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About this book

This booklet, *Craniopharyngioma*, aims to give you a basic understanding of how a craniopharyngioma is diagnosed, the types of treatment available and the sorts of problems you may encounter. It has been written to give you a brief overview of the condition and not all of the information provided will be relevant to everyone with a craniopharyngioma.

We encourage you to discuss any additional questions or areas of concern with your doctor after reading this booklet.

Merck Serono Australia is proud to bring you this booklet from the *Hormones and Me* educational series. We hope that you find it a valuable and helpful resource.

This booklet was revised in 2012 with the help of Dr Shubha Srinivasan (The Children’s Hospital at Westmead, NSW, Australia) – a Paediatric Endocrinologist specialising in childhood endocrine disorders and a member of the Australasian Paediatric Endocrinology Group (APEG).

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This booklet was first updated and reproduced for Australian and New Zealand readers in 2000 by Dr Jim Penfold, (Women's and Children's Hospital, SA, Australia). Special thanks to the original authors and editors, Dr Catherine DeVile (Great Ormond Street Hospital for Children, UK), Dr Richard Stanhope (Great Ormond Street Hospital for Children and the Middlesex Hospital, UK), Mrs Vreli Fry (Child Growth Foundation, UK), Ms Janis Clayton (Serono Laboratories (UK) Ltd, UK) and the British Society of Paediatric Endocrinology (BSPE).
**Introduction**

A craniopharyngioma is a rare form of brain tumour. This particular type of tumour is present from before birth. It may grow fast or slowly and might cause problems in childhood or not until later in adulthood.

A tumour is a collection of cells that are growing abnormally. Craniopharyngiomas are benign tumours, which means they do not spread to other parts of the body. However, as they enlarge they may push on other sensitive areas of the brain which can cause problems.

“Craniopharyngiomas are rare, but are one of the most common types of childhood brain tumour.”

Craniopharyngiomas have now been associated with 2 gene mutations:

CTNNB1 gene mutation is seen in the adamantinomatous form in childhood and adolescence. This is not currently able to be treated medically. Surgery remains the first option for treatment.

BRAF V600E mutations are seen with the papillary form in some adult craniopharyngiomas. These tumours may be able to be treated with a form of modern chemotherapy.

Although craniopharyngiomas are uncommon they account for around 9% of all brain tumours in children. This booklet focuses on childhood craniopharyngiomas, but some of the information is also relevant to adults with this condition.

*A craniopharyngioma is a rare type of brain tumour. It is NOT cancerous and will NOT spread to other parts of the body.*
What is a Craniopharyngioma?

A craniopharyngioma is a tumour that forms in the brain close to the hypothalamus and pituitary gland (diagram 1). It is thought to form early on in the developing baby, when it is still in the womb. No one knows why this happens, but it is not thought to be linked to taking medications or being ill during pregnancy. The condition is not hereditary so is not passed on from parents or grandparents.

The tumour is made up of solid parts and cysts which are filled with a thick fluid. Occasionally, a craniopharyngioma may be entirely fluid–filled or more rarely, completely solid. These tumours are often very ‘sticky’ and they cling to surrounding tissues which can make them difficult to remove completely.

Although present from birth, the impact of a craniopharyngioma may be felt at any age – from birth to old age. Symptoms start to appear when there has been a change in the size of the tumour and it begins to press against other areas of the brain. The most common of these ‘pressure’ symptoms are headaches and disturbed vision.

“Symptoms, treatment and outcomes will vary from person to person.”

Children are often diagnosed with a craniopharyngioma between the ages of 5–10 years, and they are slightly more common in boys than girls.
A Little About the Brain …

When you have a craniopharyngioma many of the symptoms you experience will be related to the pressure the growing tumour exerts on the delicate surrounding areas of the brain. To help you understand why these symptoms occur it is useful to know a little more about the different parts of the brain that can be affected.
The hypothalamus is a very important part of the brain that controls many essential body functions including:
- body temperature
- hunger and thirst
- sleep
- emotional behaviour
- memory.
The hypothalamus is also the main ‘communication centre’ for the pituitary gland. It produces chemical ‘signals’ or hormones that travel via the bloodstream to the pituitary gland. These ‘signals’ then control the production and release of further hormones from the pituitary gland, which in turn control other glands and organs around the body.

