Moya Moya disease presenting as Chorea: A case report

Quraishi SMS¹, Hussain KMF², G Rajni³, Afzal S⁴

¹Dr S M Saifullah Quraishi, Assistant professor, Kurnool Medical college Kurnool, ²Dr K M Iqbal Hussain, Associate Professor, Kurnool medical college Kurnool, ³Dr G Rajini, Post graduate Student, Department of Medicine, Kurnool Medical college Kurnool, ⁴Dr Sayeeda Afzal, Post graduate student in Paediatric, GVR children hospital Kurnool, India

Address for correspondence: Dr S M Saifullah Quraishi, Email-safu53@yahoo.co.in

Abstract
Moya Moya disease is traditionally known as disease of adults. Name is derived from angiographic appearance of cerebral vessels. Here we are reporting a case of Moya Moya disease in a young adolescent girl.

Key words: Moya Moya disease, chorea, involuntary movement.

Introduction
Moya moya disease is a rare disease characterized by multiple occlusions of the cerebral circulation with an unusual net like system of collaterals. In Japanese moya moya means HAZY. The disease derives its peculiar name from the angiographic appearance of cerebral vessels in the disease that resembles a ‘PUFF OF SMOKE’. Ischemic insult is more common in children. Chorea is rarely noted with this disease. When a child presents with chorea, clinicians may attribute this symptom to a number of diseases. Any child with involuntary movements, moya moya disease can be a possibility.

Case report
A 13 year old girl with history of sudden onset of right hemiparesis at the age of 3 years, recovered in 4 hours, later at the age of 8 years she developed involuntary movements of right upper limb and lower limb which subsided spontaneously, thereafter she is having on and off rhythmic, transient, dance like movements involving right upper and lower limbs with no loss of consciousness.

On examination, her consciousness was clear with good orientation. A detailed physical examination showed that the functions of cranial nerves were normal. No residual weakness of right upper and lower limbs. Hypotonia of right upper and lower limb present. Deep tendon reflexes were normal. No pathological reflexes. Sensory system was intact. Frequent episodes of chorea involving right upper and lower limb were observed.

Laboratory tests including a complete blood count and tests for anti streptolysin o titer, rheumatoid factor, anti nuclear antibodies, anticardiolipin antibodies, anti ds DNA, lipid profile, ESR, CRP, homocysteine, Hb electrophoresis, 2D Echo were unremarkable. CT scan brain at the time of hemiparesis showing small infarct left superior frontal region. Later when child presented with involuntary movements, MR angiography revealed diffuse narrowing of left internal carotid artery with critical stenosis of supraclinoid segment, paucity of left middle cerebral artery branches, non visualization of A2 segment of right anterior cerebral artery, narrowing of A1 segment of left anterior cerebral artery at origin. Digital subtraction angiography shows complete occlusion of left distal internal carotid artery and its branches beyond supraclinoid segment, prominent collaterals seen at the level of basifrontal lobes. Hypertrophied dural collaterals are seen arising from the ophthalmic artery suggestive of moya moya disease [figure1 ]. Patient is on medical management receiving aspirin and valproate. Episode of chorea has decreased with medicine.

Discussion
In children, moya moya disease presents with symptoms suggestive of ischemia of brain. These include headache, seizures, focal neurological deficit, recurrent TIA [1]. Headache is a common presentation due to dilatation of leptomaningial and dural collateral vessel stimulating dural nociceptor [2]. Involuntary movements [chorea] have been rarely described previously [3].
Chorea in moyamoya disease may be an early warning sign of vascular insufficiency in the brain. In children with crying and breath holding episodes can precipitate abnormal movements due to decrease blood supply in critical area [4]. There are reports of association of moyamoya disease with variety of disorder like radiotherapy of head and neck, sickle cell anemia, neurofibromatosis type 1, autoimmune disorder [Grave disease, Antiphospholipid syndrome] and congenital heart disease, but most common is idiopathic [5]. In this patient no association with other disease was found.

Vascular bypass are performed to increase perfusion and to decrease hypoxia in brain. Clinical and angiographic improvement can be seen after bypass procedure in all age group patients unless patient had permanent neurological damage in the form of infarct [6]. As the course of disease is unpredictable sometimes it is slowly progressive with intermittent neurological event or occasionally disease can have fulminant course. However clinical symptoms invariably progress irrespective of angiography finding in untreated patient [7]. Medical therapy cannot halt progression of disease [8]. Surgical treatment delays symptomatic neurological progression, that is why timing of surgery is very important. Even in presence of severe angiographic changes if revascularization surgery is done before disabling infarct, prognosis is excellent [9]. Delay in diagnosis results in delay in treatment and increasing permanent disability from stroke. Medical therapies mainly confined to treat acute episode like seizure and infarct.

Conclusion

In children and young adult presenting with stroke we should consider moyamoya disease. Early revascularization surgery will prevent permanent neurological disability.

Most important predictor of overall outcome in moyamoya is neurological status at the time of treatment.

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How to cite this article?

Quraishi SMS, Hussain KMI, G Rajni, Afzal S. Moya Moya disease presenting as Chorea: A case report. Int J Med Res Rev 2014;2(3):256- 258. doi:10.17511/ijmrr.2014.i03.15.