The number of infants with head shape deformities has risen over the past several years, likely due to increased awareness of the “Back to Sleep” program. Most of the time, the head deformity is simply positional plagiocephaly, a benign condition that does not require surgical intervention. However, some deformities are caused by craniosynostosis, a condition where skull sutures fuse prematurely.

Craniosynostosis must be ruled out because it has significant medical implications. Early diagnosis and treatment of craniosynostosis improves outcomes and reduces possible adverse effects on brain development.

**Overview: Craniosynostosis vs Positional Plagiocephaly**

**Craniosynostosis**
The sutures of the skull serve as growth plates that allow the skull to grow as the brain grows. In craniosynostosis, one or more of the sutures close early. The skull then attempts to grow parallel to the fused suture, rather than perpendicular to it, causing an abnormal head shape.

Normal sutures include the metopic (m), coronal (c), sagittal (s), lambdoid (l) and squamosal (sq).

In craniosynostosis, the anterior fontanel (af) may be open or closed.

Occurring in one out of 2,000 – 2,500 live births, craniosynostosis may be spontaneous, syndromic or familial and can present in many different forms. Left untreated, craniosynostosis can result in further cranial deformity and potentially an overall restriction in head growth, with secondary increased intracranial pressure. It can also lead to psychosocial issues as the child interacts with peers during development. Multiple types of surgical intervention for craniosynostosis exist. Early referral to a pediatric craniofacial center is essential as it allows all options to be offered.

**Positional Plagiocephaly**
In positional plagiocephaly, the skull sutures are not fused. This head shape deformation, typically of the back of the head, is caused by repeated pressure to the same area. It usually develops when a child prefers to lay his or her head on the same spot. Torticollis, or gross motor delay, can also contribute, and these conditions may be responsive to physical therapy.

Effects of positional plagiocephaly are primarily cosmetic, as the condition does not impact brain growth or development. Positional plagiocephaly can often be treated by the child’s primary care physician. It usually does not require a referral to a specialist, however specialist consultation is always available if there are concerns.

**Positional Plagiocephaly** is different than craniosynostosis. In positional plagiocephaly, the ear and possibly forehead on the side of the posterior flattening are displaced anteriorly, giving the head a parallelogram shape. Flattening may occur on the right, left, or center occiput. Symmetric positional plagiocephaly can sometimes be difficult to distinguish from bilateral coronal craniosynostosis and imaging may be considered.

**Typical Craniosynostosis Presentations**
Most often, an examination of head shape can differentiate craniosynostosis from positional plagiocephaly. Head shapes are best viewed by standing above and looking down at the top of a child’s skull. In the case of an unclear diagnosis, a 4 view skull x-ray series or head CT with three dimensional reconstructions of the bone windows can be used to see if sutures are fused. If the sutures are patent, then there is no craniosynostosis.

**Sagittal craniosynostosis, the most common non-syndromic form, causes a long and narrow head. There may be prominence, or “busing,” of the forehead and/or occiput.**
Metopic craniosynostosis causes a triangular shape to the forehead when viewed from above. Eyes may be abnormally close together. **Metopic ridging** without the triangular shape is a normal variant and does not require surgical correction.

Unilateral coronal craniosynostosis causes a rotated appearance to the face with flattening of the forehead and elevation of the orbital roof on the affected side along with rotation of the nose. Sometimes the anterior fontanel is somewhat displaced to the contralateral side.

Lambdoid craniosynostosis is very rare and the only type that would cause flattening in the back of the head similar to positional plagiocephaly. With lambdoid craniosynostosis, however, the ear and possibly forehead on the side of the posterior flattening are displaced posteriorly, giving the head a trapezoidal shape. With positional plagiocephaly, the ear and forehead displacement is anterior. In addition, lambdoid craniosynostosis can cause inferior displacement of the ipsilateral ear and mastoid which is not present in positional plagiocephaly.
Other Clinical Features of Craniosynostosis

Risk factors for craniosynostosis include fetal constraint (nulliparity, plurality, macrosomia), low birth weight, preterm delivery, maternal valproate use and shunted hydrocephalus. While the majority of cases (approximately 82%) are not syndromic or familial, craniosynostosis can be a feature of many different genetic syndromes. It is important for the child to be examined carefully for signs of an inherited genetic disorder, such as facial anomalies, limb defects, hearing loss, or cardiovascular malformations.

