Acquired Chiari malformation following spinal cord injury—a case series

David McKean1 · Umme Sara Zishan2 · Sarah Billingsley1 · Shyam S. Swarna1 · Cormac O’Neill1 · Monika Banerjee3 · Safa Siddiqi1 · Joseph Papanikitas1 · Sarah Yanny1 · Richard Hughes1 · Tom Meagher1

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Abstract

Introduction Chiari malformation is characterized by caudal descent of the cerebellar tonsils through the foramen magnum. Acquired Chiari malformations (ACM) have previously been described after a variety of pathologies including lumbar puncture, cerebrospinal fluid (CSF) drainage, lumboperitoneal shunts, and conditions causing craniocephalic disproportion. Case presentation We present four cases of ACM following spinal cord injury (SCI), which has not previously been described in the literature. Discussion ACM is rare and typically associated with abnormalities in CSF pressure or space-occupying lesions. This case series describes the potential association of SCI with ACM. We discuss the imaging findings and clinical management of these patients. Early recognition and intervention may be important to prevent progressive neurology in this vulnerable patient group.

Introduction

The development of cerebellar tonsillar descent, also referred to as an acquired Chiari malformation (ACM), has been described as a recognized but rare complication following lumboperitoneal shunting, spontaneous cerebrospinal fluid (CSF) leakage, and other conditions, which result in craniocephalic disproportion including pseudotumour cerebrii and craniostenosis [1–3]. In addition, space-occupying lesions such as meningiomas, medulloblastomas, epidermoid cysts, and arachnoid cysts have been reported in the literature as the rare causes of ACM and syringomyelia. The pathogenesis of ACM is not well understood but the predominant theory suggests that it is related to abnormal craniospinal pressure gradients [4]. Spinal cord injury (SCI) following spinal trauma is known to cause disturbances in spinal anatomy and abnormal CSF flow dynamics [5]. We report four cases of acquired tonsillar descent, which developed in patients with a history of SCI.

Case presentation

Case 1

A 3-year 4-month-old male child suffered an incomplete tetraplegia following a road traffic accident as a front seat passenger. Injuries at the time of presentation included diffuse axonal injury and complete C1/C2 dislocation (Fig. 1a.). Additional injuries included right radial and ulnar fractures and a right median nerve neuropraxia. The patient was treated for several weeks with halo stabilization.

Routine follow-up MRI of the spine at 8 months post SCI demonstrated that the patient had an ACM with tonsillar herniation of ~13 mm into the vertebral canal (Fig. 1b). There was no relevant clinical history of lumboperitoneal shunting or evidence of an intracranial space-occupying lesion. He had no significant change in his neurological status and was managed conservatively. He had annual MR imaging and clinical assessment showing no radiological or clinical deterioration. He continued to mobilize with an unsteady gait using a Kaye walker and no significant

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*David McKean
david.mckean@nhs.net

1 Stoke Mandeville Hospital, Buckinghamshire Healthcare NHS Trust, Aylesbury, UK

2 Queen Elizabeth University Hospital, NHS Greater Glasgow and Clyde, Glasgow, UK

3 Oxford University Hospitals NHS Trust, Oxford, UK

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change in his neurological function has been reported in the subsequent 18 months.

Case 2

A 1-year 10-month-old male child suffered an incomplete tetraplegia following flexion/extension SCI RTA as a rear seat passenger. Imaging at the time of presentation demonstrated cord injury at the cervicothoracic junction (Fig. 2a). He was managed conservatively with a halo device for 8 weeks. Two years following the initial injury, the patient presented with recurrent headaches. MRI at this time demonstrated tonsillar herniation of ~10 mm consistent with ACM. There was no relevant clinical history of lumboperitoneal shunting or evidence of an intracranial space-occupying lesion. The patient’s headaches persisted and after review he had a VP shunt inserted with subsequent partial relief of symptoms. The ACM has been followed up annually with MRI and clinical assessment. The patient continues to mobilize in a wheelchair and is able to stand with support. No progressive neurological deterioration has been reported.

Case 3

A 23-year-old male presented with complete paraplegia after an RTA secondary to a spinal hematoma. He had a complete lesion at neurological level T2 and had been managed conservatively. Twelve years following his injury he presented with decreased function in his right hand. MRI demonstrated tonsillar descent of 12 mm (Fig. 3). MRI revealed extensive local dural adhesions, focal cord edema and a small central cord syrinx at the level of C2. There was no relevant clinical history of lumboperitoneal shunting or evidence of an intracranial space-occupying lesion. Following discussion with neurosurgery and clinical review, the decision was made to manage the patient conservatively. Annual MRI did not demonstrate any significant interval change in the extent of ACM and there has been no further progression in the patient’s neurology.

Case 4

A 32-year-old male patient suffered an RTA in 2002 developing a T2 level complete paraplegia. Fifteen years following his spinal injury he presented with a new difficulty in swallowing, double vision, and loss of power in the upper limbs. MRI at this time demonstrated an ACM with tonsillar descent of ~12 mm (Fig. 4a). There was no relevant clinical history of lumboperitoneal shunting or evidence of an intracranial space-occupying lesion. The ACM was associated with abnormal fluid intensity signal within the cervical spinal cord extending from level of C2 to C4 consistent with new syringomyelia. At T3 there was an impression of extensive local dural adhesions, a loculated
Fig. 2  a Sagittal T2-weighted image at the time of injury. There is partial transection of the cord at the cervicothoracic junction and extensive cord oedema, with disruption of the posterior longitudinal ligament and ligamentum flavum. No Chiari malformation is present. 

b,c Sagittal T2- and sagittal T1-weighted images 3 years post injury. Myelomalacia and post-traumatic cyst formation with focal tethering of the cord secondary to dural adhesions at the site of injury. There is tonsillar descent through the foramen magnum of ~10 mm, consistent with acquired Chiari malformation.