The pituitary gland is linked to the hypothalamus by the pituitary stalk. It is about the size of a pea and is divided into two lobes – the anterior (front) and posterior (rear) lobes. These lobes produce different hormones that control many important functions and other organs around the body including:
- growth
- development in puberty
- stimulation of the thyroid gland
- stimulation of the adrenal glands for a normal response to stress
- fluid regulation.
The different hormones released by the pituitary gland and their target organs are summarised in diagram 2.
Optic Nerves
Optic nerves form the connection between our eyes and our brain. They carry all the important visual information from the back of the eye (retina) to the visual centres of the brain. The two optic nerves (one from each eye) cross over each other at a point in the brain just above the pituitary gland (optic chiasm).

Ventricles of the Brain
Ventricles are spaces inside the brain containing a constantly circulating fluid known as cerebrospinal fluid (CSF). The CSF flows through the ventricles via channels and openings, circulates over the surface of the brain and eventually drains back into the blood–stream.
Symptoms

Symptoms produced by a craniopharyngioma vary depending on the tumour’s size and location and the amount of pressure it is exerting on the surrounding structures.

Common symptoms include:

Headaches
Increased pressure from the tumour can cause the fluid channels to become blocked. The circulating CSF cannot escape and accumulates, resulting in a condition known as hydrocephalus (‘water on the brain’). The symptoms of hydrocephalus are headaches and vomiting particularly first thing in the morning or at night.

Visual Problems
Vision can be affected if the craniopharyngioma is close to the optic nerves or the optic chiasm. Pressure on the nerves can stop signals from the eyes reaching the brain and sight can become impaired or lost altogether.

Loss of eyesight is usually first noticed at the outer edges of vision, on one or both sides. If the problem is not picked up early it can eventually affect the whole field of vision. The early signs of loss of eyesight are frequently missed in children because a child can compensate by moving their head from side to side and do not notice or report the problem.
Effects on Hormone Secretion
The growing tumour can affect the pituitary gland, the hypothalamus or the connection between them. When this happens the production and release of the pituitary hormones is disrupted and the following problems may occur:

**Poor Growth**
Poor growth or slowing growth rate in children can be a problem if not enough growth hormone (GH) is produced by the pituitary gland.

**Delayed Onset of Puberty**
The changes that occur in the body at puberty (breast and pubic hair development in girls and growth of the penis, facial and body hair and muscular development in boys) are triggered by the gonadotrophin hormones. These changes will not occur if insufficient levels of these hormones are produced by the body.

**Early Onset of Puberty**
Very occasionally the growth of a craniopharyngioma can affect the hypothalamus or pituitary gland in such a way that puberty is triggered early.
**Increased Thirst and More Frequent Urination**
Sometimes a craniopharyngioma can cause pressure on the hypothalamus and the posterior (rear) lobe of the pituitary gland or the pituitary stalk. This prevents the production of the hormone vasopressin (also called anti–diuretic hormone (ADH)) which helps control water loss from the body. When this happens excessive amounts of urine are produced along with an increased thirst – causing some children to drink up to 6–10 litres of fluid per day. This condition is called diabetes insipidus (more information on this condition is available in the Diabetes Insipidus booklet from the Merck Serono Hormones and Me series).

**Other symptoms may include:**
- tiredness and frequent infections which are difficult to ‘shake off’
- intolerance to cold temperatures
- disturbed sleep – either excessive sleepiness during the day–time or increased waking at night
- weight loss or gain – with a normal, decreased or increased appetite
- behavioural problems such as becoming withdrawn, introverted, aggressive or having poor concentration
- poor balance or weakness in the arms or legs
- fits – though these rare.

Many of these symptoms can be caused by other conditions and it is important to get any symptom checked by your doctor.
Diagnosis

If your doctor suspects a craniopharyngioma to be present a combination of checks and tests will be ordered to confirm the diagnosis and exclude other possibilities. These investigations may include:

Imaging to Locate the Tumour

Brain Scans
CT and MRI scans of the head are commonly used to confirm if a tumour is present. These scans allow the brain to be seen in great detail and are used to ‘see’ where the tumour is located and which of the surrounding structures are affected. Information from these scans allows treatment, especially surgery, to be planned very carefully.

Skull X–Ray
The flecks of calcium present in the tumours make them easily seen on X–rays. However, brain scans are now becoming more widely available and give more detailed information, so head X–rays are less often taken in recent times.

