OEIS COMPLEX (OMPHALOCELE-EXTROPHY-IMPERFORATE ANUS-SPINAL DEFECTS): A CONFUSING SYNDROME.

CASE REPORT

Keywords: Meningomyelocele; Anus, Imperforate; Neural Tube Defects; Bladder Exstrophy.

Palabras clave: Meningomielocele; Ano imperforado; Defectos del tubo neural; Extrofia de la vejiga.

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Introducción. El complejo OEIS es un conjunto de defectos polimalformativos con baja incidencia y prevalencia mundial que suele estar asociado a causas epigenéticas y genéticas que ocasionan alteración al final de la blastogénesis, dando como resultado la asociación de cuatro malformaciones clásicas: omfalocele, extrofia vesical, ano imperforado y lesiones de la médula espinal. En ocasiones también se presenta espina bífida, diástasis de la síntesis púbica y anormalidades en las extremidades.

Presentación del caso. Paciente femenina de 7 meses de edad (al momento de la elaboración del presente reporte), procedente de un área rural colombiana, producto de una tercera gestación con alto riesgo obstétrico y diagnosticada prenatalmente con un defecto en el plegamiento caudal de la pared abdominal y un lipomeningocele. Durante el nacimiento se evidenció extrofia vesical, ano imperforado y disrafismo espinal, lo que permitió plantear el diagnóstico de complejo OEIS e iniciar manejo interdisciplinario pertinente.

Conclusiones. El complejo OEIS es una polimalformación fetal con signos y anomalías características, en donde los conocimientos sobre la etiopatogénesis, el diagnóstico pre y postnatal, el asesoramiento genético y las propuestas terapéuticas son primordiales para favorecer el manejo precoz de las diferentes comorbilidades, aliviar la sintomatología aguda, reducir múltiples comorbilidades y mejorar la calidad de vida del paciente.

Introduction: The OEIS complex is a group of polymorphic defects with low incidence and prevalence worldwide. It is associated with epigenetic and genetic causes that occur in early blastogenesis, resulting in 4 classic malformations consisting of omphalocele, bladder/cloaca extrophy, imperforate anus, and spinal cord injuries. Spina bifida, symphysis pubis diastasis and limb abnormalities may also be observed.

Case presentation: 7-month-old female patient (at the time of writing this report). The mother was from a rural region of Colombia, and this was her third pregnancy, which was at high risk of obstetric complications. The infant was prenatally diagnosed with a caudal folding defect in the abdominal wall and a lipomeningocele. During birth, bladder extrophy, imperforate anus and spinal dysraphism were observed, leading to a diagnosis of OEIS complex. Relevant interdisciplinary management was initiated.

Conclusions: The OEIS complex is a fetal polymorphic malformation with characteristic signs and defects. Knowledge on its etiopathogenesis, pre- and postnatal diagnosis, genetic counseling and therapeutic approaches are essential to favor the early treatment of different comorbidities, alleviate acute symptoms, reduce multiple comorbidities and improve the patient’s quality of life.
INTRODUCTION

The OEIS complex is a rare polymorphic malformation. Its worldwide prevalence has been estimated between 0.04 and 0.05% in live newborns or 1 case per 200,000-400,000 pregnancies, with a male-to-female ratio of 1:2 (1,2). According to Mallikarjunappa & Ghosh (3), the first case of this syndrome was reported in 1709 by Littre. However, as stated by Keppler-No reuil (4), it was not until 1978 that Carey et al. described a congenital syndrome with multiple abdominal wall malformations, including omphalocele, cloacal exstrophy, imperforate anus, and spinal defects, which was called the “OEIS complex” (3,4). Other conditions associated with this disorder have also been discovered over time, including spina bifida, congenital urological anomalies, renal anomalies, pubic symphysis diastasis and limb abnormalities (5).

The spinal anomaly associated with the OEIS complex is occult spinal dysraphism (6), which is defined as a group of congenital malformations of the spine and spinal cord characterized by failure of fusion (total or partial) of neural structures, bone, and midline mesenchymal fields (7).

OEIS is considered a complex as it comprises morphological defects that share a common or adjacent embryological region. Its etiology is not yet clear, but it is believed to involve genetic and epigenetic factors (8). The degree of malformation depends on the prenatal period in which the primary defect occurs (9). Its prognosis is unfavorable, so early family management and counseling is always necessary.

The present article describes the case of an infant diagnosed with OEIS complex in order to emphasize the scarcity of information on this entity, especially in Latin America, and to inform about the treatment options available to date.

CASE PRESENTATION

This is the case of a 7-month-old female patient (at the time of writing this report). The mother came from a rural region of Colombia and this was her third pregnancy. Her parents were not related by blood and were low-income farmers. The pregnancy was at high obstetric risk due to advanced maternal age and prolonged intergenesic period (>11 years). The mother did not report exposure to toxics or psychoactive substances during pregnancy and had adequate prenatal check-ups. Her TORCHS profile was negative. A prenatal ultrasound scan (at 20 weeks) allowed diagnosing omphalocele associated with abdominal wall malformations. Fetal karyotyping revealed 46XX.

