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Headlines Craniofacial Support
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1 What causes Craniosynostosis? - a general discussion with a focus on genetic aspects (syndromes)
2 Apert Syndrome
3 Non-syndromic Craniosynostosis
4 Craniofacial Surgery
5 The Surgical Treatment of Hand Anomalies associated with Craniofacial conditions
6 Crouzon Syndrome
7 Pfeiffer Syndrome
8 Saethre-Chotzen Syndrome
11 Glossary of Terms associated with Craniosynostosis
12 Coping with Facial Disfigurement
13 The Genetic Background to Craniosynostosis
14 Breathing Problems in Craniofacial Syndromes
15 Muenke Syndrome
16 Eye Aspects of Craniofacial Conditions
17 Occipital (Positional) Plagiocephaly
18 Craniofrontonasal Syndrome

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<table>
<thead>
<tr>
<th>Operation</th>
<th>Age</th>
<th>Indication</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cranioplasty</td>
<td>Infancy</td>
<td>Skull expansion and remodelling, for cosmetic benefit and to relieve</td>
</tr>
<tr>
<td>Shunt surgery</td>
<td>Childhood</td>
<td>Neurosurgical operation to reduce intracranial pressure</td>
</tr>
<tr>
<td>Facial advancement</td>
<td>Childhood</td>
<td>To protect the eyes, protect against breathing difficulty, and provide</td>
</tr>
<tr>
<td></td>
<td>Adolescence</td>
<td>cosmetic benefit. Often preceded and followed by a programme of</td>
</tr>
<tr>
<td>Hand surgery</td>
<td>Infancy</td>
<td>Operations as indicated for functional problems of thumb, grip,</td>
</tr>
<tr>
<td></td>
<td>Childhood</td>
<td></td>
</tr>
<tr>
<td>Choanal dilation</td>
<td>Childhood</td>
<td>ENT procedures to improve the airway, treat chronic ear infection, aid</td>
</tr>
<tr>
<td>Grommet insertion</td>
<td>Childhood</td>
<td>hearing</td>
</tr>
<tr>
<td>Bone-anchored hearing aid</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Squint surgery</td>
<td>Childhood</td>
<td>To correct ocular squint and improve vision, Tarsorrhaphy may be used to</td>
</tr>
<tr>
<td>Tarsorrhaphy</td>
<td>Childhood</td>
<td>protect against exposure damage to</td>
</tr>
</tbody>
</table>

**Further reading**

*Assessment and Treatment of Craniosynostosis.* Thompson D, Jones B M, Hayward R D, Harkness W; British Journal of Hospital Medicine 1994, 52 (1); 17-24.


Written by Jonathan Britto, Craniofacial Research Fellow, Great Ormond Street Hospital for Children NHS Trust
The hand and feet syndactyly and digit problems may come to need surgical care by the hand surgeon. Indications for such surgery are functional rather than cosmetic, and are aimed at improving thumb function and grip. Operations on the feet are reserved for those problems interfering with gait and wearing shoes.

The team is made complete by the geneticist, psychologist, speech and language therapist, respiratory care specialist, and specialist nursing staff.

Non-surgical aspects
Developmental delay in the absence of raised intracranial pressure in Pfeiffer syndrome is uncommon. The psychology and language therapy teams have many means of identifying and treating developmental delay early, and their important role in the overall care of the child is emphasised.

Though many new cases are spontaneous, Pfeiffer syndrome can run in families. When this happens it does so in a ‘dominant’ manner. The genetic basis of the syndrome is one of the recent research discoveries. Although the inheritance is ‘dominant’, it must be remembered that the expression of the disease in the child is variable, and most children of a Pfeiffer family will not be severely affected. The geneticist will advise about risk in subsequent generations of a family.

Summary
Pfeiffer syndrome is an inherited syndrome affecting craniofacial growth and development. The care of the child with Pfeiffer syndrome is multi-disciplinary, involving the coordinated expertise of many clinical teams. Surgical care is staged throughout life from infancy to late adolescence, and may follow the following pattern as clinical circumstances arise.

The Clover Leaf anomaly (Kleebattschädel)
This is an uncommon feature which may accompany the Pfeiffer syndrome. There is a trilobar or clover leaf head shape, which usually is the result of pan-craniosynostosis involving multiple sutures. The brain is severely compromised, and intelligence can be greatly impaired. The ears are displaced downwards and the mid-face severely regressed. High intracranial pressure is common. In addition, there may be a cluster of other craniofacial features. The child with the clover leaf anomaly may have a reduced lifespan related to the multiple and severe problems, despite multidisciplinary intervention by many specialist teams.

Introduction
Pfeiffer syndrome is an inherited syndrome of craniofacial dysmorphology, or abnormal craniofacial appearance, which was originally described in 1964 and is now well recognized.

Children who have Pfeiffer syndrome have a range of problems of variable severity, from predominantly cosmetic facial symptoms to severe symptoms affecting breathing, feeding, vision and brain development.

The child with Pfeiffer syndrome usually enters a coordinated programme of care involving many different clinical specialities integrating their various expertise, which often continues from birth to the later teenage years. In addition, there is an on-going programme of research into many aspects of Pfeiffer and its related syndromes, to constantly investigate and update the services that the specialist team provide.