Tests That Form Part of the Overall Assessment

Visual Assessment
The paediatric endocrinologist and an eye specialist will check to see if the tumour has affected the child’s sight. They will check the child’s field of vision by looking at the total area in which objects can be seen, both in front them (central vision) and to the sides (peripheral vision).

Height & Weight
The height and weight of the child will be measured to check if growth has been affected. The child’s stage of development in puberty will also be checked when appropriate.
Blood Tests
Blood tests will be carried out to check for any unusual levels of hormones.

Water Balance
If your doctor suspects your child has diabetes insipidus they may ask you to estimate how much fluid he or she drinks and urinates each day, and whether this has changed recently.

Then, if required, a test can be performed to measure the level of fluid in the body. This is done by comparing the salt concentration in the child’s blood with the salt concentration in the urine (there is a relationship between the two which should be maintained). If abnormally high levels of salt are present in the blood with a very dilute urine it may indicate that the child has diabetes insipidus. Further tests will be needed to confirm this. A high salt level occurs because excess water is being lost from the body, not because there is any abnormality of salt.

Hearing Assessment
Occasionally, hearing loss or ringing in the ears (tinnitus) can occur as an early symptom of a craniopharyngiroma and your child’s hearing may need to be checked by a specialist.

Psychological Assessment
It may also be necessary to assess the child’s stage of learning, memory and behaviour as these can be affected by the tumour and some treatments.
Treatment Options

Usually treatments will involve a combination of surgery, radiotherapy, chemotherapy and medications that help control symptoms. The types of treatments used will depend on the tumour’s location and size, as well as the age and health of the child.

Your doctor will discuss the treatment options available to you. However, most patients with craniopharyngiomas will need surgery.

Surgery
Craniopharyngiomas can be very ‘sticky’ and they can become attached to surrounding tissues. Removing the entire tumour without causing additional damage to other parts of the brain can therefore be difficult. By looking carefully at scans taken before the operation the surgeon will decide whether it is best to try and completely remove the tumour or only partially remove it. There are different ways this can be done, but it will usually require a major operation.

Sometimes a craniopharyngioma may regrow after surgery and further treatments will be needed. Surgery can be followed by radiotherapy treatment to help prevent this regrowth, but this will depend on the age of the child.

Different types of surgical procedures include:

Craniotomy
The most common form of surgery used to remove all or part of a craniopharyngioma is a craniotomy. This is a major operation that involves temporarily removing a piece of the skull in order to reach the area of the brain where the tumour is situated, so the tumour can be removed.
Trans–Nasal (or Trans–sphenoidal) Operation
If the tumour is small and confined to the area around the pituitary gland the operation can be done through the nose (trans–nasal). It may not be possible for this operation to be performed in all children. For example, the technique is not suited to very extensive tumours or a tumour in a very small child (due to the size of the nasal passages).

Cyst Aspiration
Cyst aspiration is a procedure that drains fluid from cysts that may form part of the craniopharyngioma. There may be a single cyst or several cysts that join together and press against a critical structure, causing increased pressure within the brain. A surgeon may decide to drain or aspirate the contents of a cyst before attempting to remove the tumour.

Cyst aspiration is a small operation where a hole (a burr hole) is made in the skull and a tube is passed into the cyst to drain its contents. The surgeon will then go on to perform a craniotomy at a later date. Cyst aspiration may also be done if the cystic part of a tumour regrows.

Drainage of the Ventricles
A craniopharyngioma can cause the ventricles of the brain to become enlarged due to the build–up of CSF. This happens in about one third to a half of the children with this type of tumour. It may be necessary to drain the CSF to reduce the pressure and re–establish the flow of fluid around the brain before removing the tumour.
The fluid is drained by inserting a temporary (or sometimes permanent) drainage system into the ventricles. The most commonly used system is called a ventriculo–peritoneal shunt and is shown in diagram 3. It consists of a tube (or shunt), that connects a ventricle to the abdomen so that the fluid can be safely drained away. The shunt has a valve so that the speed and pressure at which the fluid drains can be controlled.

Occasionally, a shunt may need to be inserted after the main operation to remove the tumour or if it regrows and causes obstruction of the CSF flow.
Diagram 3: Ventriculo-Peritoneal Shunt

- Ventricular catheter
- Valve
- Distal Catheter (under the skin)
- Tip of catheter in peritoneal cavity
Radiotherapy

Radiotherapy is used to target and kill tumour cells while leaving the surrounding normal tissues relatively undamaged. Although this form of treatment is more commonly used to treat malignant or cancerous types of tumours, it has been found to be effective in preventing regrowth of craniopharyngiomas.

