Prader-Willi Syndrome in Bahrain: Case Presentation

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A case of Prader-Willi syndrome (PWS) is described in a 6 year old Bahraini boy. He presented in the neonatal period with hypotonia and feeding problems. At 3 years of age he developed voracious appetite which led to obesity and psychiatric manifestations. Clinically he had dysmorphic features consistent with diagnosis of PWS. The diagnosis was confirmed by karyotyping and DNA analysis. Making early diagnosis of PWS is vital for: a) management of various manifestations, b) parental counselling and c) avoiding unnecessary investigations for simulating conditions.

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PWS is a syndrome characterized by protean manifestations presenting early in life with new features as age advances. It is also known as Prader-Labhart-Willi syndrome when described first in 1956. The early fetal and neonatal signs are nonspecific, this include hypotonia, poor feeding and genital hypoplasia, changing later on to polyphagia, obesity, small hands and behavioral and psychiatric manifestations. The prevalence of PWS is one in 10,000 births, this figure now is increasing due to improved molecular genetic laboratory studies. The diagnosis is confirmed by karyotyping and DNA analysis. In about 70% of the cases, it is caused by loss of a critical portion of the proximal part of the long arm of paternally derived chromosome 15 (del 15 (11q13)1-3. The remaining 30% of cases are of nondeletion, caused by inheritance of two maternal copies (two maternal copies of 15q and absent paternal copy). We present our experience with a diagnosed patient suffering from PWS, followed by discussion.

THE CASE

A Bahraini male newborn delivered by an assisted breech after 36 weeks gestation to a 24 year old primigravida mother. Apgar scores were normal. Parents are not consanguineous. There was no history of neurological disorder on either side of the parents. Pregnancy was uncomplicated and according to the mother fetal movements were normal. At birth he appeared to be of normal morphology, apart from small hands (Fig 1). His growth parameters were as follows: Weight 2230 gm, length 47cm and head circumference 32.5 cm, corresponding to the 5th, 50th and 40th percentiles respectively. Physical examination showed normal morphology, except of small hands. He had profound hypotonia with diminished primitive reflexes. Testes were undescended (felt above inguinal ring), penis of normal size (20.5mm). Because of respiratory distress and prematurity, he underwent septic work-up and treated with broad-spectrum antibiotics, intravenous glucose solution and 30% oxygen via head box. Five days later, because of poor sucking reflex, he was fed by nasogastric tube for two weeks. He started to suck the bottle and the breast very slowly. His blood culture was sterile. Blood glucose, urea, electrolyte and calcium were normal. Brain ultrasound was normal. Thyroid function test showed slightly elevated TSH (T3 and T4 were normal), for which he was treated with thyroxine for 3 months until a repeat test was reported normal.

Figure 1: The hands of the patient showing small hands

Over the following months and first two years he showed marked developmental delay. Social smile was first noted at 6 months, attained head control at 10 months, rolled over at 14 months, sat unaided at 24 months and spoke meaningful polysyllables at 28 months. During the first year of life, he had feeding problem. His feeding time lasted more than half an hour, mostly by bottle, but also spoon feeding was used to expedite his feeding time. At 3 years of age he developed voracious appetite for milk and sugar containing food. He insisted on eating without restrictions, attempt to do so led frequently to tantrum and raging episodes. His morphological changes changed to generalised obesity, with normal stature.

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his head was dolichocephalic. The palpebral fissure was almond-shaped. The corners of the mouth were downward-turned (Fig 2). The hands were moderately small. The hair colour was fair; skin and eyes color were normal. The parents colour of the hair is black, the skin is light brown and the eyes were black. At 4 years and 3 months he weighed 31.1 kg (> 97th percentiles) and height was 108 cm (< 90th percentiles). A dietitian was consulted to control his obesity. Compliance with dietary restriction was poor. The obesity was investigated by brain and adrenal gland CT, fundic examination and ACTH stimulation test, all of which were normal and ruling out any hypothalamic-pituitary-adrenal pathology. He underwent orchidopexy. Chromosomal study and DNA analysis, performed overseas (JSPL Lab, Bewlay House, London, UK). It showed a normal boy with 46,xy karyotype. DNA analysis showed only maternal allele at 15q11-13, typing the patient and his parents with polymorphic markers on 15q confirmed that he had maternal deletion with GABRB3 and L-S1 (D15S113).

![Profile of the patient showing facial features.](image)

At 5 years of age, he remained polyphagic and moderately obese. He developed dental caries. Intelligence Quotient (IQ) was 60 on Griffith's Scale. His behavior was labile and frequently led to temper and aggressive episodes especially if prevented from eating certain food. At the age of 6 years he slept frequently during the day with inspiratory stridorous breathing (Pickwickian syndrome). During hospital stay to assess his problem, it was noted that he became hypoxaemic and hypercarbic. He underwent adenoidotonsillecctomy to relieve his upper airway obstruction with little improvement. The family is trying to admit him in a special school for mentally handicapped children.

**DISCUSSION**

According to clinical features our case fulfill the criteria of Holmes et al. In children over three the diagnosis is satisfied if there are 5 criteria, 5 of which must be major ones. Our patient presented with abnormal craniofacial and hands morphology, initial hypotonia, difficulty in feeding, later on developed polyphagia and obesity. The diagnosis confirmed by karyotyping and DNA studies, which showed the usual (70%) abnormality of paternally deleted chromosome 15q. The etiology in the non-deleted cases (30%), the maternal disomy (with 2 maternal and no paternal allele) is presumably due to nondisjunction and duplication occurring early at the gametale stage. The mode of inheritance is sporadic in majority of cases. However, some authorities claim that paternal exposure to hydrocarbons in the causation leading to deletion in PWS. The delayed diagnosis in our case is not unusual. This is due to changing early signs and symptoms as children get older. The presence of upper airway obstruction and hyperventilation and obesity presented in our patient has been reported by others. A rare presentation, which did not present in our patient, is severe skin picking behavior, this can be complicated by self-mutilation and extending this to the rectum resulting in lower gastrointestinal bleeding. Our patient had dental caries reported by Dunn et al. He also had fair skin color compared to his family member. The presence of a common condition like nonspecific mental retardation, of about 3%, should raise high index of suspicion for other condition simulating PWS.

The purpose of this presentation is to alert the clinicians not familiar with PWS in order to diagnose them early in life. This would be beneficial for: a) provision of multidisciplinary patient care, b) avoid unnecessary investigations (for other similar conditions) and c) provide a family counseling. It is noteworthy that this condition can be diagnosed antenatally using amniotic fluid. To our knowledge this is the first case of PWS to be reported in Bahrain.

**REFERENCES**


