A diprosopus monocephalus tetrophthalmos: a case report and review of literature

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Abstract

Objective: The aim of this paper was to present and discuss a case of double-face or diprosopus of a rare variant of conjoined twins.

Case: A 24-year-old patient at 21 weeks of gestation having gravida 3, 2 living children and no significant family history was referred to our clinic with the diagnosis of hydrocephaly. Intrauterine assessment of the obstetric ultrasonography showed the fetus with one head, two faces, one body, two arms and legs, a single, unified spinal cord and vertebral column compatible with 21 weeks of gestation.

Conclusion: With the spread of prenatal follow-up, early detection of cases with conjoined twins, such as diprosopus, is essential in terms of social, economic and ethical aspects, and it will enable parents to decide at early weeks.

Key words: Diprosopus, monocephalus, tetrophthalmus.

Introduction

Conjoined twins are rare and interesting phenomenon, and the incidence reported for craniofacial duplication or diprosopus (two face, single head and single body) is 180,000/15,000,000. Conjoined twins have high mortality rate. Approximately 40% of them results in abortus, and 1/3 of the cases die less than 24 hours after birth. Although its etiology is not known well, they have a wide spectrum from double nasal structure to double face, four eyes (tetrophthalmos) and single head (diprosopus monocephalus).

The oldest diprosopus case known was first stated in the works of Ambroise Pare in 16th century. Many diprosopus cases with single body and normal extremities are complete duplication and there are severe defects in the central nervous system. On the other hand, incomplete diprosopus cases (duplication of maxilla, mandible and oral cavity) have normal central nervous system. In this study, we aimed to evaluate prenatal diagnosis and management of a diprosopus case diagnosed on 21st week of gestation, through the literature.

Bir diprosopus monosefalus tetroftalmus olgusu ve literatürün gözden geçirilmesi

Amaç: Bu makalede yapışık ikizin nadir bir varyantı olan çift yüz veya diprosopus olgusunu sunmak ve tartışmak amaçlanmıştır.


Sonuç: Prenatal takibin yaygınlaşması ile diprosopus gibi yapışık ikiz olguların erken döneme tespit edilmesi, sosyal, ekonomik ve etik açıdan önemli olup, ebeveynlerin erken haftalarda karar vermesini sağlayacaktır.

Anahtar sözcükler: Diprosopus, monosefalus, tetroftalmus.
Case Report

A 24-year-old patient at 21 weeks of gestation having gravida 3, 2 living children and no significant family history was referred to our clinic with the diagnosis of hydrocephaly from another clinic. In the intrauterine assessment by the obstetric ultrasonography (Voluson 730 PRO, General Electric, Fairfield, Connecticut, USA) of the patient at 21 weeks of gestation showed the fetus with one head, two faces, four eyes (Fig. 1), one body, two arms and legs, hydrocephaly, and compatible with intrauterine 21 weeks of gestation. The findings were confirmed by MRI (Figs. 2 and 3).

In the transverse and longitudinal views, it was found that fetus had one heart, one stomach, one bladder, one placenta and one umbilical cord. The patient and her husband were informed about the condition. The case was discussed in the ethics committee of our clinic. After the written consent of the patient and her husband was obtained, termination was decided. After termination, it was observed in the macroscopic examination that fetus had one head, two faces, one body, two arms and legs (Fig. 4). The parent did not allow an examination on fetus after delivery.

Discussion

Conjoined twins are quite rare congenital anomaly seen in 1/50,000-1/100,000 cases. Our case was a craniofacial duplication or diprosopus case which is a variant of conjoined twins. The conjoined twins are independent from race, heredity, delivery number or kinship. Although the etiology of conjoined twins is not known well, there are two theories proposed: According to the first theory, monovular embryo is divided incompletely on 13th-15th days of conception. In fusion theory, a secondary fusion occurs among two monovular embryonic disks.

When fusions with different pattern are compared, central-located conjoined twins are seen more frequently. Although complex problems within a wide
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Spectrum which may include all organ systems are observed in conjoined twins, the most severe anomalies are seen on fusion location. Diprosopus shows full duplication of face structures. Cleft palate and lip anomalies as well as central nervous system, cardiac, gastrointestinal and respiratory system anomalies are observed at a high rate. In our case, there was hydrocephaly. As in our case, generally female gender is dominant. Chromosomal analysis results of such cases are generally normal. We could not conduct a chromosomal analysis in our case since the parent did not allow.

It is believed that diprosopus occurs as a result of a fault in the neurulation of embryo. Notochord defines the axis of embryo and induces neurulation. Peripheral neural crest cells create mesenchymal connective tissue of face and mouth areas, and pharyngeal arch. A fault in the neurulation causes neural tube to fail to be closed. Central nervous system anomalies frequently accompany. In our case, hydrocephaly was observed as neural tube defect.

Prenatal diagnosis of cases with diprosopus was reported in the first trimester. Maternal alphafetoprotein levels were found to be high in some cases. Low fetoprotein was reported only in a case with cranioschisis. Although polyhydramnios is observed in conjoined twins more frequently as a prenatal sonographic finding compared to normal twins, amniotic fluid was normal in our case. Interestingly, Onuf mentioned excessive amount of amniotic fluid by the examination before ultrasound. Other sonographic findings are irregular cranium, enlargement of vertebra, and head in heart shape or with two equal parts. Facial duplication is shown clearly by sonography.

Central nervous system anomalies were detected in all reported cases with diprosopus. In our case, hydrocephaly and double faces were seen prenatally. Conjoined twins can be recognized at first trimester easily by ultrasonography. If a single yolk sac and two fetuses or monoamniotic twins are observed by ultrasonography, the possibility of conjoined twins should be kept in mind. Cases which have advanced fusion like diprosopus can be recognized in advanced weeks.

Conclusion

Consequently, diprosopus is one of the conjoined twin types and usually central nervous system anomalies accompany it. In our region, diagnosing such cases occur only in advanced weeks of gestation due to irregular prenatal follow-up. With the spread of prenatal follow-up, early detection of cases with conjoined twins, such as diprosopus, is essential in terms of social, economic and ethical aspects, and it will enable parents to decide at early weeks.

Conflicts of Interest: No conflicts declared.

Fig. 4. Diprosopus monocephalus.
References


15. Onuf B. A case of double formation of the face with craniorachischisis involving the whole vertebral column. Med Rec 1895;48: 401-4.