Radiotherapy is mainly used when only part of the tumour can be removed. A course of radiotherapy can be given as soon as the child has recovered from surgery or a ‘watch and see’ policy may be adopted. This gives the option of having the radiotherapy at a later date if scans show that the craniopharyngioma is regrowing. If the surgeon is confident that the entire tumour has been removed and this is confirmed by further brain scans, then it is unlikely that radiotherapy will be needed.

“The age of the child is important when deciding whether radiotherapy is given.”

Although radiotherapy is commonly used in adults it is not generally given to a child under 2 years of age and only rarely used for those under 5 years. This is because of the potential long–term effects radiation can have on development of young children, particularly in the area of memory processing.

Newer, more refined techniques of giving radiotherapy that produce fewer side–effects are being developed all the time – so this form of treatment may become more common.

Local radiation using a radioactive compound inserted into the cyst has been used successfully in some cases.
Chemotherapy
Chemotherapy is a treatment that uses anti–cancer drugs to stop the growth of tumour cells. It is not generally used to treat craniopharyngiomas.

However, some cystic craniopharyngiomas can be treated with certain drugs that are injected directly into the cysts. For example, the antibiotic–like drug bleomycin is occasionally used to try and shrink the cystic portion of a tumour so that it is easier to remove. Bleomycin is also sometimes used to try and control cyst growth after surgery.
What Happens During Treatment?

The length of stay in hospital will vary depending on the child and what investigations and treatments they are having.

The operation to remove a tumour may be delayed (by a few days or occasionally a few weeks) if a shunt or cyst aspiration is needed. During this period the child may stay in hospital or go home – depending on the individual.

Usually the child will be in hospital for up to 1 week after the operation. Longer hospital stays may be necessary depending on the individual, the procedures used and whether any new problems develop after surgery.

It is important to monitor the child’s recovery after surgery to make sure there are no ill–effects from the operation. The child will be closely observed over the first 24–48 hours following surgery and may be taken to an Intensive Care Unit for careful observation.

Once the child has recovered from surgery any additional treatments, such as radiotherapy, can be organised. Radiotherapy treatment takes approximately 6 weeks, with daily or second daily treatments for a short time, as only small doses of radiation are given at a time (to reduce side–effects). The child does not have to stay in hospital during these treatments.
**Possible Complications and Other Treatments**

A craniopharyngioma occurs in the part of the brain which is very close to the hypothalamus and the pituitary gland. Most of the problems experienced by patients are caused by damage to these very important areas.

**Diabetes Insipidus**

When the tumour is removed the pituitary stalk may be cut, stretched or damaged and the communication system between the hypothalamus and the pituitary gland is disrupted. This means the pituitary hormones, including vasopressin, may not be produced.

If vasopressin is not produced, or not enough of the hormone is released, then the body can not maintain its normal fluid balance. This condition is called diabetes insipidus and it is a common problem immediately or in the first few days after surgery. It is therefore very important that the child’s fluid intake and output are closely monitored both during and after an operation.

Blood and urine samples may be taken several times a day for the first few days to make sure the fluid balance is correctly maintained. A very young child with diabetes insipidus is not able to explain how thirsty he or she is and thus there is a risk of becoming rapidly dehydrated without correct monitoring and treatment.

Diabetes insipidus is not related to diabetes mellitus a condition involving high blood sugar levels.
How is Diabetes Insipidus Treated?

Diabetes insipidus is treated by replacing the missing hormone, vasopressin, with a similar but synthetic medication, called an ‘analogue’. The vasopressin analogue has the same actions as the original hormone, but lasts longer and only has to be given 2 to 3 times per day. It is commonly given in the form of tablets, but can also be given as a nasal spray or injection.

In the first few days following surgery fluid balance may fluctuate and the child may have periods of feeling very thirsty and wanting to drink frequently. During this period the hormone may need to be given by an intravenous drip. Usually these problems settle down after a few days and the vasopressin, if still required, can be given orally or nasally.

Very occasionally a patient may have damage to the ‘thirst centre’ as well as having diabetes insipidus. This means that although a lot of fluid is lost the child does not feel thirsty (adipsia), and can quickly become dehydrated. If this happens the specialist will advise you on how much fluid the child needs to take each day.

