CASE REPORT

Nystagmus in a newborn: a manifestation of Joubert syndrome in the neonatal period

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SUMMARY
Joubert syndrome is a rare disorder, usually autosomal recessive, with a prevalence of 1:80 000 to 1:100 000. This disease presents most commonly as breathing irregularities, although the two major clinical criteria are hypotonia and developmental delay, sometimes associated with ocular movement abnormalities. The severity of the presentation varies, ranging from mild cases with normal intelligence to severe developmental delays associated with early death. We report a case of a newborn who presented to the emergency department for absent ocular fixation and torsional nystagmus without other neurological abnormalities. Her cranial MR showed cerebellar vermis agenesis and a molar tooth sign. Her laboratory evaluation, and renal and abdominal ultrasound were normal. An electroretinogram showed mixed retinal dystrophy and an AH11 homozygous missense c.1981T>C mutation was identified (parents are carriers). Throughout infancy, she has shown mild developmental delay and hypotonia, but no respiratory abnormalities. Owing to variable expressivity, a high level of suspicion is required.

BACKGROUND
Joubert syndrome is an autosomal recessive disease (in rare cases X linked) characterised by altered development of the cerebellum and brainstem.1 2

The estimated prevalence is between 1:80 000 and 1:100 000 and it is probably underdiagnosed. The average age for diagnosis is 33 months but there are reports of diagnosis as late as 25 years of age, and many patients die before the diagnosis is ever made.2–7 Despite the fact that clinical characteristics are often present in the neonatal period, there is a significant delay in their recognition.8 This disease affects both genders and all races equally.9

While the underlying physiopathology is not clear, it may be related to a failure in the development of rhombomeres, and the V and X cranial nerves from the ectoderm, leading to defective cerebellar development.4 The cerebellum controls balance and motor coordination and the vermis is responsible for posture and rhythmic modulation that originate stereotyped movements such as walking.2–9

The most common presentation is intermittent effortless tachypnoea that may or may not be associated with apnoea, with no bradycardia and no cyanosis.2–8 This disease may also present as hypotonia, ataxia, oculomotor apraxia, seizures, mental retardation, autism and delayed psychomotor development.1 7 12–14 IQs vary widely, ranging from profound cognitive delay to normal intelligence.12

Cranial MR classically shows underdevelopment of the cerebellar vermis, along with the molar tooth sign, which is essential for the diagnosis, although not specific.2 This sign derives from four anomalies: increased depth and length of the interpeduncular fossa with a narrow isthmus, increased thickness of the cerebral superior pedunculae, dilated and anteriorly deviated fourth ventricle with ‘hat wing’ appearance and cerebellar vermis hypoplasia or dysplasia.2–11–15

Other syndromes that are also associated with the molar tooth sign include Debakan-Arima syndrome, COACH syndrome, Senior-Loken syndrome, Varadi-Papp syndrome, Cogan oculomotor apraxia syndrome, Bardet-Biedl syndrome and nephropathosis. These syndromes are included in the same group of diseases designated Joubert syndrome and related diseases (JSRD).3–8 The respiratory pattern, including intermittent tachypnoea and/or apnoea, is characteristic of Joubert syndrome.3–8

Joubert syndrome may be classified into six groups according to organ involvement: pure, with ocular, renal or hepatic involvement, with ocular and renal involvement, and with orofacial or digital defects.16 Alternative classification systems have been suggested, such as dividing cases according to the presence or absence of retinal dystrophy.17–18

Although cardiac screening is not currently recommended in JSRD, there are descriptions of association with aortic stenosis, bicuspid aortic valve and atrial septal defect that may be caused by an overlap with other ciliopathies, such as Bardet-Biedl syndrome.18

The first genetic locus ever associated with Joubert syndrome was in the long arm of chromosome 9. It is currently known that the genes associated with this disease are responsible for proteins expressed in cilia or centrosomes, leading to inclusion in the group of ciliopathies.3–6 Despite all efforts to find specific genetic loci, genetic defects are found in only 50% of cases. Currently, there are 19 known causative genes, including the AH11 (chromosome 6) and CEP290 (chromosome 13).19–20

CASE PRESENTATION
A previously healthy 24-day-old newborn girl presented to the emergency department for absent ocular fixation.

She was born at 41 weeks—by vaginal delivery after an uneventful pregnancy—with a birth weight

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of 3550 kg, the second child of two healthy 24-year-old parents with no history of consanguinity. She was discharged from the nursery after 48 h.

Her family history was remarkable for an uncle with trisomy 21 caused by a de novo mutation. She had no other relevant family or personal history.

She had been observed in another paediatric emergency department 2 days before, where the family was told that her absent fixation could be a part of her normal visual development.

