Prader-Willi syndrome (PWS) is a rare lifelong genetic condition which is present from birth. Caused by an abnormality on chromosome 15, it affects the functioning of the hypothalamus and other aspects of the brain and muscles. Main characteristics of the syndrome are:

- Hyperphagia – beginning around 2-4 years
- Hypotonia – from birth
- Hypogonadism – from birth
- Short stature, compared with other family members (less evident if the person is receiving growth hormone treatment)
- Delay in and difficulties with emotional and social development
- Developmental delay /learning disabilities ranging from severe to borderline. A minority of people with PWS do not have a general learning disability. Most have specific difficulties such slower processing of verbal information etc
- Challenging behaviours, (PWS behavioural phenotype), regardless of whether or not the person has a learning disability.
- A significant number of individuals with PWS also have Autistic Spectrum Disorder.

**UNUSUAL FEATURES AND POTENTIAL RISK FACTORS IN PWS**

A high pain threshold, poor temperature regulation and lack of vomiting associated with hypothalamic dysfunction can mean that a serious medical problem may present in an unusual way and so could be overlooked.

**Abnormal body composition affecting medication**

Individuals with PWS have reduced lean tissue and increased adiposity. This may affect dosages of medication, which may need to be less than usual. Psychotropic medications must always be started at very low doses. Medications that have antidiuretic effects may cause water intoxication.

**Body temperature abnormalities**

An individual with PWS may be unaware when they are too hot or too cold, or eat or drink very hot food or liquids. Hyper and hypothermia have been reported. Hyperthermia may occur during minor illness and after anaesthesia. Fever may be absent despite serious infection. In cases of hypothermia, the patient may not report feeling cold. Baseline temperature may be below 98.6°F.

**High pain threshold**

The majority of people with PWS have a high pain threshold and hence underreport pain. This may mask fractures, acute abdominal conditions, serious infections and internal injuries. Individuals may have difficulties localising pain.

**Lack of vomiting**

Vomiting rarely occurs in PWS. The presence of vomiting, particularly when the individual has a history of never or very rarely vomiting, may signal a life-threatening illness (see below). Emetics may be ineffective.
Severe gastric distension with ischaemia
This is a life-threatening situation that may also result in stomach rupture. The person with PWS may present with abdominal distension, pain and/or vomiting but may only complain of mild abdominal discomfort. This gastric inflammation with necrosis has often been seen in individuals with PWS who have had a recent binge episode and whose weight is generally under control.

Gastroparesis
Many individuals with PWS also have gastroparesis, which can become dangerous if overeating occurs or has occurred. This is more common in adults, but may affect some children.

Loss of appetite
People with PWS will usually eat even when ill, due to hyperphagia. Loss of appetite should be viewed as possibly symptomatic of a serious disorder. However, some people with PWS will refuse food as a controlling behaviour.

Skin lesions – picking and bruises
Many individuals with PWS pick at small wounds or spots, intensifying and increasing the wound, and thus the potential for infection. This can complicate the healing of IV sites and incisional wounds. It is common for individuals with PWS to have sores caused by their own skin-picking. They may bruise easily. Appearances may wrongly lead to suspicion of abuse.

Tendency to psychosis
There have been a few anecdotal reports of teenagers and adults with PWS experiencing psychotic episodes during hospital stays. If parents and carers report worsening differences in behaviour, seek psychiatric help.

GENERAL CARE WHILE AN INDIVIDUAL WITH PWS IS IN HOSPITAL

Informed consent (adolescents and adults)
Anxiety is common in both adults and adolescents with PWS and individuals benefit from receiving clear information about what is going to happen. Even those with higher functioning levels may become anxious or confused about proposed treatments (e.g. insertion of IV lines). This might trigger challenging behaviour. They will require a calm, patient and understanding approach and support from a learning disability liaison nurse if possible.

Lower calorie diet
Severe obesity can be caused by the combination of hyperphagia and hypotonia. However, in individuals whose access to food has been well controlled, obesity may not be evident. In all cases, however, energy intake should be around two thirds of the average for the individual’s age group, whilst maintaining a balanced, nutritious diet. The input of a dietitian may be required.