Diabetes insipidus is not always permanent. Sometimes diabetes insipidus can occur in the first few days after surgery and then disappear completely. Occasionally there can be a ‘three phase’ response where diabetes insipidus occurs immediately after surgery, gets better 7–10 days later and then recurs about 2 weeks after the operation.

You will usually find out if a child will need to have long- term vasopressin treatment in the first few weeks after surgery.
Anticonvulsants
Convulsions or fits are rare in children with a craniopharyngioma. However, a child who has had a fit will be treated, with tablets known as anticonvulsants, to prevent any further fits occurring when the tumour is removed.

A small proportion of children may continue to have problems with fits or develop them later on and they may need medication for a longer time to keep the fits under control.

Steroids
Steroids are always given before and after surgery. There are two main reasons for this:

1. To help reduce the swelling and build–up of pressure within the brain.
2. To help the child cope with stress.

Adrenocorticotrophin (ACTH) is one of the hormones produced by the anterior (front) lobe of the pituitary gland. It triggers the adrenal glands to make and release a steroid called cortisol. Cortisol is a ‘fight and flight’ hormone that enables us to cope with stresses such as illness and trauma.

If the pituitary gland and pituitary stalk are damaged, either by the craniopharyngioma itself or the surgery, then the ability to produce cortisol can be lost. The body will then be unable to cope with the stress of an operation unless replacement steroids are given.
Cortisol is initially given in high doses for the first few days following the operation, partly for stress and partly to reduce local swelling around the operation site. The dose is then gradually reduced until further tests of the pituitary gland function can be carried out.

**Hormone Replacement Therapy (HRT)**

Once the tumour has been removed it is important to find out how well the pituitary gland is functioning. Approximately 2–3 months after the operation a series of blood tests will be done to find out which of the various pituitary gland hormones may be affected. The results from these tests will indicate which hormones need to be replaced. Patients who need treatment at this stage will normally need to take these replacement hormones for the rest of their lives.

Hormones that may need to be replaced include:

**Thyroid Hormone (Thyroxine)**

Thyroid stimulating hormone (TSH) stimulates the thyroid gland to produce thyroid hormone (a hormone that helps regulate the body’s metabolism). Replacement thyroid hormone treatment is given in the form of tablets.

**Growth Hormone (GH)**

GH replacement treatment may be needed if the child is growing poorly. Occasionally, after surgery a child continues to grow normally despite a lack of growth hormone. If this is the case, the child will not need growth hormone treatment.
GH treatment is given by injection and the dose calculated according to the size (either body weight or body surface area) of the child.

Currently there is no financial assistance available in Australia or New Zealand for the growth hormone treatment of adults.

**Sex Hormones**

Gonadotrophins are pituitary hormones that stimulate the gonads (ovaries in girls, testes in boys) to produce sex hormones (oestrogen and testosterone) needed to trigger the physical changes that occur during puberty.

If not enough of these hormones are being produced at the appropriate time then hormone replacement treatment will be needed to start and to continue puberty. Long–term maintenance of normal adult sex hormone levels may be required.

**Multiple Pituitary Hormone Deficiency (MPHD)**

MPHD or panhypopituitarism, is used to describe a situation where several pituitary hormones are affected. Children can develop MPHD after surgery or the pituitary gland may gradually stop producing hormones over time (particularly if radiotherapy has been used). Careful follow–up is therefore always required by a hormone specialist (endocrinologist) and these checks will need to be continued on into adult life.
IT IS VERY IMPORTANT TO REMEMBER:

1. Medications MUST always be taken as prescribed. Do NOT run out of supplies.

2. Hydrocortisone is VITAL in helping the body cope with stress. At times of illness, the prescribed dose will need to be INCREASED and if vomiting occurs, the hydrocortisone will need to be given by injection.

3. Children who are deficient in pituitary hormones and/or who suffer with fits, for whatever reason, should wear an identity necklet or bracelet and carry a medical card.

More information about some of these conditions is available in the Merck Serono series of educational booklets titled *Hormones and Me* listed on page 48.
Possible Longer–Term Problems

When a craniopharyngioma is first diagnosed it may be quite large and might press against the hypothalamus and other brain structures. This can result in changes in the child’s development and behaviour including altered growth, eating and sleeping patterns.

Neurosurgeons are increasingly aware of the importance of causing as little damage as possible when they remove a tumour of this type – even if it means leaving a small amount behind. Some damage, however, may still occur during surgery and this can also lead to long–term problems. These problems may sometimes improve over time.

