Case Report

Joubert syndrome in a neonate: case report with literature review

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ABSTARCT

Joubert syndrome is a rare autosomal recessive disorder. It is characterized by congenital ataxia, hypotonia, developmental delay and at least one of the following features: neonatal respiratory disturbances and abnormal eye movements; including nystagmus and oculomotor apraxia. Molar tooth appearance is an essential finding for the diagnosis of Joubert syndrome. We report a five-days-old newborn with mild hypotonia, abnormal pattern of respiration, abnormal eye movements and molar tooth sign on brain CT scan. Joubert syndrome is an uncommon inherited condition and delayed diagnosis is usually related to its variable, non-specific presentation. Awareness of the characteristic clinical and radiological findings in Joubert syndrome will help in early diagnosis, appropriate counseling and proper rehabilitation.

Keywords:
Joubert syndrome; Hyperpnoea; Molar tooth sign; Neonate; Oculomotor apraxia, Yemen.

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INTRODUCTION
Joubert syndrome (JS) is a rare genetic heterogeneously inherited disorder, named after Marie Joubert in 1969 [1-3]. It is characterized by brain stem and cerebellar malformations and multi-systemic affection involving nervous, ocular, gastrointestinal and urogenital system in varying proportions [4,5]. Molar tooth sign (MTS) and “Bat-wing” appearance on brain CT are basic findings in JS [6,7]. We report a newborn with mild hypotonia, abnormal respiration (episodic tachypnea and apnea), oculomotor abnormalities and MTS on brain CT scan. To our knowledge this is the first case of JS reported in Yemen.

CASE REPORT
A five-days-old term, 3.4 kg, female baby was admitted to our hospital with frequent attacks, every 5-15 min, of apnea; each lasting 20-60 seconds, associated with bradycardia and alternating with hyperpnoea (up to 120 breaths/min). No history of fever, cyanosis, decreased feeding or activity, or history of trauma. There was no reported consanguinity or similar cases in the family. On examination, she was active, alert, crying normally, has low set ears, head circumference 35 cm (at 50th centile) and heart rate was 140 b/min. Systemic examination was normal apart from faint systolic murmur on cardiac auscultation. Laboratory investigations, including hemogram, blood sugar, electrolytes, C-reactive protein, liver enzymes, urea and creatinine were all normal. Blood culture was not available.

Three days later, left side upper limb seizures appeared in association with tongue protrusion and cessation of respiration but no bradycardia. An abnormal slow eye movement upon turning the head to the sides was noticed too. Chest X ray, brain and abdominal ultrasound were normal, echocardiography showed mild Epstein’s anomaly with mild pulmonary hypertension. Funduscopic examination was normal. Brain CT scan showed MTS (Figure 1). Genetic analysis was not available.

After three weeks on supportive treatment, the frequency and the duration of the apneic attacks improved, as she only had 1-2 attacks/day, each lasting < 5 seconds. The patient was discharged home on O2 therapy, as per need. On a follow up visit, at the age of three months, she had regular breathing, mild hypotonia with hand fisting and hyperreflexia. She did not follow objects on eye examination. Her head circumference was 36 cm (below the 3rd centile). Unfortunately, since that time, we have lost contact with the patient.
DISCUSSION

Marie Joubert and associates first described JS in 1969 in four siblings and one sporadic case that exhibited episodic rapid breathing, abnormal eye movements, ataxia and mental retardation with agenesis of the cerebellar vermis [2]. By 2009, approximately 200 cases have been reported worldwide [8]. Most cases are sporadic, but some families may show a recessive pattern of inheritance [9]. The clinical manifestations usually start in the neonatal period, but the correct diagnosis is often delayed months or years after birth because of its variable phenotypes [10]. The term ‘JS and related disorders’ (JSRD) refers to a group of disorders presenting the pathognomonic features of JS in association with multiple systemic congenital abnormalities. It has an estimated incidence of 1/80000 -1/100000 live birth [1,11]. These disorders have been classified as ciliopathies [12]; and were attributed to 23 genes [13]. Mutations in the AHI1 and CEP290 genes are causal in 10-15% and 10% of cases, respectively. Homozygous deletion of NPHP1 gene results in 1-2% of cases [14].

