Chiari type I malformation with cervicothoracic syringomyelia subterfuge as Flail arm syndrome

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Abstract

Chiari type I malformation with cervicothoracic syringomyelia although very common in clinical practice usually in children can progress slowly and mimic muscular dystrophies in adulthood. We present a rare adult case of Chiari type I malformation with cervicothoracic syringomyelia subterfuge as Flail arm syndrome. A 44-year-old man was diagnosed with congenital type I Chiari malformation with cervicothoracic syringomyelia about 21 years ago without surgery. His health status deteriorated over the years until 21 days prior to presentation when he had severe pain in the right knee. In his upper limbs, he had bilateral corresponding severe weakness of 0/5 proximal strength and 0/5 strength in his distal muscles. Magnetic resonance imaging (MRI) revealed an enlargement of the spinal cord from C1-C4 level with a mass that appeared hypo-dense on T1 and hyper-dense on T2. Syringomyelia is a potentially serious neurologic condition that can mimic other neuromuscular disorders. Early detection and diagnosis with MRI is crucial to avoid irreversible neurological complications. We suggest that whether asymptomatic or symptomatic, decompressive surgery should be carried out to allow for free flow of cerebrospinal fluid thereby improving the quality of life for the patient.

Case Report

A 44-year-old man presented with worsening chronic motor symptoms for 21 years. He is a known case of congenital type I Chiari malformation with cervicothoracic syringomyelia diagnosed about 21 years ago without surgery. He was not able to access surgery from the time of the initial diagnosis (21 years ago) until he presents at our facility because of financial constraints. He health status deteriorated over the years until 21 days prior to presentation when he had severe pain in the right knee. The right knee pain is aggravated by walking, climbing stairs or in sitting and relieved by lying in bed. There was no associated swelling, no neck pains, backaches or abdominal pains. He denied having dysphagia, cough, difficulty in breathing or palpitations. All routine laboratory investigations were essentially normal. Electro-physiological studies reveal bilateral absence of median, ulnar and radial nerves compound muscle action potentials (CMAP) with slightly abnormal sensory nerve action potential (SNAP) at the upper limbs. However, the all the nerves on the lower limbs have normal CMAPs and normal SNAPs. Electromyography (EMG) of his upper extremities revealed of diffuse chronic neurogenic changes in almost all the nerves but normal in all the nerves in the lower extremities.

Preoperative MRI done revealed an enlargement of the spinal cord from C1-C4 level. Within the spinal cord is a mass that appears hypo-intense on T1 and hyper-intense on T2. Syringomyelia is a potentially serious neurologic condition that can mimic other neuromuscular disorders. Early detection and diagnosis with MRI is crucial to avoid irreversible neurological complications. We suggest that whether asymptomatic or symptomatic, decompressive surgery should be carried out to allow for free flow of cerebrospinal fluid thereby improving the quality of life for the patient.

Introduction

Syringomyelia occurs when a cyst, typically made up of excess cerebrospinal fluid (CSF), cumulates inside the spinal cord either within the parenchyma as a focal dilation of the central canal,1-5 Chiari malformation type I is a form of non-communicating syringomyelia,3-6 Flail arm syndrome (FAS), also referred to as brachial amyotrophic diplegia or man-in-barrel syndrome, is a gradually progressive sporadic motor neuron disorder, depicted by severe flaccid paralysis and muscle wasting in both arms symmetrically, but comparatively sparing the legs and bulbar parts, and scarcely signs of upper motor neuron lesions.7-9 The onset of syringomyelia although commonly chronic and gradually progressive, very few case advanced into adulthood unnoticed. The severe progressive adult type is most likely to mimic muscular dystrophy’s.9 Cerebellar tonsilar ectopia bulging through the foramen magnum is frequently the imaging findings in patients with this condition.3,4,10 The gold standard for treatment for this disorder is surgery and the choice of surgical approach for Chiari type I, depending on the surgeon’s understanding of the pathophysiology of the disease manifestation.1,4 In most cases the pathology is situated at the level of the foramen magnum leading to disparities in the contents and the volume of the posterior fossa. We present an adult case of congenital type I Chiari malformation with cervicothoracic syringomyelia subterfuge as Flail arm syndrome and review of literature.

