Very Early In-Utero Diagnosis of Walker-Warburg Phenotype: The Cutting Edge of Technology

Introduction
The Walker-Warburg phenotype belongs to a group of congenital malformations recently defined as malformation due to abnormal pial basement membrane formation. Common characteristics of this group are the association of muscles, brain and eye anomalies. 3 main phenotypes, all with autosomal recessive transmission, have been described within this group: Fukuyama congenital muscular dystrophy (FCMD), muscular eye brain (MEB) and Walker-Warburg phenotype (WWP) or syndrome (WWS) [1]. Among the above, WWP is the most severe and lethal form. Affected individuals are hypotonic at birth and present with severe cerebral and ocular abnormalities.

Results: Very early diagnosis of the Walker-Warburg phenotype at 11 weeks of gestation proved possible by depicting the classic signs of this entity, confirmed by molecular genetics, post-abortion MR imaging and histopathology.

Conclusion: Advancements in ultrasound equipment and technology, molecular genetics and histopathology have made very early detection of this syndrome possible, thus shedding new light on the natural history of this malformation.

Materials and Methods
We describe for the first time a very early manifestation of Walker-Warburg phenotype (WWP) using a 6–12 MHz high-resolution transvaginal ultrasound probe (GE Voluson 8) at 11 weeks of gestation confirmed by molecular genetics, post-abortion T2 MR imaging (1.5T GE) and histopathology.

Key words
- walker-warburg phenotype
- prenatal diagnosis
- advanced technology
- transvaginal ultrasound
- ultrasound
- obstetrics

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Abstract
Background: Walker-Warburg phenotype is a severe and lethal autosomal recessive disorder, belonging to a group of congenital malformations defined as abnormal pial basement membrane formation. So far, prenatal diagnosis was considered possible only during late pregnancy. Methods: First trimester assessment of a pregnancy suspected to be affected by Walker-Warburg phenotype, using a high-resolution transvaginal ultrasound probe (6–12 MHz), T2 MR imaging (1.5T), molecular genetics and histopathology.
A fetus with a CRL of 45 mm was identified using high-resolution transvaginal ultrasound (TVS 6–12 MHz GE Voluson E8). Innovatively, detailed cranial examination depicted various brain and eye features of WWS at a very early stage of pregnancy. These findings included an enlarged 4th ventricle, high insertion of the tentorium, a prominent lateral ventricle, and a malformed midbrain characterized by the typical 2 kinks within the pons. In addition, ocular findings included cataract of the right lens and smaller left orbit (Fig. 1).

Molecular analysis of the DNA obtained by CVS revealed a homozygous mutation in the Fukutin gene (FKTN) c.1167_1168insA (p.F390fs) typical for Ashkenazi origin which confirmed the diagnosis (Fig. 2).

Following dilatation and evacuation (D&E) at 12 weeks of gestation, we were able to collect the fetal head for further evaluation. The biparietal diameter of the head was 13.6 mm and post-abortion magnetic resonance imaging (MRI) was performed and showed an enlarged cisterna magna (CM) with asymmetry of the orbits (Fig. 3). Histopathology of the fetal cortex revealed changes compatible with cobblestone lissencephaly (Fig. 4).

**Discussion**

This is the first report describing fetal ultrasonographic diagnosis of Walker-Warburg phenotype (WWP) at a very early age of gestation with confirmation of the specific causative genetic mutation. WWP is a severe lethal malformation manifested by brain and ocular abnormalities: severe hydrocephaly, cobblestone lissencephaly, cerebellar hypoplasia, and posterior encephalocele. Ocular abnormalities include microphthalmia, cataract, and retinal dysplasia [2, 3].

In the last 2 decades the routine use of fetal imaging studies enabled prenatal visualization of some of the characteristics of WWP, particularly in patients at risk due to previous affected siblings in the family [4, 5].
Hydrocephaly, which is usually the hallmark for the diagnosis of WWS, appears on the prenatal ultrasound mostly in the third trimester [6]. Although ventriculomegaly has been reported early in gestation between 13–15 weeks, this finding was considered non-specific and therefore a more definitive diagnosis could not have been achieved until late in gestation [7, 8].

With the widespread use of fetal MRI, brain imaging studies may reveal a disorganized cortex, shallow gyri (lissencephaly-previous type II), dysplastic cerebellum and characteristic brainstem anomalies.

The typical findings in the brainstem that were described by MRI include pontine hypoplasia, an enlarged quadrigeminal plate with fusion of the superior and inferior colliculi and a distinctive “kink” in the dorsal pons. A second “kink” may be present at the ventral cervicomedullary junction, forming the typical z-shape appearance of the brainstem. However, fetal MRI is applicable mainly late in gestation usually after 28 weeks in the third trimester [9].

