A New Entity: Chiari Zero Malformation and Its Surgical Method

INTRODUCTION

When Hans Chiari first wrote about the Chiari malformations, he identified four different types based upon what brain structures were displaced out of the skull (3). Today, defining exactly what Chiari means is not an easy task (11,12). Most people define Chiari as tonsillar herniation of greater than 3mm-5mm. Herniation is measured as the distance below the foramen magnum (11,12).

While this definition is still widely used, authors showed that the amount of tonsillar herniation is not related to severity of symptoms or clinical outcome in the Chiari malformation. Conversely, there are some patients with with syringomyelia who exhibit classic Chiari-type symptoms with little to no herniation (3,11,12).

Recently, Iskandar and Tubbs et al. used “Chiari zero malformation” to characterize this kind of syringomyelia disorder and advised craniovertebral decompression (CVD) as the surgical modality (8,9,16,17). There is a very limited number of papers on the “Chiari zero malformation”, and there is a query about the truth of this pathology (8,9,10,14,16,17,18).

We presented a patient with a diagnosis of Chiari zero malformation who was treated with combined syringopleural shunting and CVD.

CASE REPORT

A 38-year-old-man was admitted to our hospital with headache, throbbing nuchal pain and weakness of the left leg. The neurological examination presented hypoesthesia below T7-(reduced sensation of vibration and position ipsilaterally,
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and of pain and temperature contralaterally below the T7 level together with inability to feel hot or cold, monoparesis at the left leg without spasticity and marked hyperactive patellar and Achilles reflexes. MRI revealed cervical syringomyelia extending from C1 to Th5 and associated small posterior fossa. There was no tonsillar herniation (Figure 1A,B).

He underwent syringopleural shunting via C5 and C6 laminectomy. Postoperatively, the syrinx collapsed and his clinical status was improved (Figure 2A). Nine months after surgery, his clinical status deteriorated. He was also suffering from headaches at times that were especially aggravated while coughing and straining while defecating. In his neurological examination, there was hypoesthesia below T7 level together with slight paresis of his left leg. MRI revealed that the cervical syringomyelia regrowth extended from C1 to C5 although the shunt was functional (Figure 2B). There was no tonsillar herniation, but the cerebellum and cerebellar tonsils filled the posterior fossa to capacity and the brain stem and cistern of the posterior fossa was compressed. Posterior fossa linear and volume measurements were performed according to the Tubbs and Furtado studies (5,16). The distance from the basion to the opisthion was 38 mm, and the tip of the obex was found to be at 4.0 mm superior to the foramen magnum. The midsagittal horizontal distance of the spinomedullary junction at a midpoint on a line connecting the basion to the opisthion was 13 mm. The distance from the fastigium to the floor of the fourth ventricle was 9.0 mm. The horizontal distance from the sphenoccipital synchondrosis to the basis pontis was 3.0 mm. The width of the posterior cranial fossa was 9.5 cm, the anteroposterior dimension was 6 cm and the height was 5 cm. The PFV was 205.3 cm³ and ICV was 1002.66 cm³. The PFV to ICV ratio was 0.204. (rev6)

Cine MRI showed that there was a CSF flow block at the craniovertebral junction. Brainstem evoked potentials (BAEP) also showed that there was low brainstem dysfunction manifesting prolongation of I-III interpeak latency (IPL).

We accepted the patient to have a Chiari Zero malformation. CVD was performed with the patient in supine position, and dural grafting was performed with bovine pericardium. Postoperatively, the clinical status was improved. Immediately after surgery, the headache and nuchal pain remitted. The postoperative MRI depicted the large created cisterna magna, enlargement of the fourth ventricle as well as of the superior and prepontine cistern. During the follow-up, MRI six months after surgery revealed that the syrinx had collapsed again (Figure 3A,B).

One year later, the patient continued without headache, nuchal pain and left monoparesia but the hyperactive patellar and Achilles reflexes remained unchanged. Cine MRI showed that CSF flow was normal in the craniovertebral junction, and BAEP also revealed improvement in I-III IPL.

**DISCUSSION**

In 1997, Iskandar et al. presented five children with syringohydromyelia, where clinical and radiological resolution was demonstrated after posterior fossa decompression, at the 47th Meeting of the AANS/CNS Joint Section of Paediatric Neurosurgery (6). None of the patients had hindbrain herniation. Their course was similar to that of patients

Figure 1: A) T1-weighted sagittal MR images of the craniocervical junction, showing the midline structures of the posterior cranial fossa and the brainstem and the cerebellum, cervical syringomyelia. The cisterna magna is completely obliterated by the thickened posterior rim of the foramen magnum. Note the lack of hindbrain herniation. There is a CSF block at the inferior pole of the cerebellar tonsils. B) Preoperative sagittal T1-weighted craniocervicospinal MRI shows impacted cisterna magna, diminished fourth ventricle and cervicothoracal syringomyelia.
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In cases in which preoperative cine–MR imaging was performed, decreased or normal-to-decreased flow was demonstrated (6,7,8). Four years later, Tubbs, et al. added a sixth case and found that even in the absence of tonsillar ectopia, the obices were located more than two standard deviations below normal (16). “Chiari zero malformation” was the name they chose to characterize the disorder. In 2004, Tubbs added 7th case to this series [17].

