drainage from the superficial brain part is carried out through sinus sagitalis superior, and the deep veins – through sinus cavernous and straight sinus. Cortical veins serve as a liaison between the two vein systems that flow in an internal jugular vein

In SWS the cortical “connecting” veins are dysplastic and the normal venous drainage is disrupted, where at the available veins are “overfilled” and the blood flow in them is bidirectional. Blood can now be returned into the superficial veins (leptomeningeal, facial) or deep vein/sinuses. This inturn is a prerequisite for venous thrombosis formation with subsequent deposition of calcium and loss of nerve cells.

The clinical manifestation in SWS can be found at birth by the presence of facial angiomata along the nervus trigeminus (in14% of the patients cutaneous angiomamay not be present). The ocular manifestation of the syndrome is the presence of choroidal or scleral angiomata, which inturn causes increased intraocular pressure that is expressed with congenital glaucoma. Concerning the central nervous system seizures are most expressed (70-90% ), then hemiparesis (30-66% ), “stroke-like” episodes, and developmental delays.

2. Clinical Case

We present you the child H.H. at the age of 10, having severe pharmacoresistant epilepsy since he was 3 months old, that cannot be controlled by the administered antiepileptic drugs in various combinations. The partial motor seizures were expressed as clonic seizures by the right limbs accompanied by speech blockage and fear. Partial onset seizure frequency was 3-4 times a week, and secondarily generalized seizures emerged once a week. The child was diagnosed with Sturge-Weber syndrome – capillary hemangioma (nevus flammeus) affecting the left facial half and scalp, cognitive deficits (IQ = 51) and severe right-sided hemiparesis.

Figure 1: Facial nevus visualization

During the imaging diagnostic with native CT scan an expressed brain atrophy at the left and frontoparieto with massive calcification at the convexity was visualized.
Subsequently, MRI examination was performed, which confirmed the expressed brain atrophy and parenchyma at the left, and decreased signal intensity of the white brain matter.

The performed Wada test found out, that the right hemisphere is dominant for speech and the right-sided hemiparesis is not exacerbated during the functional blocking of the left hemisphere.

After a multidisciplinary discussion it was decided that the child is indicated for left hemispherotomy. A parasagittal vertical hemispherotomy by Delalande was performed and as second stage it was needed to place a lumbar-peritoneal anastomosis due to the development of communicating hydrocephalus. After the hemispherotomy the child has no epileptic seizures, without aggravation of the preoperative neurological deficit and with improved cognitive deficits as the follow-up period is 2 years.
3. Discussion

The case presented in this publication is the first hemispherotomy performed in Bulgaria.

Hemispherotomies are one of the most successful interventions in relation to the epileptic seizures control. Delalande reports about full control over seizures in 74% of the cases (60 patients in a series of 83 patients). These interventions have the greatest potential to successfully discontinue the antiepileptic drugs administration –77% of the patients have achieved full control over the epileptic seizures.

The performed imaging examinations as CT, MRI and WADA test are essential when choosing the therapy and the preoperative surgical planning.

SWS is a rare syndrome with a high degree of lobar involvement and cerebral atrophy leading to severe pharmaco resistant epilepsy. For their part, seizures cause further brain damage with progressing hemiparesis. Hemispherotomies are an important part of the neurosurgical interventions range that can significantly improve the severe pharmaco resistant epilepsies control.

4. Conclusion

The imaging examinations as CT and MRI contribute to the correct diagnosis and treatment regimen determination for the treatment of patients with SWS. Also, they take an integral part in the treatment and follow-up throughout the postoperative period.

References


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