The patient was delivered through a cesarean section at 34 weeks gestation due to preterm labor caused by pre-eclampsia with severity criteria: APGAR 5/10 per minute, 7/10 at 5 minutes, and 8/10 at 10 minutes. At birth, her weight was 2,215g; head circumference was 32cm; height was 43.5cm. Induration in the lumbosacral region, bladder exstrophy due to a defect in the midline of the abdominal wall, bilateral talipes equinovarus and cloacal malformation were observed. Therefore, a possible OEIS complex was considered and other differential diagnoses such as gastroschisis, limb-body wall complex and pentalogy of Cantrell were ruled out.

On the third day of life, the baby was taken to skin vesicostomy, omphalocele closure, tubularization of the colonic pouch and intestinal bypass, with favorable postoperative evolution. During her hospital stay, a cranial ultrasound was performed, finding no alterations. A urinary tract ultrasound also showed grade 4 hydronephrosis, necessitating a right nephrostomy. Two days later, the infant was assessed by the
pediatric neurology and neurosurgery services, which established that she presented with hypotonia and a lumbosacral mass of approximately 4x4cm, paraparesis, and hyporeflexia of lower limbs and talipes equinovarus. As a result, MRI of the neuraxis was performed, revealing lumbosacral lipomeningocele, type II diastematomyelia (Figure 1), and sacral agenesis. Occult spinal dysrafism was considered that did not require immediate intervention was considered. However, this condition had to be followed up on an outpatient basis and treated through rehabilitation with physical and occupational therapy.

At 6 months, the patient was taken back to surgery for a bilateral pelvic osteotomy and colostomy remodeling by the pediatric orthopedic and pediatric surgery services. These procedures allowed for proper positioning and bone consolidation in the pelvis and did not cause further complications.

At the time of writing the present case report, the patient, aged 7 months, did not present any significant clinical deterioration, had adequate response to the treatment provided to alleviate her symptoms, and did not develop any postoperative complications despite the uncertainty of her prognosis. Furthermore, she was awaiting spinal anchoring and interdisciplinary follow-up by the pediatric neurology, neurosurgery, urology, pediatric surgery, interventional radiology, and physical and occupational therapy services. Clinical genetics requested genomic hybridization + microarray to identify any pathogenic variant, deletion or copy number variation that could explain the etiology, but no report has yet been received.

**DISCUSSION**

The first case of a patient with an OEIS complex was reported in 1709. However, Carey et al. characterized this entity for the first time in 1978 (4,9,10) after identifying 175 children with one or more of the following malformations after reviewing the medical records from a California hospital: omphalocele, cloacal exstrophy, imperforated anus, and spinal defects. According to Austin et al. (11), Meizner was the first to perform an ultrasound diagnosis of the OEIS complex in 1985.

OEIS is defined as a complex since it comprises a group of defects that share an embryological region and stage. It occurs at the end of blastogenesis (fourth gestational week) (12), when important embryological processes occur, such as closure of the neural tube, transverse and longitudinal folding of the embryo (formation of the anterior chest and abdominal wall), development of the midline and laterality, disappearance of the chorionic cavity due to the expansion of the amniotic cavity, formation of the umbilical cord and the cardiovascular system, onset of kidney development and initial growth and patterning of the limbs (13,14), explaining the characteristic polymorphic malformations of the entity.

Concerning pathogenesis, there are several theories that postulate four major defects: 1) failure in the formation of the urorectal septum,
which prevents the separation of the urogenital and anorectal tract; 2) total rupture of the cloacal membrane and failed junction of genital tubercles and pubis branches; 3) alteration of ventral abdominal wall closure secondary to abnormal lateral folding, and 4) incomplete development of the lumbosacral vertebrae and failure of cranial neural tube closure (15,16). The reported patient showed phenotypic effects associated with the above-mentioned mechanisms. The first caused persistent cloaca, imperforate anus and genitourinary alterations such as hydronephrosis; the second caused cloacal exstrophy; the third led to an abdominal wall defect, and the fourth led to occult spinal dysraphism with diastematomyelia and tethered cord syndrome, as well as sacral agenesis.

Most OEIS complex cases are isolated and caused by a multifactorial alteration involving environmental factors such as smoking and exposure to benzodiazepines during pregnancy (17) and genetic factors such as deletion of 9q 34.1-q, 1p36 and 3q12.2-3q11.2, trisomy 18, mosaic Turner syndrome, mutations in mitochondrial 125rRNA and mutations in homeobox genes such as HLXB9 since recurrence has been reported in some families with monozygotic conjoined twins (2,18-20). Almost all cases are premature with low birth weight; however, there are reports of patients with an average gestational age of 37.5 weeks and proper weight, height, and head circumference for age (21).

Cohen (22) suggests that the proximity of the neural tube to the cloaca during embryonic development may explain cloacal abnormalities related to occult spinal dysraphism, an entity that includes a broad spectrum of congenital fusion abnormalities of one or more dorsal midline structures. These abnormalities can affect the skin, subcutaneous tissue, vertebral bodies, meninges, and neural tissue.

