Chiari 1.5 malformation, accessory odontoid synchondrosis, and ventral compression: case report

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Abstract: The pathogenesis of Chiari 1 malformations has been explained in several different ways, but extensive evidence suggests a relationship between loss of volume within the posterior cranial fossa and Chiari 1 presentations. It is important to be able to differentiate Chiari 1.5 from Chiari 1 malformations as they have similar clinical presentations, but the latter have progressed further and are characterized by caudal herniations of the brain stem through the foramen magnum. Despite the similarities of presentation, Chiari 1.5 malformations have greater rates of complications following posterior decompression surgeries, which are typically performed to relieve ventral compression. An improved understanding of the odontoid synchondroses could lead to better understanding of Chiari malformations and lead to improved treatment of patients with these presentations. Here we present a rare case of an accessory odontoid synchondrosis in a patient with a Chiari 1.5 malformation and ventral compression.

Key words: Dens, Odontoid process, Anatomy, Chiari malformations, Odontoid synchondroses

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Introduction

The odontoid synchondroses are cartilaginous articulations separating the odontoid process from the body of the axis (Fig. 1) [1]. They normally become fully fused between the ages of five and seven years, usually forming a line on radiographs until 11 years; so most cases involving fracture of the synchondrosis are in children [1-3]. In this report, we present a rare case of an accessory odontoid synchondrosis in a patient with a Chiari 1.5 malformation and ventral compression.

Case Report

A 13-year-old boy presented with diplopia and occasional headaches. The symptoms were ongoing for three months and were more pronounced in upon wakening in the morning. There was no past surgical or medical history. On examination, a left abducens nerve palsy was identified and his gag reflex was absent. The remainder of his motor and sensory exams were normal as was his cerebellar and deep tendon reflex exams. Magnetic resonance imaging (MRI) revealed a Chiari 1.5 malformation without syringomyelia (Fig. 2). Additionally, ventral compression (8 mm) of the anterior brainstem was noted. There was no hydrocephalus. An additional synchondrosis of the odontoid process, superior to the normal subdental synchondrosis, was identified (Fig. 2). This synchondrosis

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was located at the midportion of the odontoid process inferior to where the apical synchondrosis would have been located and gave the odontoid process and elongated appearance. A posterior fossa decompression procedure was offered but was declined by the family.

Discussion

The odontoid synchondroses are present at birth and serve as cartilaginous separations between the odontoid process, neural arches and body of the axis [1]. They are categorized as dentocentral (subdental), between the odontoid process and the body of the dens, neurocentral, between the odontoid process and the neural arches, and apical just below the apex of the odontoid process [1, 2, 4]. Typically, the odontoid process has fused fully with the body and the neural arches by 5 to 7 years of age, leaving adults with a dentocentral remnant within the cancellous bone that can be visualized on MRI as a hypointense ring between the superior surface of the body of C2 and the inferior surface of the odontoid process [1-3]. The apical synchondrosis fuses by ten years [3]. Owing to these fusion patterns, fractures of the odontoid synchondroses usually affect children under the age of 10 years.

Our case shows no sign of traumatic injury, but it displays a Chiari 1.5 malformation. Chiari 1.5 malformations exhibit similar tonsillar herniations to Chiari 1 malformations, but they are distinguished by caudal brainstem herniations through the foramen magnum [5-8]. The diagnosis of Chiari 1.5 malformation has posed challenges because of the similarity in clinical presentation to Chiari 1 malformation, including (but not limited to) dyspnea, jaw pain, drop attacks, dysarthria, opisthotonos, perioral numbness, negative gag reflex, lethargy, headaches, and limb weakness [6-8]. Despite the similarity in presentations, it is important to identify Chiari 1.5 malformations early and differentiate them from Chiari 1 malformations as their operative outcomes can differ [5, 6]. In a study of 130 pediatric patients with Chiari malformations, Tubbs et al. noted a difference in the incidence of unresolved or recurring episodes of syringomyelia following posterior fossa decompression between patients presenting with Chiari 1 (6.8%) and Chiari 1.5 (13.6%) malformations [6, 7, 9].

Many different explanations of the pathogenesis of Chiari malformations have been proposed, but because a wide variety of anomalies can arise, no definitive account has yet been agreed [10]. Nevertheless, extensive evidence demonstrates that reduction of the posterior cranial fossa volume contributes to hindbrain herniation and the development of a Chiari 1 malformation in many cases [11, 12]. Such decreases in volume have been associated with developmental abnormalities disrupting the rotation of the tentorium cerebelli and petrous bone during the embryonic, fetal, and postnatal periods, leading to an overcrowded posterior cranial fossa [11].
Although ventral compression can be seen in patients with Chiari 1.5 malformation, commonly associated vertebral variations include atlanto-occipital fusion and Klippel-Feil syndrome and not an accessory synchondrosis of the odontoid process as seen in the present case (Fig. 2).

Furthermore, there is a high incidence of craniovertebral junction (CVJ) abnormalities among the Chiari 1 malformation population. Many hypotheses have linked the development and migration of the odontoid process, axis, and clivus to shrinkage of the posterior cranial fossa, exacerbating the conditions and causing further tonsillar herniation [5, 13]. Studies of pediatric populations have revealed a correlation between retroflexion of the odontoid process and the presence of a Chiari 1 malformation or other skeletal disorder with CVJ abnormalities [1, 14, 15]. In fact, excessive retroflexion of the odontoid process has been classed as a causative agent of ventral compression among Chiari 1 patients and this was seen in the present case [15]. Currently, it is standard protocol to perform a posterior decompression on Chiari 1 patients exhibiting ventral compression [15]. In view of the differing surgical outcomes between Chiari 1 and 1.5 patients, it is essential for physicians to distinguish between them and consider their variable anatomy [6, 7, 9, 15].

Although variations and pathology such as atlanto-occipital fusion and Klippel-Feil syndrome have been reported in a patient with a Chiari 1.5, to our knowledge, an accessory synchondrosis of the odontoid process with a Chiari 1.5 malformation has not been described in the literature. Owing to the lack of available information, the clinical implications of a duplicated odontoid synchondrosis are still unclear, and it is difficult to be certain that a concomitant Chiari malformation contributes to its development. Elucidating this mechanism could give greater insight into the pathogenesis of Chiari malformations and provide us with a more comprehensive plan for diagnosing and treating patients with these presentations. Since there are differences between the surgical outcomes of Chiari 1 and 1.5 malformation patients, differentiating the two is imperative for ensuring the greatest patient benefit.

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Conceptualization: JI, RST. Data acquisition: ASD, RST. Data analysis or interpretation: DH, CJB. Drafting of the manuscript: DH, JI. Critical revision of the manuscript: ASD, RST. Approval of the final version of the manuscript: all authors.

**Conflicts of Interest**

No potential conflict of interest relevant to this article was reported.

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