Case Report

ADULT PRESENTATION OF DYKE DAVIDOFF MASSON SYNDROME WITH SCHIZOAFFECTIVE DISORDER — A CASE REPORT

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ABSTRACT

Dyke Davidoff Masson syndrome (DDMS) is a rare disease with characteristic radiological features, seizures, mental retardation, facial asymmetry, and psychiatric manifestations. Here, we report a case of a 50-year-old female who had a refractory seizure disorder and schizoaffective disorder. Radiological investigation revealed the features of DDMS.

Keywords: Dyke Davidoff Masson Syndrome, Cerebral hemiatrophy, Schizoaffective disorder

INTRODUCTION

In 1933, Dyke et al.¹ reported the characteristic bone changes associated with Dyke Davidoff Masson Syndrome (DDMS) in their series of nine patients who had infantile hemiplegia. They described cerebral hemiatrophy which consisted of ipsilateral thickening of the cranial wall on the same side as the cerebral lesion. These lesions were accompanied by hypertrophy of the frontal and ethmoid sinuses and the air cells of the temporal bone. Encephalogram studies of the patients showed enlargement of lateral ventricles. The adult presentation of this syndrome is quite rare and few of them present with psychiatric comorbidity. The psychiatric manifestations could be depression² or schizoaffective disorder.³,⁴

CASE REPORT

A 50-year-old unmarried lady, belonging to the low socio-economic status, presented with her mother who reported that the patient, who is on treatment for seizure disorder since childhood, has behavioral problems for a few months. The patient has been complaining that she could hear voices which were abusive and commanding her to commit suicide. At times, the patient had threatened to kill her mother and to end her own life.

The patient had normal development in her first year, till she had a serious brain infection for which she was hospitalized for two weeks. Details regarding this were not available. Subsequently, the patient developed seizure disorder for which she has been on antiepileptics. She also showed much slower development in her childhood and could not complete her primary education. She could dress on her own and maintain hygiene. She was on treatment for hypothyroidism. Currently, she was on Phenytin sodium, Phenobarbital, and Thyroid hormone. Interviewing her was difficult as she was uncooperative, and she often stated that she continuously heard the voices of her dead relatives abusing her and asking her to end her life. She also expressed that she had lost hope for living, that she was depressed, and that she wanted to die. The mother reported that she had disturbed sleep.

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Considering the presence of persistent psychotic symptoms and the history of periodic affective, obsessive, and depressive symptoms which used to last more than two weeks, a diagnosis of schizoaffective disorder was made based on the ICD-10 criteria.

Evaluation by a neurologist did not reveal any major neurological deficits. The patient was not cooperative for a detailed examination. She had difficulty in walking, but her gait was steady, and no visible ataxia or other gait disturbances were noted. On neurological examination, the cranial nerves function was normal. She had a power of 5/5 in all her limbs, and her tone was normal. She had sluggish deep tendon reflexes throughout her upper and lower limbs. Plantars were flexor on both sides.

She was initially treated with atypical antipsychotics, Risperidone (up to 12 mg) and Quetiapine 100 mg. She was given Escitalopram for episodic depressive features. As she did not improve, Haloperidol 10 mg was initiated. After two weeks, she exhibited features of tardive dyskinesia, and hence Haloperidol was discontinued. At present, she is maintained on Clozapine 200 mg. She was continued on Phenytoin sodium 300 mg and Phenobarbital 60 mg for seizure control. Even with this regime, she was not seizure-free. An attempt to switch to Carbamazepine was made but had to be terminated as she developed severe drug-induced rashes. She was not cooperative for psychological evaluation.

The radiological findings are provided in Figures 1-3.

DISCUSSION

DDMS is considered a rare disease which often presents in childhood, adolescence or young adults. After an extensive literature review, presentation beyond 50 years was found to have been reported only once. The predominant clinical presentation includes seizures, mental retardation, hemiplegia, and facial asymmetry. The probable suggested mechanism for DDMS is a cerebral insult which may have occurred in-utero or during early life which may cause the associated calvarial changes. Causes implicated include birth trauma, vascular events, hypoxia, and infections. Psychiatric manifestations have also been described, which often include psychotic features and episodic mood disorders. Cases of refractory schizoaffective disorders are frequently reported.

The case discussed here had psychiatric features suggestive of schizoaffective disorder, refractory seizure disorder, and mental retardation. The illness in the first year of her life would have contributed to development of the characteristic brain changes. The management of the patient will focus on control of seizures and her psychiatric symptoms. She may benefit from domiciliary physiotherapy.

Loss of neurons and their connections in the cerebral hemisphere is known as cerebral atrophy. Cerebral atrophy may be bilateral as in senile patients (diffuse atrophy) or neurocognitive diseases like Alzheimer’s disease (lobar predominance of atrophy), or unilateral as in diseases causing ischaemic, traumatic, inflammatory, or infective insult to the brain parenchyma. The presence of unilateral hemiatrophy is a pointer to a narrow group of differential diagnoses like DDMS, Rassmussen’s encephalitis, and Sturge-Weber syndrome. Asymmetry in size may also be due to an enlarged hemisphere as in unilateral hemimegalencephaly. In DDMS, the insult to the brain parenchyma occurs during the foetal period or in early childhood. It is characterized by hemiatrophy with ipsilateral midline shift, dilation of the ipsilateral lateral ventricle and dilated ipsilateral paranasal sinuses called pneumosinus dilatans. Ipsilateral calvarial thickening is also noted. The closest imaging differential diagnosis is Rassmussen’s encephalitis, which does not show calvarial changes. Sturge Weber syndrome can be differentiated by the presence of pial angiomias and ipsilaterally enlarged choroid plexus. In hemimegalencephaly, which is a hamartomatous overgrowth of one side of the cerebral hemisphere, the affected side shows a dysmorphic lateral ventricle, with a normal contralateral ventricle in the relatively smaller and normal cerebral hemisphere.
**Figures 1a-c:** Axial T2 weighted images of the brain shows hemiatrophy with the involvement of the left frontal (arrow-head), parietal (small arrow), and temporal regions (curved arrow) with dilation of the ipsilateral lateral ventricle (broken arrow).

**Figure 1a**

**Figure 1b**

**Figure 1c**

**Figure 2:** Sagittal T1 weighted image of the brain shows dilation of the left frontal sinus, suggestive of pneumosinus dilatans (small arrow).

Our patient is supported by her widowed mother who has great difficulty in affording the cost of treatment at the higher centres, which could be the reason for the delay in identifying her condition. Socioeconomic factors and the lack of information regarding government support for such patients have played a significant role in the delay in the diagnosis and treatment of this patient. Patient education campaigns and outreach programmes should strive to identify patients suffering from mental health diseases and provide their families necessary information on governmental assistance for treatment of such diseases. This will also help to identify any underlying organic cause for the psychiatric symptoms. The failure to do so significantly reduces the quality of life for these patients and their families.
**Figure 3:** Coronal T2 weighted image of the brain shows dilation of the left frontal sinus, suggestive of pneumosinus dilatans (small arrow)

**REFERENCES**