Craniosynostosis Surgical Treatment

The goals of craniosynostosis surgery are to unlock and reshape the bones. Historically, craniosynostosis has been treated using surgical methods that involve an incision from ear to ear and the removal, reshaping, and reattachment of affected bones. However, at Nationwide Children’s, advances in technology are allowing us to conduct more of these procedures in a minimally invasive manner.

Traditional open surgery

With traditional surgery, the procedure lasts approximately four hours and is performed in conjunction with a craniofacial plastic surgeon. A blood transfusion is usually necessary. The child is typically observed overnight in the ICU and then an additional three days on the regular neurosurgical floor before discharge. Pronounced periorbital edema is normal, but generally resolves before discharge. To reduce surgical risk, this procedure is performed around the age of six months. Younger infants are very unlikely to experience increased intracranial pressure before then. Syndromic and multiple suture cases are more frequently treated using traditional surgical techniques. Because reshaping occurs at the time of surgery, no further interventions are required.

Minimally invasive surgery

Minimally invasive surgery involves two small incisions and the removal of only the fused suture to unlock the bones. The surgery lasts approximately one hour and rarely requires a blood transfusion. Postoperatively, the child is observed overnight on the regular neurosurgical floor, and is then discharged. Usually there is no periorbital swelling. Minimally invasive surgery produces the most successful outcomes when performed on children before the age of six months.

With minimally invasive techniques, reshaping occurs postoperatively with the assistance of either a cranial molding helmet or implanted custom springs.

The cranial molding helmet has a hard outer shell with moldable foam on the inside. It is worn 23 hours per day until the child’s first birthday. The helmet does not press the skull into shape, but rather directs the growth of the skull into a more normal shape. Because the helmet relies on the high rate of skull growth in the first year of life, helmet-assisted surgery should be done between 10 to 14 weeks of age. The helmet requires frequent visits to an orthotist, but no additional surgery.

Stainless steel cranial expander springs are implanted after the fused suture is resected. The springs are then removed three months later. The level of spring tension is selected based on the patient’s age, bone thickness, and deformity severity. Spring-assisted surgery is performed between the ages of three to six months. The springs require a second surgery for removal but not the use of the helmet.

Cranial distraction

In rare cases, when the majority of sutures are fused, cranial distraction can be employed to create more intracranial volume. After the bones are unlocked, distractors are implanted spanning the bone cut. At a rate of 1 mm per day, the sides are separated by turning the screw that connects them using posts that project through the skin. After 30 days, distraction osteogenesis results in 3 cm of new bone. The posts are then removed and the skin closed in the office. Three months later, the distractors are removed at a second surgery.

Prevention and Treatment of Positional Plagiocephaly

Positional plagiocephaly may be prevented by regularly checking head shape and educating parents during well visits. Strategies to prevent positional plagiocephaly include encouraging parents to:

- Increase the amount of supervised tummy time for their infant while he/she is awake.
- Place colorful items on the outside of the crib to encourage baby to look in the direction opposite the flattened side.
- Vary the position of the head when the child is being held.

Parents should be advised to continue placing their baby on his or her back to sleep, even if the child has positional plagiocephaly. The flattening is likely to diminish as the child obtains gross motor milestones and lies on the area less. It should also be noted that after the age of one, it becomes increasingly difficult to modify a child’s head shape non-invasively. If head shape is not responding to traditional conservative treatment, parents can discuss the pros and cons of using a helmet with their pediatrician.
Differentiating Craniosynostosis from Positional Plagiocephaly