Accordingly, ocular anomalies, such as hyperplastic primary vitreous, retinal detachment and dysplasia, which are typical in WWP, have been anecdotally reported at an advance stage of gestation, thus preventing early counseling and management [10–12].

A previous case report in 2005 [6] described a prominent hindbrain vesicle at 12 weeks and dilatation of the lateral ventricle at 14 weeks of gestation, while hydrocephaly and lissencephaly appeared only at 30 weeks. Amniocentesis was performed but without genetic diagnosis since at that time molecular diagnosis was not available [8].

WWP presents with a relatively homogenous phenotype, but it is genetically heterogeneous. The most common etiologies of WWP are mutations in 2 glycosyltransferase, protein O-mannosyltransferase 1 and 2 (POMT1 and POMT2), POMT1 and POMT2 attach the first sugar in the O-mannose-linked glycan moiety of the α-dystroglycan protein, a transmembrane glycoprotein expressed on the surface of muscle cells and neurons. β1-dystroglycan protein interacts with several extracellular matrix components in the basal membrane, and disruption of its function is thought to underlie the severe defects in muscle, eye and brain development in WWP. However, mutations in POMT1 were identified in 7–20% of WWP cases and mutations in POMT2 were found only in 7% of patients. Mutations in POMGNT1, FKRP LARGE and FKTN have been reported at much lower frequencies among WWP patients [13].

Manzini et al. [14] analyzed the most frequent mutations in 43 individuals with WWP. They observed striking differences in the geographic distribution of mutations. Notably, a specific mutation was found in the FKTN gene in all 3 individuals of Jewish Ashkenazi origin with WWP. Furthermore, a carrier rate of 0.7% for the (FKTN) c.1167_1168insA (p.F390fs) mutation was found in Ashkenazi Jews in Israel, suggesting a founder effect in this specific population.

Chang et al. [15] identified the same FKTN mutation in 4 families with WWS and also among 2 of 229 normal American Ashkenazi Jewish adults (0.7%), thus confirming the previous report.

In the present case we used cutting edge prenatal diagnostic tools (imaging, genetics and pathology) in order to provide a diagnosis very early in pregnancy. The transvaginal ultrasound performed prior to CVS at 11 weeks of gestation revealed significant features consistent with WWP. The brain findings of prominent lateral ventricles, a giant cisterna magna and the double kink of the mesencephalic-pontine junction are typical for WWP. In addition, the ocular finding of orbital asymmetry with unilateral cataract strongly suggests a fetus affected by WWP. All these anomalies are usually described on fetal MRI in the third trimester.

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**Fig. 2** Molecular genetic: The c.1167insA mutation in the FKTN gene was identified as homozygous state in the fetus and heterozygous state in the mother.

**Fig. 3** Post-abortion axial T2-weighted brain MRI showing enlarged cisterna magna (CM) with asymmetry of the orbits. Note that the BPD is 13.6 mm.
We have shown that the posterior fossa abnormalities appear first, while dilatation of the lateral ventricles which progresses to overt hydrocephalus is a secondarily developing process. Similarly, we have noticed that the anomalies of the eye may start with a smaller orbit and cataract early at gestation, while the retinal detachment and HPVP may appear later on [8, 11].

We have already shown that Dandy Walker malformation and eye development could be imaged at a very early gestational age [16, 17]. Therefore, it is not surprising that we were able to describe these typical findings for WWP so early in gestation.

We have shown that with appropriate preparation we can obtain a good specimen of a small fetal head for further evaluation even when D & E is performed in early gestation. Indeed this is the first report of using MRI on the product of abortion at 12 weeks gestation and the first to describe the microscopic cortical abnormalities in a fetus with WWP. Additionally, the specific molecular genetics evaluation of the FKTN enabled definitive diagnosis. All the above information led to early decision making before most of the typical signs usually appear. This is a great advantage for patients who wish to terminate pregnancy long before fetal viability when it is still possible according to local law.

Our case is unique since it describes very early changes in the brain and the eyes in a fetus with a priori risk of having WWP. Thus it sheds some light on the time sequence of the appearance of the anomalies in this particular genetic disease. We show that close collaboration between obstetricians, geneticists and pediatric radiologists improve our ability to manage complicated cases for the benefit of our patients. A thorough search for the molecular basis of this syndrome should follow any fetal brain and eye anomalies.

In addition, the FKTN mutation should be added to the genetic screening programs offered to the Jewish Ashkenazi population.

References
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