Case Report

The OEIS complex – clinical & radiological evaluation

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Abstract

The OEIS complex comprises a constellation of complex and severe malformations of the abdominal wall, gastrointestinal and genitourinary tracts, and spinal cord. The malformation results from improper closure of the ventral abdominal wall due to failure of convergence of cephalo-caudal and lateral folding of the embryo during early gestation. The rarity of the condition suggests etiologic heterogeneity and the possible role of environmental and genetic factors. We present clinical and imaging findings of the OEIS complex in a neonate.

Introduction

OEIS complex includes a spectrum of complex and severe malformations involving multiple organ systems. The anomaly comprises omphalocele, cloacal exstrophy, imperforate anus, and spinal abnormalities. Cloacal exstrophy is the most severe form of the exstrophy–epispadias complex. The first description of the anomaly was published by Littre in 1709 and Carey et al first coined the term OEIS complex [1,2]. Reported incidence of OEIS varies from 1 in 2,00,000 to 4,00,000 pregnancies [3–5]. The rarity of the condition suggests etiologic heterogeneity and the possible role of environmental and genetic factors [6].

Case report

A 34 weeker, low birth weight baby born by normal vaginal delivery to a second gravida mother presented on day 2 with lower abdominal wall defect with exposed urinary bladder, through which bowel loops were protruding out and imperforate anus [Figure 1A]. A small exomphalos with a central cord was present just above the abdominal defect. The baby had a bifid penis and well developed scrotum with a descended and one undescended testis. An anal opening was absent and the patient was passing stool through the protruding bowel loop. There was a skin covered raised area with the consistency of fat noted in the midline over the lumbosacral region. The lower limbs below the knees showed minimal movement and without any reflexes. No abnormalities were noted in the upper abdomen, chest, upper limbs, and head. The baby was on breast feed.

Radiological findings

X-ray AP view of pelvis and lumbosacral spine revealed a wide pubic diastasis and defects involving a body of multiple sacral vertebrae (Figure 1B).

USG of Lumbo–sacral region showed:

- Hypoechoic cystic lesion containing echogenic nerve roots within and tethered cord in sacro–coccygeal region with intact overlying skin and subcutaneous tissue (Figure 2A).
- Proximal syrinx formation in the region of the lower lumbar and sacral spinal canal was also noted.
MRI of lumbo-sacral spine confirmed the above findings suggesting a closed spinal dysraphism -- terminal myelocystocele with tethered cord at sacro-coccygeal junction and syrinx formation (Figure 2B).

Discussion

Although the first description of a case of OEIS complex was reported in 1709, it was only in 1978 that Carey et al first used the term OEIS complex [1,2]. The anomaly is thought to be due to a defect in early blastogenesis or of mesodermal migration during the primitive streak period [7,8]. These defects lead to improper closure of ventral abdominal wall due to failure of convergence of four ecto-mesodermal folds (a cranial, a caudal, and two lateral) of an embryo with associated defects in the development of cloaca and urorectal septum during 4th gestational week. The cloacal membrane is composed of ectodermal and endodermal layers and it covers the region from the umbilicus to the tail portion of the embryo at the 4-mm stage. In subsequent stages, the primitive streak mesoderm invades the membrane to form the lower abdominal wall. Lack of mesoderm in the infra-umbilical abdominal wall results in omphalocele. During the 8– to the 16-mm stage, the urorectal septum meets the cloacal membrane and divides the cloacal chamber into an anterior urogenital and posterior alimentary systems. Cloacal extrophy results when the membrane ruptures before the formation of the urogenital septum. The coexistence of cloacal extrophy and spinal dysraphism may be explained by a single insult in the embryonic tail in early pregnancy [6]. Caudal dysgenesis interferes with somite formation resulting in defective vertebrae [9].

Because of its rarity and sporadic occurrences, some environmental and genetic factors may be responsible for its causation [6]. One study reported maternal exposure to diphenyl hydantoin and valproic acid and cigarette smoking as possible risk factors [2,6,9]. Higher incidence of OEIS in monozygotic twins than in dizygotic twins suggesting a possible genetic contribution to the occurrence of these defects was reported [10].

The OEIS complex involves multiple body systems. The classical presentation consists of omphalocele superiorly with an open plate mucosa inferiorly consisting of two posterior walls of hemibladder on either side with a central strip of the tunestinal mucosa [11]. Similar findings were noted in our case with the passage of meconium and stool from the protruding bowel loop in the upper part of the central bowel strip. Ambiguous genitalia is common with bilateral cryptorchidism in males. In the present case, one testis was found in the hemiscrotum, but the other remained high in the inguinal canal. Renal anomalies such as agenesis, rudimentary kidney, ectopic kidney pelviureteric junction obstruction etc have been reported [12]. The congenital vertebral malformations associated with cloacal extrophy vary from congenital scoliosis, kyphosis, abnormal lumbosacral segmentation, sacral agenesis, interpedicular lumbar widening to hemivertebra, spina bifida occulta, and myelomeningocele [13,14]. In the present case, there was terminal myelocystocele with tethered cord and syrinx formation. The baby had a partial neurological deficit in both lower limbs below the level of the knee. Talipes in association with OEIS complex is related to a tethered cord. Central nervous system anomalies are not commonly encountered in OEIS complex.

OEIS complex is difficult to diagnose prenatally and all the abnormalities in the fetus may not be clear until a thorough postnatal evaluation [15]. The major criteria for the prenatal diagnosis include non-visualization of fetal bladder, infraumbilical anterior abdominal wall defect, omphalocele and myelomeningocele [15,16]. The minor criteria include lower extremity abnormalities (talipes), renal anomalies, ascites, widened pubic arches, narrow thorax, kypho-scoliosis, hydrocephalus and single umbilical artery. Presence of omphalocele and spinal defects on prenatal ultrasonography strongly suggest OEIS complex.

Overall prognosis of babies born with OEIS complex depends on severity of the anomalies and presence of infection. With surgical intervention and improved medical therapy, the survival of these patients has been greatly improved. Infants who are medically stable may be considered for surgical reconstruction with closure of omphalocele, separation of ileocecal connection from the extrophied bladder plate, and reapproximation of an end colostomy with preservation of all bowels [14]. Approximation of the pubis usually with osteotomy is beneficial in reconstruction of the pelvic ring and bladder closure [16]. Bowel and urinary incontinence may persist even after good reconstruction because of neurological abnormalities. Spinal cord untethering may be required years after closure of the spinal defect [14,17]. Early evaluation with
magnetic resonance imaging will allow rapid treatment and prevention of further loss of function. Male patients may have infertility or subfertility issues and erectile dysfunction.

**Conclusion**

Babies born with OEIS complex require the care of a multidisciplinary team of neonatologists, pediatric surgeons, and pediatric urologists. These patients need multiple surgeries, with many potential complications. There is no definite etiological factor responsible for this anomaly. Prenatal diagnosis is possible based on specific ultrasonologic criterias.

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