SCHIZENCEPHALY WITH CITROBACTER SEPTICEMIA AND ABO INCOMPATIBILITY IN A PRETERM NEONATE

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ABSTRACT

Neuronal migration defects are rare causes of seizure disorder and developmental problems. Schizencephaly, the most severe form is an extremely rare entity. Presentation and outcome of Schizencephaly are variable, but it typically presents with hemiparesis, seizures and developmental deficit. Its prevalence in India is 1.5 per one lakh population. Schizencephaly is a congenital condition, characterized by cerebrospinal fluid-filled clefts that extend from the pia surface of the cerebral hemisphere to ependymal surface of the ventricle of the brain. Magnetic Resonance Imaging is the modality of choice for its diagnosis. We present a female preterm neonate of 32 weeks gestation 1.35kgs born to non consanguinous parents had reduced tone of the right upper and lower limbs. Ultrasonogram of Cranium showed prominent left lateral ventricle. MRI brain presented with left open lip Schizencephaly.

KEYWORDS: Schizencephaly, Brain malformation, Magnetic resonance imaging.

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INTRODUCTION

Schizencephaly (split brain) is a rare congenital disorder of cell migration with defect in sulcation. It is characterized by cleft in cerebral mantle, communicating between the subarachnoid spaces laterally and ventricular system medially. This disorder was originally described by Wadsworth and Yakolev in the year 1946. Two types are recognized, have prognostic significance. In type - I or closed-lip Schizencephaly, the cleft walls are in apposition and type - II or open lip Schizencephaly, in which the walls are separated. Schizencephaly type - II more common than type I. In either instance, the cleft is lined by heterotopic gray matter. In most cases, the gray matter along the cleft is polymicrogyric; in some instances, it is more dysplastic than polymicrogyric. The clefts can be unilateral or bilateral, symmetric or asymmetric and can appear anywhere in the brain, although they usually are perisylvian. In unilateral cases, perusal of the contralateral hemisphere is warranted, as subtle clefts of polymicrogyria are common. 32 weeks female singleton baby 1.35kgs was born to a III para, hypothyroid, anemic, and PIH mother aged 24 years SRM Medical college Hospital admitted to NICU for preterm care. Other 2 elder siblings also preterms with 1.9kg and 1.8kg subsequently. The antenatal profile was normal. The spine and cranium normal. Parents of this neonate were phenotypically normal.

MATERIALS & METHODS

About1ml of blood was drawn using a sterile syringe, which was inoculated aseptically into a culture bottle containing 5 to 10 ml culture media. Our hospital Microbiology laboratory with availability of BACTEC and BACT/ALERT blood culture system. After the collection of blood, it is inoculated into a blood culture bottle containing 10ml of Brain Heart Infusion broth, thus making a dilution of 1 in 10 to nullify the natural bacteriostatic/bactericidal activity of blood. After inoculation, the blood culture bottles were incubated at 37°C under aerobic conditions in the incubator/Bactec automated systems for 24 hours to 7 days and bacterial identification was performed with standard bacteriological techniques. The growth was identified by colony characteristics, Gram’s staining and standard biochemical tests described in Mackie and McCartney, Practical Medical Microbiology and Bailey and Scott’s Diagnostic Microbiology. Culture which did not yield any growth following three subcultures were reported negative at the end of 7 days. Antimicrobials sensitivity was performed by modified Kirby Bauer's Disk Diffusion Method.