The child with Pfeiffer syndrome
Pfeiffer syndrome predominantly affects the appearance of the head and face. The hands and feet are also involved, and may require surgical care.

The skull, or calvarium, is made up of flat plate-like cranial bones which are connected by seam-like joints, or cranial sutures. There are many such sutures, but the most clinically important are

- the metopic and sagittal sutures running from front to back and interrupted by the anterior fontanelle (soft spot),
- the coronal sutures running from side to side from the anterior fontanelle to the temple and
- the lambdoidal sutures running from the posterior fontanelle to the back of the base of the skull

Neighbouring cranial bones are thus mobile against each other, and this first allows normal birth, and then growth of the brain inside the skull without restriction.
During normal childhood and into adulthood, the sutures fuse, becoming bone in a seemingly pre-programmed fashion, protecting the brain within. The growth of the face and skull are, of course, closely integrated, and facial bones are also joined by sutures which slowly fuse throughout life.

In Pfeiffer syndrome, either or both of the skull and face may be affected. In the skull, the cranial sutures may fuse prematurely, and this is called craniosynostosis or craniofacial synostosis. This alters the pattern of skull growth, and thus the shape of the skull, with consequences for the developing brain. There are characteristic skull shapes depending upon the pattern of sutural fusion. Pfeiffer syndrome may involve premature fusion of any combination of cranial sutures. Common terms for the resulting head shapes are brachycephalus – giving rise to a boat-shaped skull and turricephalus – a tower-shaped skull; and these may result in different pressures on the growing brain.

A severe form of Pfeiffer syndrome is characterized by the ‘clover leaf’ deformity or klebbatschädel (see page 6).

Craniosynostosis begins during pregnancy or the first year of life and is complete by three years of age. Raised intracranial pressure may become a clinical concern.

In the face, the commonest features are a regressed mid-face and shallow orbits (eye sockets), which may be present at birth or become more evident as the childhood progresses. The arrangement of the teeth, or dentition, is also affected, and this requires specialist orthodontic care, which is dependent on good general dental care at home. Rarely, there may be palate problems. Seen from the side the face has a concave appearance, and the shallow orbits result in prominent eyeballs or proptosis. Occasionally the nasal passages are completely blocked, called choanal atresia, resulting in the need for surgical intervention.

The hands and feet in Pfeiffer syndrome are involved to variable degree. The thumbs and big toes are broad and deviated toward the mid-line. There may be mild soft tissue webbing, or syndactyly, between the second, third and fourth digits of either/or both hands and feet. The digits may be short and misshapen, with consequences for an adequate grip (hand) or footwear (feet). Anomalies of other systems have been described, but are very uncommon.

Clinical problems and programme of care

The child with Pfeiffer syndrome may thus have a range of clinical problems. Although the head shape is often the most striking initial feature, from the outset the major concerns are the ease of breathing and potential feeding problems.

The regressed mid-face, or maxillary hypoplasia, results in a small larynx and pharynx behind the nose and mouth. This restricts the passage of air into the trachea (windpipe) and lungs, and causes respiratory distress, particularly at night when snoring and snuffling can interrupt sleep. The degree of airway obstruction and quality of sleep is assessed by a ‘sleep study’, and if necessary, treatment takes the form of CPAP (Continuous Positive Air Pressure) devices at home or surgical intervention. Similarly, the passage of food is restricted and regurgitation may result in aspiration of food into the lungs.

The shallow orbits and proptosis may threaten the cornea, or surface of the eyeball, with exposure keratitis; and surgical measures may become necessary to protect the exposed eyes. All children have regular ophthalmic review, as other ocular problems may uncommonly occur. Ear, nose and throat (ENT) follow-up is also recommended, as some children have hearing difficulties, and grommets may be advised to treat chronic infections and improve hearing. The common surgical approaches are given in the table overleaf.

The abnormal skull shape may require surgery to protect the constricted brain and help relieve raised intracranial pressure, which is most commonly revealed by headaches or visual changes identified by the ophthalmologist. The aim of this surgery called cranioplasty is to provide a more normal head shape and increase the volume of the skull. Examples of cranioplasty include frontal and fronto-orbital advancement, vault expansion and frontal or posterior remodelling. For more information, see Headlines leaflet Craniofacial Surgery HL4). Pressure is thus taken off the growing brain. Another surgical method of reducing intracranial pressure is to insert a ventriculoperitoneal shunt, and either or both may be used at different times during childhood, as the skull shape of Pfeiffer syndrome may be only one of many contributing factors to raised intracranial pressure.

At the current time it is impossible to predict how the skull and face abnormalities will progress as the child grows, and surgical decisions are made in the light of circumstances as they arise. In addition, ‘variability of expression’ characterizes the condition – some children may have mid-face problems only, others may have craniosynostosis only, whilst others may have severe craniosynostosis and mid-face regression together. Surgery may be indicated for cosmetic reasons only, or for the more serious conditions described.

Any surgical intervention is closely planned with all the teams involved. The surgical side of the craniofacial team usually consists of the craniofacial surgeon, neurosurgeon, ENT and ophthalmic surgeons. In addition the orthodontist co-ordinates very closely with the surgical teams.