Joubert syndrome is characterized by hypotonia, congenital ataxia, developmental delay and at least one of the following supportive features: neonatal respiratory abnormalities and abnormal ocular movements; including nystagmus and oculomotor apraxia [4,15]. Valente, et al, reported six clinical subgroups of JS [9], which were recently replaced with a more practical clinical – genetic classification [11,16,17]. Seizures and cardiac anomalies were rarely reported in JS [6,11].

Typical dysmorphic features (hypertelorism, small ear lobes, broad forehead, arched eyebrows, broad mouth with intermittent tongue protrusion, ptosis and anteverted nostrils) are present and hypotonia is important only if associated with other features [4,18]. Respiratory manifestations are characterized by alternating episodic attacks of hyperpnoea and central apnea or episodic hyperpnoea alone, which intensifies when the patient is stimulated. It starts typically in the neonatal period, gradually diminishes with age and disappears around the 6th month of life [2,11,19]. Ergun et al [20] reported persistence of the respiratory symptoms till the age of 17 years. The severity of respiratory irregularities can range from mild to severe and prolonged attacks of apnea [11,19]. Abnormal eye movements are characterized by nystagmus and oculomotor apraxia [15]. The similarity in clinical and radiological aspects between JS and both hypotonic cerebral palsy and rhombencephalosynapsis can lead to delayed or erroneous diagnosis [21,22]. Characteristic radiological findings in CT and MRI can help to guide and establish the diagnosis of JS [7]. The molar tooth sign is an essential part in the diagnosis of JS and JSRD [11]. It results from a combination of midline hypo-dysplasia of the cerebellar vermis, deepened interpeduncular fossa (at the level of isthmus and upper pons) and horizontal, thick, elongated superior cerebellar peduncles [1,4,12,15,23]. Batwing or umbrella sign results from the dilated or distorted fourth ventricle [2,7,12]. “Buttock sign” appears in the coronal images, resulting from the absence of the posterior vermian lobe, leaving the normal cerebellar hemispheres separated by a midline cleft [7]. The cerebrum is not affected but moderate lateral ventricular enlargement was noticed in 6%–20% of cases, with corpus callosum dysgenesis in 6%–10% [24]. Features resembling Dandy-Walker malformation were observed in 10% of cases with JS [5]. Recently, “Shepherd’s Crook sign” was reported. It appears in the sagittal views of the posterior fossa where the brain stem and the pons form the shaft of the crook and the abnormal superior cerebellar peduncles and cerebellar hemisphere complete the arc of the crook [25]. Multidisciplinary approach with supportive and symptomatic treatment is needed depending on the presence of neurological deficits and cognitive
impairment, considering respiratory and feeding problems a priority [11,12,23,26]. Apnea monitoring is required for possible intensive care management and assisted ventilation, especially; peri-operatively as JS patients are very sensitive to the respiratory depressant effects of anesthetic agents [19,26,27]. In general, the prognosis of JS is poor [8]. It largely depends on the severity of different organ systems involvement [12]. Steinlin et al [28] described a variable course for developmental outcome in JS: some patient’s die early in infancy, some have severe developmental handicaps and others survive with mild developmental delay. The 5-year survival rate is 50% [8]. Death is usually due to feeding difficulties and respiratory infections [7]. Genetic counseling is an important part of management. The recurrence risk for JS is 25%. Once a diagnosis is made in a neonate, serial antenatal ultrasonography screening should be performed in subsequent pregnancies combined with fetal MRI at 20–22 weeks gestation to maximize the accuracy of prenatal diagnosis [18,27,29,30]. Annual screening for progressive JSRD complications is recommended [12,27].

CONCLUSION
Joubert syndrome is an uncommon inherited condition. Delayed diagnosis is usually related to its variable, nonspecific presentation. Awareness of the characteristic clinical and radiological findings in JS will help in early diagnosis, appropriate counseling and proper rehabilitation.

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REFERENCES


