Case Report on Subacute Sclerosing Panencephalitis

Sweta Chavhan a#, Archana Maurya b† and Amruta Kothe a

a Smt. Radhikabai Meghe Memorial College of Nursing, Sawangi (Meghe), Wardha, Datta Meghe Institute of Medical Sciences (Deemed to be University) Maharashtra, India. 
b Department of Child Health Nursing, Smt. Radhikabai Meghe Memorial College of Nursing, Sawangi (Meghe), Wardha, Datta Meghe Institute of Medical Sciences (Deemed to be University) Maharashtra, India.

Authors’ contributions

This work was carried out in collaboration between both authors. Both authors read and approved the final manuscript.

Article Information

DOI: 10.9734/JPRI/2021/v33i60B34607

Open Peer Review History:

This journal follows the Advanced Open Peer Review policy. Identity of the Reviewers, Editor(s) and additional Reviewers, peer review comments, different versions of the manuscript, comments of the editors, etc are available here: https://www.sdiarticle5.com/review-history/78380

Received 07 October 2021
Accepted 15 December 2021
Published 21 December 2021

ABSTRACT

Introduction: Subacute sclerosing panencephalitis (SSPE), frequently referred as Dawson disease, is a kind of sclerosing panencephalitis, and is an uncommon long-term, continuous cerebral inflammatory condition triggered by a gradual infection with particular faulty types of hyper mutated measles virus.

Main Symptoms and/or Important Clinical Findings: Patient came with the complaint of gait abnormalities, sudden fall while walking, loss of speech, staring look, abnormal smile and loss of bladder control, hemiplegia and contracture of all four limbs, seizures, Grade III Malnutrition.

The Main Diagnoses, Therapeutic Interventions, and Outcomes: Patient undergone investigation i.e. Blood Test: Hb - 9.8 gm%, TLC - 14,700/mm, Platelet - 2.24 lacs /cu.mm, Total Protein - 8.4 g/dl, HCT - 29.8, Serum Bilirubin - 1.0mg/dl, ALP - 99IU/L, Serum Urea - 32mg/dl, Serum Creatinine - 0.5mg/dl, Serum Sodium - 153meq/L, Serum Potassium - 2mmol/L. CSF Igg Measles test was done EEG which revealed burst of periodic complexes with well-preserved background activity. Doctor manages with Tab. Valparin 200mg BD, Tab. Frisium 5mg BD, Tab. Baclofen 10mg BD, Tab. Samion D3 OD, Syp Q-Carni 5ml BD, Tab. Augment 375mg BD.
Chavhan et al.; JPRI, 33(60B): 214-218, 2021; Article no.JPRI.78380

**Outcome:** After treatment, the child showed some improvement. The patient was admitted to Pediatric Ward No-22, AVBRH with known case of Subacute sclerosing panencephalitis (SSPE) and he had complaint of swelling over right jaw with carries tooth since 4month.

**Keywords:** Subacute sclerosing panencephalitis; Dawson disease; hyper mutated; measles virus; cerebral inflammatory condition; seizures; infection.

1. INTRODUCTION

Subacute sclerosing panencephalitis (SSPE) is indeed neurodegenerative entropy that affects youngster and adolescent central nervous systems. It’s a gradual and sustained infectious illness produced by a faulty measles virus [1]. It is also known as Dawson's Disease, Dawson's Encephalitis. Dawson described for the first in stance in 1933 kid having gradual mental decline with spontaneous motions whom, at necropsy, was discovered to also have a prominent engagement of grey matter with numerous neuronal inclusion bodies [2].

In emerging nations such as India and Eastern Europe, the prevalence of SSPE tends to be high. Males have a greater prevalence than females (male/female: 3:1) [3]. This was triggered by the measles virus's interaction in the brain, which destroys neurotransmitters. SSPE's Pathophysiology is still a mystery, although wild strains, not vaccines trains, are to blame, according to genetic investigations, which further corroborate this conclusion [4].

The process of SSPE evaluation is generally complex. Dyken criteria were first designed to aid in the diagnosis of SSPE. This series of criterion was revised in 2010 due to the fact that SSPE presentations might vary widely. Two main and one minor criterion are required for diagnosis when using this new recommended diagnostic criteria, which is part of the SSPE. Histomorphological and genetic investigation scan be used if clinical findings or accompanying requirements are lacking but the diagnosis is still likely. High anti-Measles levels of antibodies in CSF and standard or unusual patient findings are two major criteria. Radermecker complexes are typical EEG results typically feature regular, unspecified, two folded concurrent, and well-proportioned weak waves of great amplitude that reoccur every 5–15s. Amount of Cerebrospinal fluid globulin containing more than 20% CSF protein. Brain sample histopathological features, Test to identify the altered genome of the wild-type measles virus using molecular diagnostics. In most cases, two major criteria and one minor requirement are necessary [5].