RESULT & DISCUSSION

The spine and cranium normal. The parents of this neonate were phenotypically normal. Routine investigation: Hemannogram normal, COOMBS negative, CRP +ve (192mg), blood culture sent, indirect bilirubin total 15mg%, direct 1.2mg% , blood group of the baby - B+ve , mother - O + ve, blood smear showed toxic granules. Started on intravenous 10% Dextrose 80ml/kg and IV antibiotics Amikacin and cefotaxim Baby had apnoeic episodes and ABO incompatibility hence caffeine citrate and double surface phototherapy given. Blood culture report came as Citrobacter positive hence antibiotic was changed to Meropenem and Amikacin. On day six, the baby had decreased tone of right upper and lower limb Ultrasonogram of the cranium showed prominent left lateral ventricle. In view of this finding, MRI brain done which revealed left open Schizencephaly. Pediatric neurlogist opinion sought open lip Schizencephaly, communicating with left lateral ventricle- A developmental anomaly. Suggested monitoring of vital signs, watch for seizures, delayed milestones, neurodevelopmental follow up and serial Doppler studies. The baby was discharged after 17 days of stay in NICU with a weight of 1.25kgs. The baby was feeding well, bilirubin normal, neonatal reflexes were normal. Advised Breast feeding to be continued, regular follow up and immunization. Schizencephaly is an extremely rare developmental disorder of the brain and Listed as RARE DISEASE by office of RARE DISEASES OF NATIONAL INSTITUTE OF HEALTH.Type2/open lip schizencephaly is more common than type 1/fused cleft/closed cleft with 60% of unilateral Schizencephaly being of the open type. It is, however more common in the parietal and frontal lobes. Children with unilateral schizencephaly had a mild or moderate outcome more frequently than those with bilateral lesions (62% versus 28%, p<0.05). Unilateral closed-lip schizencephaly was associated with the best neurodevelopmental outcome; in contrast, 11 of 12 children with bilateral open-lip clefts had severe disabilities. Language development was significantly more likely to be normal in those children with unilateral Schizencephaly than in those with bilateral cleft (48% versus 6%, p=0.002). Our patient is a open lip

![Figure 1](https://via.placeholder.com/150)

MRI brain showing left side open lip schizencephaly
schizencephaly communicating with left lateral ventricle. Schizencephaly is often associated with other congenital abnormalities in 50-90% of cases such as agenesis of the septum pellucidum and corpus callosum, polymicrogyria (an excessive number of small, partly fused gyri), pachygyria (unusually thick convolutions of the cerebral cortex), heteropias (ectopic gray matter), septo-optic dysplasia and optic nerve hypoplasia. Our patient presented with small corpus callosum. The exact etiology is not known. The likely causes may be Genetic (thought to be associated with EMX2 Gene) . These expressed genetic factors are believed to damage the periventricular germinal matrix impairing cellular migration at 6-7 weeks of intrauterine growth. Physical insult like infection, infarction, hemorrhage, toxin and mutations. In our case, there were no stigmata of congenital infections. Schizencephaly is probably a disorder of normal neuronal migration during second trimester of intrauterine development, when primitive neuron precursors (germinal-matrix) migrate from just beneath the ventricular ependyma to the peripheral hemispheres where they form the cortical grey matter. Gray matter contains neuronal cell bodies and dendrites whereas white matter contains axons which are coated in myelin. Individuals with clefts in both hemispheres, or bilateral clefts presents with i) developmentally delayed ii)delayed speech and language iii) corticospinal dysfunction iv) Microcephaly, mental retardation v) Seizures and spastic quadriaplegia. Individuals with unilateral clefts presents with i) hemiparesis ii)average or near average intelligence iii) seizure disorder. Treatment of Schizencephaly involves i) Physiotherapy to involved extremity in hemiparesis ii) Occupational Therapy for rehabilitation iii) Treatment of seizures with oral antiepileptics and iv) Surgical management in the form of Shunt for hydrocephalous. Complications of Schizencephaly are i) Optic nerve hypoplasia ii) Skull deformity iii) Learning disability iv) Seizures.

**CONCLUSION**

Schizencephaly is a rare central nervous system developmental disorder, is very often associated with other severe brain malformations and in most of the cases subsequent multiple neurological symptoms. The method of choice in diagnosis of Schizencephaly is MRI which shows the degree and type of cleft, coexisting abnormalities and allows differential diagnosis.

**CONFLICT OF INTEREST**

Conflict of interest declared none.

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