Long Term Monitoring of Growth

A child’s growth and development can be affected by a craniopharyngioma and height and weight must be measured regularly. Pubertal development will also have to be carefully monitored.

“It is common for children to gain some weight after craniopharyngioma surgery, but the amount varies from child to child.”

Children can sometimes grow normally following surgery even though the pituitary gland is no longer producing growth hormone. Normal growth may continue for many months or years, but can then slow down. Growth hormone injections may need to be started at this point to re–establish normal growth. In other children growth hormone deficiency can cause poor growth both before and after surgery, and they will need hormone treatment earlier.

Long term growth monitoring is usually carried out at a growth clinic by a paediatrician or paediatric endocrinologist.
Increased Food Intake and Excess Weight Gain
Excessive weight gain may result from damage to the hypothalamus and its hormones. The ‘eating control’ centre is disrupted causing a persistent feeling of hunger, so that the child feels hungry even immediately after a meal. This change in eating behaviour can eventually cause children to become overweight or obese if not monitored carefully.

Increased appetite can develop very soon after surgery and the child can become totally preoccupied with thoughts of food. It is important to discuss the issue openly and to encourage the involvement of the child as much as possible in the ideas of how to deal with this problem. Early referral to a dietician may be helpful to guide you through healthy eating options for the child.

Understandably, this change in behaviour may cause distress to the child and their family, particularly if there is a sudden change in body appearance due to rapid weight gain. Problems at school may also occur both from poor concentration and from teasing by other children. Talking to a psychologist can be helpful.

Other causes of excess weight gain include:
• a change in the metabolic rate of the body (possibly due to a change in levels of the hormone leptin)
• high doses of steroids given around the time of the operation
• other pituitary hormone deficiencies (such as growth hormone deficiency or thyroid stimulating hormone deficiency).
**Sleep Disturbance**
Sleep disturbance can be caused by a large tumour pressing on other areas of the brain or via damage during surgery. A child may wake up several times during the night or fall asleep at odd times during the day. The problem may continue as the child gets older.

The main concerns associated with these altered sleeping patterns is the impact on the child’s education (they may be very tired when they are at school) and the disturbance to other members of the family, particularly if the night-time wakefulness is spent looking for food! Daytime sleepiness can be treated effectively in some children with an amphetamine derivative.

**Impaired Thirst**
Although rare, the inability to feel thirsty is a serious problem that can sometimes occur when a child’s hypothalamus is damaged. It is particularly serious when it is associated with diabetes insipidus. If both conditions are present, carers need to use a ‘fluid prescription’ with accurate daily measurements of fluid intake and variations in volume depending on the weather.

**Memory Disturbance**
Memory may be affected either by damage to the hypothalamus or disturbance to other nearby areas of the brain. The type of memory loss will vary from child to child, but short-term memory is usually affected rather than long-term memory.
Memory ‘processing’ can become a problem for some children. This is where new information is stored briefly but cannot be transferred into the long-term memory. This may cause problems at school, especially high school, where learning in some subjects, such as mathematics, builds on previously learned information. This problem needs to be recognised early, so the child or teenager can be helped or they can choose appropriate subjects where this type of learning is not so important. Amphetamine derivatives can be helpful at reducing the impact of this problem.

**Impaired Temperature Regulation**

This is a rare but serious problem. Family and friends may notice that the child is not sensitive to temperature extremes. When everyone else is piling on jumpers, the child might be walking around in T-shirts or on a hot day ask for a jumper!

Very occasionally a child may have difficulty in maintaining a normal body temperature and need an electric blanket or hot water bed to prevent hypothermia. The problem requires very careful management, particularly in hot weather, as the body temperature may rise uncontrollably, causing a seizure to occur. Sometimes it may be necessary to keep the child in an area with reasonably constant temperature when the outside weather is extreme.

Impaired temperature regulation is often associated with other severe hypothalamic problems, such as impaired thirst. This can be a very difficult situation to manage as carers need to vigilantly monitor both the child’s fluid needs and environmental temperature.
Mood Swings
Children treated for a craniopharyngioma occasionally have a much ‘shorter fuse’ than before the operation and have temper tantrums which are quite out of character. This may be due to the fact that they have been ill or have damage to the hypothalamus. In some cases it may be necessary to seek advice from a psychiatrist.