Fig. 3 Sequential T2-weighted sagittal images of the cervical spine 12 years post injury demonstrate focal myelomalacia and atrophy at the site of decompression at the C7/T1 level. The cerebellar tonsils extend 10 mm below the level of the foramen magnum consistent with acquired Chiari malformation. Abnormal fluid signal intensity within the spinal cord is seen at the level of C2 consistent with associated syringomyelia.
syrinx and myelomalacia of the atrophic cord at the site of prior injury.

The patient’s neurological symptoms continued to deteriorate and after a multidisciplinary discussion with the neurosurgical team the patient was put forward for operative management. He underwent a T1-T3 laminectomy and cordectomy. His new neurological symptoms resolved and he returned to his baseline of T2 complete paraplegia. No further deterioration of the patient’s neurological condition was reported and follow-up MRI demonstrated resolution of the ACM.

Discussion

A Chiari malformation is a condition whereby cerebellar tissue extends into the spinal canal through the foramen magnum and can be either congenital or acquired. The most
common type of Chiari malformation is a type 1 Chiari malformation, a structural abnormality of the posterior fossa causing cerebellar tonsillar descent through the foramen magnum. Symptoms most commonly present in adolescence or early adulthood. Type II and type III are present from birth; type II is associated with spina bifida and type III, the most severe form of ACM, is often diagnosed at prenatal ultrasound and has the highest mortality rate [6]. A number of studies have investigated the degree of cerebellar ectopia necessary to make a diagnosis of Chiari I malformation. Aboulezz et al. in 1985 reported that the average distance of the tonsillar tips was 2.9 ± 3.4 mm above the foramen in 82 patients without posterior fossa abnormality, and 10.3 ± 4.6 mm below the foramen in 13 patients with Chiari malformations (p < 0.005). They suggested that extension of the tonsils up to 3 mm below the foramen magnum was normal, may be considered of borderline significance between 3 and 5 mm, and was pathologic when it exceeds 5 mm [7]. In their study into the radiographic findings of Chiari malformation, Milhorat et al. reported that while tonsillar herniation less than 5 mm does not exclude the diagnosis, the cerebellar tonsils were at least 5 mm below the foramen magnum in 91% of symptomatic patients in a study of 364 symptomatic patient, though there is no direct correlation between how low the tonsils are lying and clinical severity [8]. Meadows et al. reviewed the MRI images of over 22,000 patients [9], demonstrating that 0.77% of patients had cerebellar tonsillar ectopia of at least 5 mm, however, of these patients, 14% were asymptomatic. The author emphasized that the identifying tonsillar herniation in isolation is of indeterminate clinical significance and that this must always be considered in correlation with the patients symptoms and clinical examination findings. The frequency of spinal cavitations, such as syringohydromyelia, in Chiari type 1 patients varies in the literature between 40 and 75% [8]. The prevalence of hydrocephalus associated with type 1 Chiari malformations is ~10% [10]. Mikulis et al. describe changing position of the cerebellar tonsils with age, with cutoffs of 6 mm up to 10-year old, 5 mm in ages 10–30 years, and 4 mm in those aged 30 years and older [11].

ACM may occur when a craniospinal pressure gradient develops naturally or iatrogenically. Conditions in which the normal cephalic absorptive pathways are disrupted whilst normal absorptive pathways in the spine are maintained may lead to a downward pressure gradient. This can occur iatrogenically secondary to lumbar punctures or CSF drainage, for example if patient has a VP shunt in situ [12–14]. ACMs can also occur following spontaneous spinal CSF leakage [15]. Spontaneous CSF leakage occurs in patients who have a combination of underlying weakness of the meninges and a minor traumatic event at the cervicothoracic junction or thoracic spine. Radiologically these patients often have images mimicking a Chiari type I malformation.

The mechanism by which ACM occurs following spinal trauma is uncertain. It is possible that posttraumatic dural tethering may result in hydrodynamic changes in CSF circulation causing a downward displacement of the cerebellar tonsils. The resolution of symptomatic ACM, syrinx and cervical cord edema in case 4 of our report after the patient underwent a cordectomy lends credence to this theory.

The incidence of symptomatic ACM is reported to be rare in patients with perioperative lumbar drainage [16]. However, it is known that an ACM can cause catastrophic results including cardiopulmonary arrest or quadriplegia when it contributes to the development of acute foramen magnum syndrome. In this scenario the cerebellar tonsils acutely compress the brainstem leading to cardiovascular and neurological compromise [12]. Given that assessment of progressive neurology may be especially challenging in the context of patients with traumatic cord injury, in the case of pediatric patients or complete cervical lesions, the authors believe that awareness of the potential association of ACM following SCI is important. Awareness of this potential association may also be important for patients who require CSF lumboperitoneal shunts, when over drainage should be avoided due to the risk of brain stem compression.

**Conclusion**

This case series is the first to describe ACM in patients following traumatic SCI. The pathophysiology of ACM following cord injury is uncertain but may result from altered CSF circulation and a subsequent disruption of the craniospinal pressure gradient. Progressive or unacceptable neurological symptoms may necessitate intervention; as described by the authors, cordectomy resulted in the resolution of a patient’s ACM, cervical cord edema and syrinx. Clinicians should be aware of the potential association between ACM and SCI as early recognition could prevent progressive neurological deterioration in this patient group.

**Compliance with ethical standards**

**Conflict of interest** The authors declare that they have no conflict of interest.

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