Cerebellar Mutism After Pineal Tumour Excision: A Case Report

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Abstract

Cerebellar mutism syndrome (CMS) was first described by Rekate et al in 1985. This syndrome is a common complication of posterior fossa surgery in children with range of 11-29% and usually manifests as diminished speech, hypotonia, and ataxia. The cause is due to bilateral perturbation of the dentate nuclei and their efferent pathways by edema, perfusional defects, axonal damage or metabolic disturbances. Other rare causes of CMS like acute subdural hematoma of the posterior fossa, head injury, brainstem glioma surgery, meningitis and basilar artery occlusion have also been reported. CMS after supracerebellar resection of the pineal tumor is a very rare with very few cases reported. We report such a case in a 10-year old boy who underwent excision of a pineal tumor through the infratentorial supracerebellar route.

Key Words: Cerebellar mutism, hydrocephalus, pineal tumor, posterior fossa.

Introduction

Cerebellar mutism syndrome (CMS) was first described by Rekate et al in 1985. This syndrome is a common complication of posterior fossa surgery in children with range of 11-29% and usually manifests as diminished speech, hypotonia, and ataxia. The cause is due to bilateral perturbation of the dentate nuclei and their efferent pathways by edema, perfusional defects, axonal damage or metabolic disturbances. The sequelae may be permanent if the right cerebellar hemisphere is involved. Other rare causes of CMS like acute subdural hematoma of the posterior fossa, head injury, brainstem glioma surgery, meningitis and basilar artery occlusion have also been reported. CMS after supracerebellar resection of the pineal tumor is a very rare with very few cases reported.

Case report

A 10-year old male child presented to the emergency with history of severe headache and vomiting for the last 2 months. There was associated difficulty in walking and decreased appetite. On examination the child appeared sick and there was early papilloedema bilaterally with no cranial or motor deficits. The routine hematology and biochemical parameters were normal. Computed tomogram (CT) revealed gross hydrocephalus with an enhancing tumor in the pineal region which was further defined by magnetic resonance imaging (MRI). The tumor was heterogeneously enhancing and displacing the great veins superiorly and compressing the brainstem inferiorly (Figure 1). Infratentorial supracerebellar approach in sitting position along with a right posterior external ventricular drain was used for complete tumor removal with intraoperative problems. Postoperative the child recovered well and was able to ambulate and feed himself after 48 hours. On the fifth postoperative day the child suddenly stopped talking, responding to commands and there was generalized spasticity. There were episodic abnormal cry, shouting, difficulties in swallowing and generalized tonic contractions. Repeat CT showed complete tumor excision, resolution of the hydrocephalus and no postoperative hematoma (Figure 2). No biochemical cause could be found to explain these new onset symptoms. The histopathology revealed as a benign pineal gland tumor. A diagnosis of CMS
was made, the family counseled and he was managed with a nasogastric feeding, physiotherapy and muscle relaxants. The symptoms slowly disappeared and at the end of 4 months the child started to respond with return of speech. Presently he is independent in activities of daily life and goes to school but still has learning difficulties.

Various studies have found abnormalities in the postoperative imaging in the left temporal lobe, the left and right basal nuclei, and the right frontal lobe\textsuperscript{6,9}. Speech impairment in CMS can be investigated based on standardized acoustic speech parameters and perceptual criteria and has been divided as dysarthria in post mutistic phase and children with mainly behavioral disturbances\textsuperscript{6-9}. This syndrome has been associated with medulloblastoma and other posterior fossa tumors. CMS after pineal surgery seems to be very rare but as shown by this case is still a probability. The cause may need further investigation but one probable reason may be continued pressure on the cerebellum to cause damage to the superior cerebellar peduncle. Why the CMS developed in this child on the fifth day when he had recovered completely is unknown.

All the neurosurgeons must be aware of CMS and its self limiting course. The family members need to be counseled regarding CMS prior to the surgery. Meticulous technique during surgery and prevention of continued retraction or manipulation of the superior cerebellar peduncle is must to avoid this complication postoperatively.

**References**


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