The commonest typical age of onset is 5–15years, usually 5–8years after infection with measles. The most prevalent symptoms include myoclonus, cognitive deterioration, poor academic performance, and behavioral problems. The route is a steady downhill that may be divided into four distinct segments (Jabbour staging). Stage 1 is characterized by behavioral shifts and cognitive deterioration, both of which impair academic achievement. Stage 2 begins with the appearance of myoclonus. In the third stage, 6543 individuals acquire a variety of pyramidal and extrapyramidal symptoms such as stiffness, dystonia, trembling, spasms, and hemiparesis. Akinetic-mute condition with periods of profuse sweats, blood pressure fluctuations, and irregular breathing rates constitute Stage 4.[6].

There has been presently none treatment for SSPE and elimination through an efficient vaccination programme is thought to be much highly beneficial along with expense than other distinguished modes of control. The cornerstone is supportive care, which includes seizure stabilization and associated problems. One of the most commonly prescribed antiepileptic drugs is divalproate sodium. For the treatment of SSPE, there are no other are no established therapy procedures. In the therapy of SSPE, antiviral medicines and immunosuppressive agents are employed [7].

2. CASE REPRESENTATION

2.1 Patient Identification

A male child of 12-year-old from Yavatmal district admitted to pediatric ward no- 22 in AVBRH on 25th June 2021 with a known case of Subacute sclerosing panencephalitis (SSPE). He is 22kg and height 125cm.

2.2 Present Medical History

A male child of 12 years was brought to AVBRH on 25th June 2021 by her parents with the complaint of swelling over the right jaw with carries tooth since 4month. After the
investigation she was diagnosed as extra-oral abscess (cutaneous fistula).

2.3 Past Medical History

Patient was diagnosed to have Subacute Sclerosing of Panencephalitis (SSPE) since 7 years of age.

2.4 Birth History

Full term normal vaginal delivery, place of delivery was at home. He did not cry immediately after birth. Cried after 5 minutes of birth. Birth weight was 3.5 kg.

2.5 Immunization History

Patient did not receive all vaccines according to IAP (Indian Academy Pediatrics) schedules. Measles vaccine not received.

2.6 Developmental History

Achieved all milestone still 7 years of age.

2.7 Family History

Not significant.

2.8 Past Interventions and Outcomes

Patient was diagnosed with Subacute Sclerosing of Panencephalitis (SSPE) since 7 years of age. As the condition of the patient deteriorating they went to Nain hospital, Mumbai at the age of 7 years for 1 and half month. Since 5 years for which he is on medication: Tab. Sodium valproate 200 mg BD, Tab. Frisium 5 mg BD, Tab. Baclofen 10 mg BD, Tab. Samion D3 OD, Syp Q-Carni 5 ml BD.

2.9 Clinical Findings

Gait abnormalities, sudden fall while walking, loss of speech, staring look, abnormal smile, and loss of bladder control, hemiplegia and contracture of all four limbs, seizures. Generalized gingival growth over teeth. Gingiva bleeds on manipulation. Patient is in vegetative state. AEBE present and rhonchi positive.

2.11 Diagnostic assessment

2.11.1 Blood Test

Hb 9.8 gm%, TLC 14,700/mm, Platelet-2,241acs/cu.mm, Total Protein-8.4 g/dl, HCT-29.8, Serum Bilirubin-1.0mg/dl, ALP-99IU/L, Serum Urea-32mg/dl, Serum Creatinine-0.5mg/dl, Serum Sodium-153mEq/L, Serum Potassium-4.2mmol/L. CSF IgG Measles test was done.

2.11.2 EEG

The EEG revealed burst of periodic complexes with well-preserved background activity.

2.11.3 Therapeutic Intervention

Doctor prescribed Tab. Valparin 200 mg BD, Tab. Frisium 5 mg BD, Tab. Baclofen 10 mg BD, Tab. Samion D3 OD, Syp Q-Carni 5 ml BD, Tab. Augment 375 mg BD.

3. DISCUSSION

A male child of 12 years was brought to AVBRH on 25th June 2021 by her parents with the complaint of swelling over the right jaw with carries tooth since 4 month with a known case of Subacute sclerosing panencephalitis (SSPE) associated with Gait abnormalities, sudden fall while walking, loss of speech, staring look, abnormal smile, and loss of bladder control, hemiplegia and contracture of all four limbs, seizures.