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Ethics approval and consent to participate: The ethical committee of the hospital fully approved our case study. The patient was informed about our intention to involve him in a case study and he agreed to partake in the study. He signed the concern form before the operation was carried out according to all surgical protocols.

Conflict of interest: the authors declare no potential conflict of interest.

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intense on T2. These findings are consistent with Chiari malformation type I with prominent syringomyelia (Figure 2A and B). MRI of lumbar vertebral and adjacent structures showed no abnormalities. There is no disc space narrowing, no abnormal vertebral body and no cord stenosis. The patient was scheduled for suboccipital decompression of the syringomyelia.

The patient was on park-bench position after general anesthesia with the surgical incision area marked. A straight median incision was made from the occipital protuberance up to C3. Subcutaneous tissues and muscle was dissected up the C3. Suboccipital craniectomy and laminectomy of C1 and C2 was done using the high-drill and further exposor with a Kerrison rongeur. Resection of the atlanto-occipital dura was done to free the compression. The dura was peeled off with a Y shaped incision and the arachnoid membrane was intact. After securing total hemostasis, the fascia, muscle, subcutaneous tissues and skin was closed in layers. Estimated blood lost was 50 mL. We noticed during surgery that there were no obvious abnormalities on the skin and subcutaneous tissues but however, there was thickening of paraspinal muscle with retracted occipital plane and thickening of atlanto-occipital fascia. The patient was nursed in the recovery word and later to neurosurgery word. Post-operative management was uneventful. His amyotrophy did not improve much while on admission. He was discharged home two weeks after the operation. Outpatient visits were arranged at 3 months’ interval. MRI done at the 2-year follow up showing the syringomyelia was shrinked significantly (Figure 2C and D). His amyotrophy remained stable over the years afterwards.

**Discussion**

The precise pathogenic link between syringomyelia and Chiari type 1 malformation is still a matter of debate among neurosurgeons and quite a number of authors have proposed various theories as the formation of syrinx.\(^1,3,4\) Most of the authors have come to the conclusion that malformation of the mesodermal occipital somite is accountable for the tiny nature of the posterior fossa leading to the anomalous CSF flow at the level of the foramen magnum.\(^1,11,12\) Therefore, most disorders originate from the cervical spinal cord and can enlarge superiorly into the brainstem leading to the formation of syringobulbia or inferiorly into the thoracic and sometimes the lumbar regions.\(^3,4\) Apart from Chiari type 1 malformations, Syringomyelia can...

*Figure 1. Physical examination showing patient’s upper extremities with severe atrophy of forearm extensors (A) and flexors (B).*

*Figure 2. Preoperative magnetic resonance imaging scans demonstrating the syrinx from the level of C1 extending inferiorly into the thoracic spinal cord (A and B), and at 2 years’ follow-up showing the shrinking syrinx (C and D).*
occur together with diverse number of congenital anatomic anomalies as well as acquired structural abnormalities such as scoliosis, spina bifida, tumors and hemorrhage, post-infectious, post-inflammatory and post-traumatic conditions. 4,5

FAS characteristically presents with progressive upper limb weakness and wasting that is often symmetric and proximal, without significant functional involvement of lower limbs or bulbar muscles.4,5,9 Hu et al used FAS to describe this disorder8,13 although many other authors have also described the same disorder and given it various names such as Vulpian-Bernhardt syndrome,5,9,14 hanging-arm syndrome,9,15 neurogenic man-in-a-barrel syndrome,8,9 and brachial amyotrophic diplegia.8,9 The presentation of our case typically mimics the above condition with diverse names. Amyotrophic lateral sclerosis (ALS) is also a degenerative motor neuron disorder, with progressive loss of both upper and lower motor neurons in motor cortex, spinal anterior horn cells and motor neurons in brain stem.7,16 Our patient disease presentation does not correspond to ALS because his lower limbs was spared. Syringomyelia is general classified based on the pathological and magnetic resonance physiognomies of numerous spinal cord cysts. They are put into five categories: non-communicating, communicating, primary parenchymal cavitations, atrophic cavitations and neoplastic cavitations.4,6 The symptomatology of syrinx can be relatively complicated and diverse among patients because of the different syrinx types, the spinal levels involved, the degree of extension as well as the diameter of lesion.4,17 The potential permutations of the clinical symptoms comprise of headache, severe segmental central and dysesthetic pain, loss of temperature and pain sensation, down-beating nystagmus, vocal cord dysmotility, urinary frequency and incontinence, stiffness, weakness, and sclerosis.4,17-19 Mora et al noticed central cord syndrome as one of the cardinal presentation of this condition.4

