Case Reports & Case Series (CRP)

Management of craniosynostosis at an advanced age: Clinical findings and interdisciplinary treatment in a 17 year-old with pansuture synostosis

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Abstract

Craniosynostosis is the premature fusion of cranial sutures, occurring at a rate of approximately 1 in 2000 live births; it is usually diagnosed and treated within the first year-of-life. Some diagnoses are delayed and only detected later in childhood or adolescence when symptoms of increased intracranial pressure (ICP) arise such as headaches and vision changes. We present a case of occult craniosynostosis in which a relatively normocephalic 17-year-old male presented with debilitating headaches, optic nerve edema, and developmental delay consistent with probable ICP elevation. CT scan demonstrated pansuture craniosynostosis. Invasive monitoring confirmed increased ICP for which he underwent cranial remodeling and expansion. While the functional benefits of cranial remodeling are still vigorously debated, this patient’s headaches resolved postoperatively. Clinicians should be cognizant of cases of occult craniosynostosis, obtain the appropriate preoperative evaluations, and recognize the utility of cranial remodeling in appropriately selected patients. © 2015 The Authors. Published by Elsevier B.V. This is an open access article under the CC BY-NC-ND license.

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Introduction

Craniosynostosis is the premature fusion of one or more cranial sutures. It occurs at an overall rate of 1 in 2000 births [1]. Head shape anomalies or a syndromic diagnosis usually alerts the parent or pediatrician early in infancy to the potential underlying bony pathology. Single suture synostoses result in characteristic head shapes, which facilitate clinical diagnosis. Surgical correction is typically performed within the first year-of-life to expand and normalize cranial shape to accommodate the rapidly growing infant brain [2]. One central controversy in craniofacial surgery is whether cranial remodeling is an aesthetic or functional procedure. Evidence supports opinions on both sides of this contentious debate [3,4]. Cases of untreated craniosynostosis provide some insight into the natural history of this condition [5]. We present a case of pansuture craniosynostosis with relative normocephaly, which eluded timely diagnosis and treatment. These rare cases serve as a reminder that craniosynostosis can be associated with severe refractory headaches and visual changes in the absence of significant head shape anomalies.

Case report

A 17-year-old male presented with recurrent debilitating headaches and ear and mastoid pain, leading to frequent school absence, grade repetition, and eventual expulsion. As an adolescent he was diagnosed with a Chiari malformation, but the finding of pansuture synostosis remained elusive. He underwent Chiari decompression via a suboccipital craniectomy with a Cl laminectomy. The dura was opened in a Y-shaped fashion. Tonsil tips were found to be ischemic and were elevated with coagulation. Duraplasty using an allogdem patch was performed. The patient was 12-years-of-age at the time of Chiari decompression and did not experience improvement in his headaches postoperatively. He was then evaluated by a second neurosurgeon and was diagnosed with occipital neuralgia and referred for medical management of his headaches. Oral medications failed to improve his symptoms, but illicit marijuana provided some relief. His early medical history was unremarkable; the product of a full-term vaginal delivery, his family reported a normal head shape at birth. He first began to report headaches at 10-years-of-age. Family history was notable for migraines, particularly in his maternal grandmother, but no history of craniofacial anomalies.

At presentation his head circumference was 55 cm (<50th percentile). He had subtle contour abnormalities and asymmetry of the head, with some flattening of the left frontoparietal region, mild
bilateral mastoid bulging, and a prominence of the right frontal bone. Computed tomography (CT) and magnetic resonance imaging (MRI) revealed pan-suture synostosis and a bony spur at the site of his prior Chiari decompression (Fig. 1). The bony spur was not considered clinically significant as MRI demonstrated adequate decompression with establishment of the cisterna magna. Ophthalmologic evaluation revealed blurring and elevation of the inferior optic disc margins in both eyes with absence of spontaneous venous pulsations, concerning for bilateral optic nerve edema secondary to elevated intracranial pressure (ICP) (Fig. 2A). Optical coherence tomography (OCT) was also performed to determine the thickness of the retinal nerve fiber layer (RNFL), which can aid in assessing the degree of optic nerve edema. OCT revealed a slight blurring of the inferior margin of the optic disc in our patient, consistent with mild optic nerve edema.

Given the controversial indications for delayed cranial remodeling and the potential for confounding causes of his headaches, formal intraparenchymal pressure monitoring was performed using the Codman © Microsensor for 48 h. This revealed elevated intracranial pressures in excess of 20 mm Hg, A wave activity and rhythmic B waves consistent with elevated ICP (Fig. 2B). These findings coupled with the refractory headaches and abnormal ophthalmological exam provided the indication for cranial vault expansion.