Metabolic Syndrome and Cardiovascular Problems Later in Life
Metabolic syndrome is a collection of disorders, including weight gain, high insulin levels and high cholesterol, that increase the chance of developing diabetes, high blood pressure, heart disease and stroke in later life. Craniopharyngioma patients can have many of the risks factors for metabolic syndrome. It is therefore very important to monitor these risk factors and treat these conditions if they arise.
Additional Long-Term Follow-Up

Eyes
Eye sight problems often improve after surgery and may even return to normal. However, sometimes visual loss can become permanent. Eye tests will be needed at regular intervals to monitor progress. Special help may also need to be organised at school if the child's sight is very restricted in one or both eyes.

Brain Scans
A brain scan will usually be performed within 24–48 hours of surgery or before the child is discharged from hospital to make sure all is well. A repeat scan will probably be arranged 6 months after the operation to check for any regrowth. Depending on the results of this, scans will be repeated every 6–12 months for the next 2–3 years and then as recommended by the specialist.

School Progress
Some children may experience some learning difficulties following treatment of a craniopharyngioma. It is very important to monitor the child's progress at school and to provide extra assistance if a problem is recognised.

Problems may include:
• difficulty in catching up on missed school work
• difficulties keeping up due to poor sight
• difficulties with concentration because the child is distracted by thoughts of food
• difficulties due to learning or memory problems.
Ideally, a psychologist should be involved early on in the child's care so learning and memory testing can be done at intervals after the operation or any radiotherapy treatment. Any special needs can then be picked up early and prompt referral can be made to an educational psychologist so that appropriate support and supervision can be organised at school as soon as possible. Rarely the child’s needs may be better met in schools geared to special educational aspects and it is crucial not to let children struggle for several years before finding this out.
Questions and Answers

Will the craniopharyngioma regrow?
There is a possibility that the craniopharyngioma will regrow. This is why it is important to have regular brain scans after the initial treatment. If it was not possible to remove the entire tumour then the chances of regrowth are increased.

What are the signs that a craniopharyngioma is regrowing?
Sometimes a child will show no signs that the tumour has regrown and it will only be picked–up during a routine follow–up scan. At other times the regrowth may trigger the return of previous symptoms. One early sign that the tumour may be regrowing is poor growth in children who are on hormone replacement therapy.

What happens if the tumour regrows?
Treatment options will depend on the individual child. Radiotherapy is sometimes used. Further surgery may also be necessary.

Will a child's brothers and sisters have the condition as well?
This is unlikely. Craniopharyngiomas are not hereditary (not passed on from one generation to the next). They happen by chance as a result of a problem that occurs in the developing unborn child.

What will happen when a child treated for a craniopharyngioma becomes an adult?
Most of the hormone treatments a child receives will need to be continued into adulthood. Some hormone deficiencies may also develop later and it is very important that the child is referred on to an adult endocrinologist in their mid to late teens for continued ongoing care.
Will the child be able to have children when they grow up?
Yes. Some will need hormone replacement therapy (testosterone or oestrogens) in order to go through puberty normally and will need to stay on this treatment to maintain sexual development.

For those who need HRT on a long term basis, special replacement treatment will be needed to induce fertility, when it is desired. This is quite quick and simple for women but takes longer for men to achieve normal sperm numbers. For both men and women the treatment involves daily injections for several months. This is one of the reasons why they will need to remain under the care of an endocrinologist even as an adult.

Why is this tumour different in adults?
Craniopharyngiomas that occur in adults tend to have a different structure compared to those found in children. Childhood craniopharyngiomas can be more difficult to treat because they are growing faster – especially the ones that cause symptoms in children at a very young age.
Glossary

Cerebrospinal Fluid (CSF)
The fluid, which flows through the ventricles and over the surface of the brain, eventually draining into the blood–stream.

Chemotherapy
Medications used to treat various forms of cancers or tumours.

Computerised Tomography (CT Scan)
A type of X–ray which shows a three dimensional view of the area being examined.

Congenital
A feature or condition that is present from birth, but not necessarily hereditary.

Convulsion
Involuntary muscular contractions and relaxations, also known as a fit.

Cortisol
A steroid hormone produced by the adrenal cortex. There are a number of hormones made by the adrenal gland and they are called corticosteroids.

Dehydrated
Condition resulting from excessive fluid loss by the body, i.e. when fluid intake fails to replace fluid lost by the body.