On physical examination, she had conjugated erratic ocular movements suggestive of congenital torsional nystagmus, with a positive blink reflex, associated with tremor of the upper limbs. The ocular media, fundus and optic disk were normal. Her pupils were reactive to light symmetrically. The neurological examination revealed rapid, jerky and irregular conjugated eye movements; the patient was alert and had normal flexor and extensor tone, head response to traction and response to ventral suspension; her tendon and primitive reflexes were normal, as well as her posture, and she had a positive auditory response. Cranial nerve, motor and sensory function testing was normal.

INVESTIGATIONS
Cranial MR revealed cerebellar vermis agenesis with a mesencephalic molar tooth appearance (figures 1 and 2), as well as a thickened cerebral superior pedunculae and a narrow isthmus (figure 3).

Laboratory examinations revealed haemoglobin 13.1 g/dL, 12 000/μL leucocytes, 44.6% neutrophils, 297 000/μL platelets, aspartate aminotransferase 42 U/L, alanine aminotransferase 29 U/L, creatinine 0.4 mg/dL and urea 34 mg/dL. Abdominal and renal ultrasound was normal.

An electroretinogram showed mixed retinal dystrophy comprising attenuated amplitude of cone and rod potentials with preserved macular function.

Gene sequencing was positive for an AH11 missense c.1981T>C homozygous mutation. The patient’s parents are carriers of the same genetic mutation.

OUTCOME AND FOLLOW-UP
The patient is now 24 months old and has developmental delay, as she does not stand or walk. She sits with support, has a social smile, babbles and responds to simple commands. She maintains torsional nystagmus and acquires a tilted head position when trying to fixate objects, although she no longer manifests tremor. There is no report of respiratory irregularities.

She has been on a physical therapy and rehabilitation programme since she was 6 months old, and will maintain follow-up with Neurology, Ophthalmology and Child Development.

DISCUSSION
Criteria for diagnosis of Joubert syndrome include the presence of hypotonia in infancy, developmental delay/mental retardation and at least one of two features including breathing irregularities and abnormal eye movements.17

This case was diagnosed in the neonatal period fulfilling one clinical criterion, abnormal eye movements. The presence of the molar tooth sign in the cranial MR confirmed the diagnosis. The most common manifestation in this age group is respiratory abnormalities, which our patient never developed.2

This case illustrates how important it is to listen to parents’ concerns. Newborns may fixate on a face and have ‘doll’s eye’ movements on turning, but nystagmus and complete absence of ocular fixation are never normal.21

Joubert syndrome is associated with several ocular abnormalities, including nystagmus (horizontal, vertical, torsional, pendular or see-saw pattern) and oculomotor apraxia, which are usually present at birth and may improve with age. Strabismus, ocular coloboma, visual loss, ptosis, pigmentary abnormalities of the fundus and decreased vestibulo-ocular reflexes are also manifestations.22

Retinal dystrophy is also associated with polycystic renal disease,23 for which our patient has tested negative thus far. AH11 gene mutations are associated with retinal abnormalities and renal progressive disease.24 25 AH11 missense c.1981T>C is considered possibly pathogenic according to Mutation Predictor.

Joubert syndrome can evolve variably in three major groups of severity: early death, severe developmental delay with multiple visual and motor deficits, or mild cognitive delay (even normal intelligence).5

Figure 1 Horizontal MRI T2 section of the brain illustrative of the molar tooth sign.

Figure 2 Horizontal MRI T1 section of the brain illustrative of the molar tooth sign.
There are characteristics to this disease that should be considered, namely the progressive nature of retinal and renal dysfunction, which require frequent follow-up and repeat ophthalmological examinations, as well as renal ultrasounds.15 Patients may have adverse reactions to anaesthetics based on opiates and nitrous oxide and these should thus be avoided.26 The early diagnosis in this case allowed us to initiate rehabilitation at an early age. We are hoping that this will potentiate the patient’s psychomotor development and minimise her deficits. An early diagnosis is also important to ensure family genetic counselling, allowing decisions regarding future pregnancies.27

**Learning points**

- Joubert syndrome is a rare disease that is caused by hypoplasia or dysplasia of the cerebellar vermis.
- Breathing irregularities, hypotonia and developmental delay are the most common signs, although early on, patients may not fulfil all the clinical criteria.
- This disease is phenotypically heterogeneous and a high level of suspicion is required to successfully diagnose mild cases.
- Mutations in the AHI1 gene are associated mainly with retinal abnormalities.
- Early diagnosis is essential to ensure timely initiation of rehabilitation measures and genetic counselling.

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**REFERENCES**


**Figure 3** Vertical MRI section of the brain and cerebellum illustrative of thickened superior cerebral superior pedunculae and a narrow isthmus.


