Prenatal diagnosis of isolated bilateral anophthalmia

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DESCRIPTION

The patient is a 28-year-old G2P1, with no relevant medical history and a non-consanguineous partner. She presented with an uncomplicated pregnancy, without known history of consumption of teratogenic drugs and a low risk first trimester combined screening for aneuploidies with normal nuchal translucency. At the 20-week fetal morphology scan, in 2D mode, an asymmetry of the eye balls was obvious, with reduced interorbital distance, the right orbit appeared small and hypoplastic and the left orbit was bigger with complete absence of lens bilateral aphakia (figure 1). The rest of the morphological examination was completely normal. TORCH screening completed at the time of diagnosis was negative and the patient had no family history of such congenital anomalies. Amniocentesis was suggested to obtain fetal karyotype but the patient refused. Fetal MRI was corroborative for these findings, also showing normal encephalic anatomy, with normal ventricular system and reassuring the presence of normal corpus callosum and cavum septum pellucidum. The rest of the fetal anatomy evaluated in the MRI was also normal. The couple decided to continue the pregnancy. At 30 weeks, 3D ultrasonography (figure 2) showed sunken eyelid appearance and hypoplastic orbits. She underwent spontaneous labour at term and delivered a 3250 g female new-born, with normal APGAR score. Bilateral anophthalmia was confirmed after birth, with suspected presence of a small cystic malformation located at the left inferior eyelid. Genetic testing after birth, using new generation sequencing panel for microphthalmia/anophthalmia confirmed heterozygotic autosomal recessive mutation for ALDH1A3 gene. The child has revealed normal growth and neurodevelopment so far. Echocardiography performed at 6 months was completely normal. ALDH1A3 encodes a protein (retinaldehyde dehydrogenase) involved in retinoic acid synthesis, playing an important role in eye development.1 Congenital bilateral anophthalmia is a rare condition, affecting 0.6 per 10 000 births, with only about 10% cases appearing isolated. Mutations in numerous genes, including RAX, PAX6, SOX2, OTX2, RARB and ALDH1A3 have been described in association with anophthalmia, being the SOX2 mutations the major single-gene mutation causing about 10%–15% of anophthalmia cases. Anophthalmia can also be a part of trisomy 13 (in about 1% of the cases). However, in about 50%–60% of the cases, no underlying genetic cause is determined.2 Prenatal diagnosis of this condition, both by 2D and 3D ultrasonography, plays a major role in providing continuation vs termination options for parents and allowing psychological preparation and planning for management of this condition, although prognosis for this fetuses is frequently uncertain.3 Although eye examination is part of routine ecanatomy assessment, more and more 3D...
imaging is considered key for early diagnosis, especially when the fetal head position is not favourable.3

Contributors AMV was responsible for clinical research and elaboration of the clinical case. IP was responsible for scientific revision and approval of the final version for publication.

Funding The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.

Competing interests None declared.

Patient consent for publication Obtained.

Provenance and peer review Not commissioned; externally peer reviewed.

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