Restrict access to food
Hyperphagia causes food-seeking and food-stealing behaviour. Ensure that the individual with PWS is not left alone with food or drinks trolley or drinks vending machines, nor has access to other patients’ food and plates. Be aware that an individual may say that they have not had a meal or break when they have already eaten.
Comprehension and language – receptive and expressive ability

Many people with PWS can give the impression that they understand everything said to them. This is not always the case. Others may have articulation problems or dyspraxia and hence be difficult to understand. Children and adults may interpret information literally. Keep instructions clear and simple. Use visual aids to help with both comprehension and communication. Give them plenty of time to process information and respond.

Time frames
Be very clear about time frames. If you say you will be back in a few minutes, make sure you do so. Not sticking to times given can result in a rapid increase in anxiety and escalating challenging behaviour.

Parental support
Individuals with PWS require structure to their lives and can find unexpected events or change difficult. The hospital environment can be a very challenging time for them. Parental or carer support and access may be needed much more frequently than usual, especially for adults in wards with restricted visiting times.

For children, take parental advice about their child’s routines and expectations particularly about food. Small deviations in expectations around food can have significant consequences. Take parents’ lead on ways of reducing anxiety and responding to any behavioural difficulties.

Behavioural and emotional outbursts
Children and adults with PWS are prone to emotional lability and outbursts. The reasons for an outburst are not always clear but may include hunger, tiredness, uncertainty, unexpected events, conflict etc. Parent or carer guidance can help to minimise these.

TREATMENT

Anaesthesia recommendations for patients suffering from Prader-Willi syndrome
www.orpha.net/data/patho/Pro/en/Prader_Willi_EN.pdf

Cortisol levels
Individuals with PWS are probably more likely that the rest of the population to have cortisol deficiency. If there is clinical suspicion of cortisol deficiency during acute illness, immediately take a blood sample to check serum cortisol level. This can then be used to help with diagnosis even if you need to start hydrocortisone as an emergency. Please discuss results with your local endocrinology team.

Medication
People with PWS may have abnormal reactions to standard doses of medication (see Body Composition above)

Patients receiving growth hormone replacement
Growth hormone (GH) replacement is often used in patients with PWS. If felt necessary, it can be temporarily stopped during an acute illness without any likely acute adverse consequences. Patients and families may however wish to continue GH to help maintain routine. Please discuss with your local endocrinology team if any questions or concerns.

Oxygen levels
Oxygen levels may be normally lower than usual. Obese patients who have been chronically hypoxic may not tolerate fully corrective use of oxygen and are likely to start retaining CO2.
ADDITIONAL INFORMATION

Guidelines for postoperative monitoring of patients with Prader-Willi Syndrome

Guidelines for postoperative monitoring of paediatric patients with Prader-Willi Syndrome

Obesity in PWS

Clinical Presentation of Obesity Hypoventilation and Right Heart Failure in Prader-Willi Syndrome

Psychiatric alert for Prader-Willi syndrome
www.pittsburghpartnership.com/handouts/Psychiatric%20Alert%202009.pdf

Medical alerts in several languages are also available at http://www.ipwso.org/#1medical-alerts-booklet/czxp

WEBSITES WITH EXTENSIVE INFORMATION ABOUT PWS

PWSA UK www.pwsa.co.uk
International PWS Organisation www.ipwso.org – includes Medical Alerts in several different languages
PWSA USA www.pwsausa.org

PWSA UK

PWSA UK is in contact with medical specialists with an expertise in PWS. Please do not hesitate to contact us for further information at the address below.

ACKNOWLEDGEMENTS

With grateful thanks to the following for their help with compiling this information:

- Dr Nicola Bridges, Consultant Paediatric Endocrinologist, Chelsea and Westminster Hospital
- Dr Anne Livesey, Consultant Community Paediatrician, Sussex Community Trust. (Brighton PWS Multidisciplinary Clinic)
- Dr Tony Goldstone, Consultant Endocrinologist, Imperial College Healthcare NHS Trust
- Manjeet Singh Riyat, Consultant in Emergency Medicine, Royal Derby Hospital