This study supported to the study which clears the idea that background of fever, seizures that began at two months of age. Symptoms were associated with myoclonic twitches in the axial direction, head dips, along with diminished span of concentration for the last two months. Benchmarks were met properly, although he had not had any of his vaccines, with the exception of a dose of diphtheria, tetanus, and pertussis vaccine. The family background was devoid of mental or neurological disorders. This resulted in epileptic seizures that prolonged 30 minutes. Due to a one-hour febrile generalized status epilepticus, his treatment was supplemented with valproate when he was three years old. The
patient's physical assessments for physiological parameters as well as laboratory investigations were also normal. EEG indicated a sluggish back drop with generalized cyclic complexes compose of two folded concurrent, well-proportioned weak waves of great amplitude that did not vanish after diazepam induction. Because the EEG image suggested sub-acute sclerosing panencephalitis sample of CSF was collected to be tested for anti-measles antibodies. [8] Another study supported to the case of an 11-month-old newborn came to the hospital complaining of right-sided partial seizures for three days, followed by muscle spasm and impaired nervous system throughout the previous month. Previous to this sickness, the newborn was healthy and meeting age-apt developmental achievement. During the assessment, myoclonus of the limbs was observed. The antenatal and postnatal periods were uncomplicated, and the mother had no incidence of measles throughout gestation or at the delivery date. At eight months of age, he had background of temperature and coughing, and acute rhinitis, which was followed by a maculopapular rash (initially detected on the forehead, then descended lower), which was diagnosed as measles by a pediatrician. Other investigations were normal. Because the EEG image suggested SSPE, a sample of CSF and serum was taken to test for anti-measles antibody. An ELISA test for Ig Gantimeasles antibody utilizing commercially available kit revealed good outcomes in both CSF and serum. While anti-measles IgM antibodies were negative in both CSF and serum. The child was treated with isoprinosine (100mg/kg/day), but interferon therapy was not financially feasible. Myoclonus was controlled with sodium valproate. [9,10]

4. CONCLUSION

Subacute sclerosing panencephalitis (SSPE) is still a frequent disease in India, characterized by progressive mental impairment, myoclonus, periodic encephalographic abnormalities, and an elevated anti-measles antibody titer in CSF fluid. The measles virus infects the brain and promotes neuronal death. The patient has made substantial progress since seeking treatment, and the treatment is still continuing as of my last visit.

CONSENT

While preparing case reports for publication guardian informed consent has been taken from parent of client.

ETHICAL APPROVAL

As per international standard or university standard written ethical approval has been collected and preserved by the author(s).

COMPETING INTERESTS

Authors have declared that no competing interests exist.

REFERENCES

1. Subacute Sclerosing Panencephalitis [Internet]. NORD (National Organization for Rare Disorders). [cited 2021 Dec 14]. Available: https://rarediseases.org/rare-diseases/subacute-sclerosing-panencephalitis/
2. Jafri SK, Kumar R, Ibrahim SH. Subacute sclerosing panencephalitis – current perspectives. Pediatr Health Med Ther. 2018;9:67–71.
3. Subacute Sclerosing Panencephalitis Information Page | National Institute of Neurological Disorders and Stroke [Internet]. [cited 2021 Dec 14]. Available: https://www.ninds.nih.gov/Disorders/All-Disorders/Subacute-Sclerosing-Panencephalitis-Information-Page
4. Measles infection and encephalitis [Internet]. The Encephalitis Society. [cited 2021 Dec 14]. Available: https://www.encephalitis.info/measles-infection-and-encephalitis
5. Manning L, Laman M, Edoni H, Mueller I, Karunajeewa HA, Smith D, et al. Subacute Sclerosing Panencephalitis in Papuan New Guinean Children: The Cost of Continuing Inadequate Measles Vaccine Coverage. PLoS Negl Trop Dis. 2011;5(1):e932
6. Gutierrez J, Issacson RS, Koppel BS. Subacute sclerosing panencephalitis: an update. Dev Med Child Neurol. 2010;52(10):901–7.
7. Measles (for Parents) - Nemours KidsHealth [Internet]. [cited 2021 Dec 14]. Available: https://kidshealth.org/en/parents/measles.html
8. Vaccine-safety-E-course-manual.pdf [Internet]. [cited 2021 Dec 14]. Available: https://www.who.int/vaccine_safety/initiative/tech_support/Vaccine-safety-E-course-manual.pdf
9. Epilepsy and Disorders of Consciousness | Adams and Victor’s Principles of Neurology, 11e | AccessNeurology | McGraw Hill Medical [Internet]. Available:https://neurology.mhmedical.com/content.aspx? book id=1477 & section id=11718422

10. Manning L, Laman M, Edoni H, Mueller I, Karunajeewa HA, Smith D, et al. Subacute Sclerosing Panencephalitis in Papua New Guinean Children: The Cost of Continuing Inadequate Measles Vaccine Coverage. PLOS Neglected Tropical Diseases. 2011; 5(1):e932.

Peer-review history:
The peer review history for this paper can be accessed here: https://www.sdiarticle5.com/review-history/78380