There are two categories of occult spinal dysraphism: open spinal dysraphism, associated with skin defects and exposed neural tissue, and the closed spinal dysraphism, characterized by subcutaneous masses (Figure 2).

![Occult spinal dysraphism classification](image)

**Figure 2. Classification of occult spinal dysraphism**

* It accounts for 98% of all cases of spinal dysraphism.
† It accounts for 2% of all cases of spinal dysraphism.

Source: Own elaboration based on Copp et al. (23) and Wallingford et al. (24).
Although myelocystocele is the most common spinal dysraphism in patients with OEIS (25), the reported patient had a lipomeningocele, which is a herniation of cerebrospinal fluid, meninges, and neural tissue through the posterior spinal bone defect. It is also associated with the presence of lipomas that extend from the subcutaneous cell tissue to the spinal canal and with diastematomyelia, which is characterized by a "splitting" of the spinal cord in one or more segments. According to Tortori-Donati et al. (26), it is classified into two types:

**Type I:** The arachnoid mater and the dura mater do not divide. There is only one dural sac for both hemicords. It represents 60% of diastematomyelias and 50% occur without bony spur.

**Type II:** The arachnoid mater and the dura mater divide into two and contain both hemicords, so each has its own subarachnoid space that joins up and down forming a single subarachnoid space. 95-100% occur with bony spur (27,28).

Alterations associated with the OEIS complex include cardiac anomalies; kidney anomalies, as in the patient reported here, who presented with grade 4 right ureterohydronephrosis that required percutaneous nephrostomy, increased nuchal translucency, and elevated serum alpha-fetoprotein (29,30).

For prenatal diagnosis, several ultrasound criteria have been classified as major and minor (11) (Table 2). Nevertheless, despite the existence of such criteria, not all anomalies can be identified prenatally and are often mistaken for some differential diagnoses such as omphalocele or gastroschisis. Therefore, the diagnosis is usually confirmed with other imaging aids such as magnetic resonance or color Doppler, used to ratify bladder extrophy and differentiate it from the omphalocele (10,31).

<table>
<thead>
<tr>
<th>Major criteria</th>
<th>Minor criteria</th>
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<tbody>
<tr>
<td>Difficulty in bladder visualization</td>
<td>Lower limb defects</td>
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<tr>
<td>Infraumbilical defect of the anterior abdominal wall</td>
<td>Kidney abnormalities</td>
</tr>
<tr>
<td>Omphalocele</td>
<td>Ascites</td>
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<tr>
<td>Lumbosacral myelomeningocele</td>
<td>Widened pubic arches</td>
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<td></td>
<td>Hydrocephalus</td>
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<td>Narrow thorax</td>
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<td></td>
<td>Single umbilical artery</td>
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<td></td>
<td>Prolapse of intestinal ileum into the amniotic cavity</td>
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</tbody>
</table>

*Observed in more than 50% of patients.

Source: Own elaboration based on Copp et al. (33)

Patients with OEIS complex require immediate multidisciplinary care, followed by individualized surgical treatment to close the abdominal wall safely, prevent short bowel syndrome and urinary and fecal incontinence, preserve kidney function, and achieve functional and aesthetic genital reconstruction. While most cases are sporadic, family cases have been reported, which could suggest one or more specific genes that may play a key role. This makes genetic counseling and study of the utmost importance (8,34).

The prognosis of patients with OEIS complex varies depending on the severity of structural defects, the extent of cloacal extrophy (due to its renal and pulmonary complications), and the severity of the neural tube defect. Thus, an adequate interdisciplinary management of the less severe forms may improve prognosis and lethality of this entity.

The patient in the case described here had the advantage that her OEIS complex was not lethal and that she was treated in a hospital with various medical specialties where she could undergo diagnostic imaging and receive the required medications, which allowed for a more comprehensive diagnosis-treatment with greater benefits. However, it should be noted
that at the time of the writing this article the patient had not undergone all the procedures indicated since this is an entity that requires life-long interventions, management, and surveillance.

CONCLUSIONS

The OEIS complex is a rare and complicated congenital condition that can only be detected in less than 25% of cases by ultrasound in the second trimester of gestation due to the wide spectrum of anatomical variants. This occurs because malformations depend on the degree of cloacal septation and, therefore, other imaging resources such as MRI or color Doppler are often required to confirm the diagnosis.

The prognosis of the OEIS complex is unfavorable, so an interdisciplinary team of neonatologists, geneticists, pediatricians, urologists, pediatric surgeons, neurosurgeons, orthopedists, pediatric neurologists, radiologists and maternal-fetal specialists is needed to provide parents with comprehensive counseling to define pre-birth management, plan appropriate perinatal management, and achieve a better quality of life for patients. For this reason, it is important to raise awareness of this entity since an adequate clinical approach will help to better guide treatment and prognosis.

ETHICAL CONSIDERATIONS

The informed consent of the patient’s parents was obtained for the present case, while privacy and anonymity were preserved.

CONFLICTS OF INTEREST

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