Today, Chiari 0 is a kind of syringohydromyelia, but with minimal findings of hindbrain herniation, or it is a term used when a patient is symptomatic but the herniation extends less than 5mm and is associated with syringomyelia. Synonyms are “Chiari-like pathophysiology”, and “borderline Chiari” (6,7,12,14,15,18). Some authors describe this disorder as “tight cisterna magna” (10).

Figure 2: A) Early Postoperative T1-weighted sagittal MRI showing collapsed syringomyelia after syringopleural shunting. There is no CSF block at the inferior pole of the cerebellar tonsils. B) Postoperative T2-weighted sagittal MRI showing regrowth of syringomyelia 9 months after SP shunting. There is a CSF block at the inferior pole of the cerebellar tonsils.

Figure 3: A) Postoperative sagittal T1-weighted cranial MRI showing the newly created cisterna magna, opening of the fourth ventricle and enlargement of the cisternae magna, and superior and preponine cisternae. and collapse of the syringomyelia 6 months after craniocervical decompression was added to syringopleural shunting. B) Postoperative sagittal T2-weighted craniovertebral MRI showing the artificial mega cisterna magna, and collapsed syringomyelia, and functional shunt 6 months after craniocervical decompression.
There are some questions and dilemmas regarding the subject: 1. Is it a physiological or anatomic abnormality?; 2. What is the method of detection or diagnosis?; 3. What is the prevalence?; and 4. What is the suitable treatment? (8,18).

In 1999, Milhorat, et al. studied 364 symptomatic patients with Chiari I malformation (12). They found tonsillar ectopia of less than 5 mm in 32 (9%) of the 364 symptomatic patients, but compression of the CSF spaces posterior and lateral to the cerebellum was present in all cases. In fact the tonsils were at the level of the foramen magnum in the illustrative case they presented.

In 2000 Meadows, et al. published a review of 22,591 patients in whom MR imaging was performed (11). Tonsillar herniation extending more than 5 mm below the foramen magnum was found in 0.77%. Fourteen percent of those patients were asymptomatic (with 7–25 mm of ectopia) and 25% of those had peg-like tonsils. The authors concluded that “isolated tonsillar herniation is of limited prognostic utility, and should be considered in the context of all available clinical and imaging data.”

This limited usefulness of tonsillar herniation in defining Chiari has led many experts to base their diagnoses not solely on MRI’s, but rather on a combination of MRI, cine MRI, symptoms, neurological examination, and their own experience (1,2,4,6,7,11,13,15,18).

Based on the review of all these papers the following remarks can be made. 1) Radiologically significant tonsillar ectopia may be completely asymptomatic. 2) There are patients in whom the tonsillar herniation extends less than 5 mm from the foramen magnum, whose clinical behaviour is similar to the remainder of the Chiari population, including even those with radiological evidence of syringohydromyelia.

Dr. Sekula and Dr. Jannetta, have extended this line of research and shown that so-called Chiari like patients (or Chiari 0 Malformation) tend to have small posterior fossa, similar to Chiari I patients (13). In this group, MRI reveals syringomyelia without tonsillar herniation, and a small posterior fossa. The obex is significantly lower than normal. Other reported abnormal anatomic findings in Chiari 0 malformation are increased AP diameter of the foramen magnum, increased AP distance of spinomedullary junction to the foramen magnum, and increased angle between the floor of the 4th ventricle and clivus. The measurements of our patient were similar to this study. This and other studies have led to the theory that Chiari is not a disorder of the brain, but rather a result of the hindbrain herniation extending less than 5 mm from the foramen magnum, whose clinical behaviour is similar to Chiari I malformation. In this group, MRI reveals syringomyelia without tonsillar herniation, and a small posterior fossa. It suggested that craniovertebral decompression is beneficial.

Conclusions

Measurements of posterior fossa and CSF flow at the craniovertebral junction, and BAEP can be used to diagnose “Chiari zero malformation”. Craniovertebral decompression seems to be the best initial approach for this entity as it addresses the underlying pathophysiology of an alteration in normal CSF flow patterns. We think that this is a valuable description of the Chiari zero malformation but more prospective reported cases with long term outcome are needed.

References


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