A Girl with Joubert Syndrome: Association of Endocrinological Abnormality and New Radiological Findings

The Editor,

Sir,

An 8.6-year old girl was referred to the Endocrinology Outpatient Clinic due to the onset of puberty. There was consanguinity between parents who had no history of such disease. She had ventriculoperitoneal shunt due to congenital hydrocephaly. The patient was receiving sodium valproic acid since she had a seizure at two years of age. The patient had mental-motor retardation with facial dysmorphism and ataxia, and tent-shaped gait. There was also facial asymmetry, arched eyebrows, nystagmus, ocular hypertelorism, broad forehead, as well as open and tent-shaped mouth [Figure 1]. According to Tanner staging, there was stage 3 breast development. At the presentation, body weight was 34 kg [68th percentile] and height was 138 cm [59th percentile]. Her height-adjusted age and bone age were 9.96 and 10.5 years, respectively. Laboratory findings were as follows: free T4, 0.6 ng/dL; TSH, 1.1 mIU/mL; FSH, 2.19 mIU/L; LH, 6.43 mIU/L; estradiol, 606 pg/mL; ACTH, 30 pg/mL; cortisol, 12 mcg/dL. There was precocious puberty and hormone profile consistent with central hypothyroidism, while kidney and liver function tests were normal. On magnetic resonance imaging (MRI), bilateral kidneys were localized at lower abdominal quadrant, while multiloculated, hypertrophic ovaries were observed at bilateral iliac anterior pararenal areas, and no uterus was seen. On cranial MR imaging, occipital horn of right lateral ventricle was asymmetrically dilated and there was brainstem atrophy [Figure 2a], and asymmetrical dilatation in occipital horn of right lateral ventricle-atrium [Figure 2b]. It was seen that there was a molar tooth sign [Figure 3a], hypoplasia in cerebellar vermis, and bat wing appearance
on fourth ventricle [Figure 3b]. In addition, corpus callosum posterior splenium was thinner [Figure 4a]; there was macroadenoma in the pituitary gland [Figure 4b].

In almost all cases with Joubert Syndrome (JS), there is a molar tooth sign, varying degrees of developmental delay/mental retardation, and history of hypotonia during infantile period. Supportive findings include nystagmus, abnormal ocular movements, and irregular respiration at infantile period. In addition, some authors propose that dysmorphic facial characteristics can be considered within diagnostic criteria (5, 8).

In this case, supportive findings for JS were history of shunt due to congenital hydrocephaly at childhood, hypotonicity, walking delayed until three years of age, inability to speak and nystagmus. Although hydrocephaly has been reported, up to 10%, development of hydrocephaly requiring shunt is extremely rare (9, 10). However, there was shunt in our case. The most common ocular finding is oculomotor apraxia. There was also nystagmus and exophthalmia in our case. Hypothyroidism was previously reported in JS (1). In this case, JS was associated with pituitary macroadenoma, ectopic kidneys and ovaries, precocious puberty, hypothyroidism and uterine agenesis. The findings other (SOMETHING SEEMS MISSING)hypothyroidism has never been reported in association with JS. We presented this case to emphasize that screening patients with JS could help early diagnosis; thus, improve clinical outcomes.

**Keywords:** Ectopic ovaries and kidneys, Joubert syndrome, puberty, uterine agenesis

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Figure 1: Facial asymmetry, arched eyebrow, nystagmus, ocular hypertelorism, broad forehead.

Figure 2: T2-weighted sagittal image; (a) brainstem atrophy on axial image; (b) asymmetrical dilatation in occipital horn of right lateral ventricle-atrium.
Figure 3: T2-weighted axial image; (a, b) molar tooth appearance at superior cerebellar pedicles and bat wing appearance at fourth ventricle (arrow).

Figure 4: T2-weighted sagittal image; (a) dysgenetic appearance of corpus callosum, posterior splenium is thinner than normal (b) macroadenoma at pituitary gland.