Dietician/Nutritionist
A health professional who specialises in dietary advice.
**Endocrine Gland**
A gland that makes hormones and releases them into the blood. The pituitary, thyroid, adrenal, testes (testicles) and ovaries are all endocrine glands. All of the glands together make up what is termed the endocrine system.

**Endocrinologist**
A doctor who specialises in the disorders of the endocrine glands.

**Gonadotrophins**
The sex hormones, i.e. follicle stimulating hormone (FSH) and luteinising hormone (LH).

**Gonads**
A term that refers to the sex glands, the ovaries in females and the testes in males.

**Growth Hormone (GH)**
A hormone released by the pituitary gland, which promotes growth.

**Hereditary**
When a genetic characteristic is passed on from parents to their children.

**Hormones**
Blood chemicals that stimulate growth and sexual development and help to regulate the body’s metabolism. Hormones are produced by endocrine glands and carry messages from one cell to another via the bloodstream. Normally the body carefully controls the release of hormones as too much or too little may disrupt the body’s delicate balance.
**Hyperthermia**
A term used to describe an unusually high fever, where body temperature is raised above normal.

**Hypothalamus**
Part of the base of the brain that controls the release of hormones from the pituitary gland.

**Magnetic Resonance Imaging (MRI Scan)**
A technique for obtaining high–resolution scans of the brain and other parts of the body. There is no exposure to X–rays and they can therefore be repeated in the same person many times.

**Nerves**
Transmit impulses from the brain to the rest of the body to make the body respond in a certain way.

**Neurosurgeon**
A doctor who specialises in surgery on the brain.

**Oestrogen**
A group of female hormones that are produced by the ovaries (from the onset of puberty until menopause) and which control female sexual development.

**Optic Chiasm**
The area just above the pituitary gland where the optic nerves from both eyes meet and cross one another.
Paediatric Endocrinologist
A doctor who specialises in the disorders of endocrine glands in children.

Pituitary gland
A pea–sized gland at the base of the brain, which releases a number of important hormones related to normal growth, development and fertility.

Puberty
Essentially the period in a young person’s life, both male and female, where they become physically capable of reproduction.

Subcutaneous Injection
An injection given beneath the skin.

Testosterone
Most potent male sex hormone, which is produced in the testes (testicles) and controls male sexual development.

Thyroid Gland
A butterfly–shaped gland in the front of the neck below the larynx, which makes the hormone thyroxine.

Tinnitus
Constant ringing in the ears.
**Urinary Catheter**
A tube inserted into the bladder to remove urine from the body. It is used when the bladder is unable to function properly, or for a hospitalised patient who cannot leave their bed.

**Womb**
Uterus.
Support Organisations and Further Reading

Australian Pituitary Foundation Ltd
www.pituitary.asn.au

Australasian Paediatric Endocrine Group (APEG)
www.apeg.org.au

The Endocrine Society
www.endo-society.org

John Hopkins University (Information on Radiosurgery)
www.hopkinsmedicine.org/neurology_neurosurgery/specialty_areas/brain_tumor/treatment/radiation-therapy.html

The Magic Foundation
www.magicfoundation.org

Pituitary Foundation UK
www.pituitary.org.uk

Pituitary Network Association (USA)
www.pituitary.org

UK Child Growth Foundation
www.childgrowthfoundation.org

UK Society for Endocrinology
www.endocrinology.org
References for Text


Merck is proud to bring you this booklet from the *Hormones and Me* educational series. We aim to provide readers with a better understanding of the issues relating to endocrine disorders particularly in children. We hope that you find it a valuable and helpful resource.

Please ask your doctor or nurse for further information on the resources available to you.

The *Hormones and Me* series includes:
1. Growth Problems in Children
2. Turner Syndrome
3. Craniopharyngioma
4. Diabetes Insipidus
5. Puberty and its Problems
6. Delayed Puberty
7. Multiple Pituitary Hormone Deficiency (MPHD)
8. Congenital Adrenal Hyperplasia (CAH)
9. Growth Hormone Deficiency in Adults
10. Management of Emergency or ‘Stress’ Situations where Hypoglycaemia or Cortisol Deficiency Occur
11. Intrauterine Growth Retardation (IUGR)
12. Congenital Hypothyroidism
13. Klinefelter Syndrome
14. Disorders of the Thyroid Gland in Children and Adolescents

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This booklet is valuable reading for anyone with Craniopharyngioma.

It is also recommended reading for their family and friends.