The patient underwent craniectomy and posterior cranial vault expansion. Intraoperatively, he was found to have a “tight” brain with immediate expansion and relaxation of the brain upon craniotomy. Preoperative and intraoperative findings were consistent with pansuture synostosis (Fig. 3A). Cranial reconstruction was performed using stair-step osteotomies to both significantly expand the cranium as well as to maintain cortical coverage (Fig. 3B and C) [6]. The patient spent one night in the ICU and recovered well without any postsurgical complications. He was discharged on postoperative day four. Postoperative CT demonstrated an 18% increase in intracranial volume with complete expansion of the brain to fill the new space (Fig. 4). The change in intracranial volume reflects the bony changes, and was calculated as the difference in volumes contained by the cranium on the pre and post op CT scans.

At his four-week follow-up his headaches had abated, though he still had intermittent mastoid pain. Follow-up ophthalmology evaluation four weeks after surgery demonstrated no change in the subtle optic nerve edema detected preoperatively. At six months follow-up he remained free from headaches, but noted occasional discomfort in his ears and mastoid areas bilaterally.

### Discussion

Cases of occult craniosynostosis with relative normocephaly have been previously reported. Martinez-Lage and colleagues documented two such cases in which the diagnosis of craniosynostosis was missed at birth due to negligible skull deformity [7]. Both patients presented years later with elevated ICP and underwent cranial expansion, which alleviated their symptoms.

Our case resembles these accounts of delayed presentation, though this child was older at presentation and had previously undergone Chiari decompression for his symptoms. Our patient had a relatively normal head shape at birth, with only subtle changes apparent in adolescence. The timing of his suture fusion is unclear, though retrospective review of his CT scans at 11-years-of-age demonstrates the synostoses.

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**Fig. 1.** Pre-operative superior (A) and lateral (B) views of the calvarium on CT scan demonstrating suture fusion and a bony spur at the site of a prior Chiari decompression. T1-weighted MRI scan (C) revealing a Chiari malformation five years prior to his craniofacial workup and post-operative MRI (D) demonstrating decompression of the Chiari malformation.
Ophthalmologic findings in cases of elevated ICP can be absent or difficult to detect. The initial ophthalmologic workup included a fundoscopic exam as well as an optical coherence tomography scan, which quantifies retinal nerve fiber layer (RNFL) thickness. This latter metric has been used as a noninvasive surrogate for conventional intracranial pressure measurements in children with pseudotumor cerebri; RNFL thickness was increased by nearly 75% on average when compared to measurements from healthy control subjects [8]. Optical coherence tomography showed a slight increase in inferior RNFL thickness in our patient, consistent with clinically observed mild optic nerve edema. One paper suggested that optic nerve edema is a sensitive indicator of elevated ICP in children with craniostenosis over eight years of age, though the specificity decreases with increasing age [9]. Nevertheless, many patients with elevated ICP have a normal ophthalmological exam. In this case, clinical suspicion of elevated ICP in children with craniostenosis was over eight years of age, though the specificity decreases with increasing age [9]. Nevertheless, many patients with elevated ICP have a normal ophthalmological exam. In this case, clinical suspicion of elevated ICP in children with craniostenosis was over eight years of age, though the specificity decreases with increasing age [9].

Furthermore, MRI studies cannot definitively rule out conditions such as idiopathic intracranial hypertension. MR venous angiogram can show stenosis of the distal areas of the transverse sinus, bilaterally or unilaterally. This is certainly not a criterion “sine qua non” to establish the diagnosis of IIH. The diagnosis is supported if found, but is not a consistent finding. MRI can show an empty sella as a sign of chronic pressure. However, this is neither a unique nor a specific finding for IIH. Although IIH can occur in conjunction with Chiari malformations, this patient did not fit the typical clinical situation of IIH, as he was a thin male. Moreover, the coexistence of “IIH-like pathology” in craniostenosis and the associated skull base deformities provide risk factors for venous outflow obstruction. Therefore, an abnormal finding in the MRV studies could be attributed to the presence of craniostenosis rather than a distinct IIH condition.

Once elevated ICP has been identified, cranial expansion can be recommended, though there are limited data on surgical outcomes in this population. Scott and colleagues published their cohort of older children with craniostenosis experiencing symptoms of increased ICP and found that cranial vault remodeling alleviated symptoms in the vast majority of patients [5]. At 6-month follow-up our patient also reported dramatic functional improvement with alleviation of his headaches. This case demonstrates that craniosynostosis may be occult, without microcephaly or significant contour anomalies. Although radiographic and ophthalmologic findings may suggest elevated ICP, intraparenchymal monitoring is more definitive, particularly when the findings are as dramatic as those seen here. In this case, the functional sequelae of the synostosis with elevated ICPs were significant, and cranial expansion resulted in considerable improvement in symptoms. While treatment of older children remains somewhat controversial, cases such as this suggest that cranial expansion and remodeling can be functionally beneficial, even at an advanced age.
Fig. 4. Superior (A) and lateral (B) CT scans of the calvarium post-operatively.

References