MRI is the gold standard for the diagnosis of a Chiari type 1 malformation with or without Syringomyelia.20 Patients with Syringomyelia and Chiari type 1 malformation should have MRI of their brain and spine during their evaluation. CAT scan (CT) although valuable does not provide accurate diagnosis of this disorder. It is however the best in visualizing the bone anatomy of the foramen magnum.20 MRI of the brain and spine will tell whether or not a patient has Chiari type 1 malformation and/or syringomyelia, or any other abnormality of the brain that may cause similar

descendence of the cerebellar tonsils approximately 3-5mm or more below the opening in the bottom of the skull.20-22 T1W image produces a more accurate reflection of cyst size than the T2W image, because it is less sensitive to flow effects. Septa in the cyst can best be seen in the T1W image.10 Zamel et al. proposed the use of brain stem auditory evoked potentials during posterior fossa decompression to assess functional integrity of the central auditory system during surgery.23

Surgery is the main treatment for patients with syringomyelia. Nevertheless, not all patients need surgical intervention. Patients who are incidentally diagnosed with MRI studies but do have any clinical manifestation of the disorder are mostly followed with serial imaging in some institutions. Some author thinks these kinds of patients should be operated.1,4 Some authors have indicated that most patients with mild symptoms remain stable and in some instances their disorders unexpectedly disappear on radiology while other are still of the view that such patient should have surgical correction of the anomaly right way to avoid further complications.4,24 We are of the view that syringomyelia whether asymptomatic or mild should be surgical corrected putting the history of advancement of our case into perspective. Surgery is the usually the required treatment option for patients with obvious symptoms. Decompression to reinstate regular CSF flow is the goal of surgery. Posterior foramen magnum decompression with or without dura opening or anterior foramen magnum decompression and shunting are some of the surgical techniques use by varies authors.1,4,17

Navarro et al are of the view that the best surgical approach to this disorder is to recreate the mesodermal section at the level of the foramen magnum. They indicated that bone and dura should be reconstructed without disrupting the structures emanating from the neuroectoderm, that is the leptomeninges and cerebellum.3 This approach would reduce most surgical complications that are associated with CSF release and adhesive arachnoiditis by enlarging the posterior fossa and redirecting the flow of CSF hence minimizing morbidity.3 Most authors including Navarro et al agree with the use of suboccipital craniectomy and laminectomy to the level of the tonsillar herniation. They are of the view that after resecting the dura thickened band, careful dissection of the dura is commenced by protecting the inner layer of the dura and avoiding arachnoid laceration. This maneuver releases the pressure on the dura and grant passage of the posterior fossa contents because the dura and bone together provides opposition to the cerebellum.1,25-27 Most patients experience mild improvement to restoration of function after surgical decompression the Chiari malformation depending on the magnitude of their symptoms before surgery.4 In adults, the average recurrence rate after decompressive surgery is about 6.7% in most patience.4,17

Conclusions

Syringomyelia is a potentially serious neurologic condition that can mimic other neuromuscular disorders. Early detection and diagnosis with MRI is crucial to avoid irreversible neurological complication. We suggest that whether asymptomatic or symptomatic, decompressive surgery should be carried out to allow for free flow of CSF thereby improving the quality of life for the patient.

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[Neurology International 2017; 9:7336] [page 73]
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[Neurology International 2017; 9:7336]