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Mehnoosh Torkzaban  
Department of Radiology, Thomas Jefferson University Hospital, Philadelphia, PA, United States

Nesa Rajabpoor Nikoo  
Shohada Hospital, Qom, Iran

Ateeh Kalateh  
Department of Obstetrics and Gynecology, North Khorasan University of Medical Sciences, Bojnurd, Iran

Mahboobeh Shirazi  
Maternal, Fetal and Neonatal Research Center, Tehran University of Medical Sciences, Tehran, Iran

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Analysis of Ultrasonographic Misdiagnosis of Cephalothoracopagus Janiceps Conjoined Twins: A Case Report

Mehnoosh Torkzaban1, Nesa Rajabpoor Nikoo2, Ateeh Kalateh3, Mahboobeh Shirazi4*

1. Department of Radiology, Thomas Jefferson University Hospital, Philadelphia, PA, United States
2. Shohada Hospital, Qom, Iran
3. Department of Obstetrics and Gynecology, North Khorasan University of Medical Sciences, Bojnurd, Iran
4. Maternal, Fetal and Neonatal Research Center, Tehran University of Medical Sciences, Tehran, Iran

Introduction

Conjoined twining (CT) is a rare complication of monozygotic twinning. The frequency of CT is approximately 2:100,000 gestation, while the true incidence is around 0.5: 100,000 live births (1,2). The incidence is three times higher among female than male twins (2). In humans, monozygotic monoamniotic twins develop when an implanted blastocyst splits at about 9–13 days after fertilization. Failure to undergo a complete spillage of the embryonic disc at or later than 14 days results in conjoined twin development (fissure theory) (2,3,4). The etiology of CT is obscure and none of race, consanguinity, hereditary, birth order, maternal age, acute and chronic diseases during pregnancy, or maternal exposure has revealed any association with CT (3,5). Conjoined twining has been reported in twine and triplet pregnancies following assisted reproductive techniques such as in-vitro fertilization and intracytoplasmic sperm injection (6,7). Conjoined twins are classified according to the most prominent site of attachment, plus “pagus”, a term that means “fixed” in Greek (2). Cephalothoracopagus is the rarest subtype of CTs and occurs once in every 58 sets of conjoined twins or once in every three million births (8,9); it refers to twins with fused head and thorax but separate pelves and limbs. If the single fused head has two faces each looking in opposite directions, the twins are referred to as Cephalothoracopagus janiceps (8,10). The term janiceps has its origin in Janus, the two-faced Roman god. When the two faces are asymmetrical and only one normal face can be seen alongside with a second incomplete face (single naris, cyclopic eyes, synotic ears, a proboscis, or two small eyes in a single palpebral fissure), it is called cephalothoracopagus.
janiceps monosymmetros (10). Here we present such a case.

**Case Report**

A 33-year-old G2P1 pregnant woman, who had conceived spontaneously and had an unremarkable medical history with no history of drug, alcohol or cigarette smoking, underwent her early pregnancy ultrasound. The scan revealed one fetal pole, one yolk sac, and a normal fetal heart rate (FHR). The crown-rump length (CRL) was 12 mm equal to 7 weeks 3 days of pregnancy. She was scanned next time for aneuploidy screening ultrasound at 11 weeks, 4 days of the last menstrual period (LMP). The gestational age (GA) based on the CRL was equal to 12 weeks, 1 day of pregnancy. The thickness of the nuchal translucency (NT) was reported as 4 mm. The radiologist had reported possible technical errors in NT measurement and inability in nasal bone (NB) visualizing due to unfavorable fetal position. A hyperechoic herniated mass was also visualized at the level of the anterior abdominal wall and GastroSchisis was suspected. The patient was referred to a tertiary health center, (Yas Hospital, Tehran, Iran), for further evaluation. The next ultrasound exam was performed at 13-14 weeks of pregnancy. Unexpectedly, two alive fetuses were reported fused in head, thorax, and abdomen. The ultrasound diagnosis of cephalothoracopagus CTs was made. Because of the poor prognosis, counseling was provided for parents and elective medical pregnancy termination by misoprostol was carried out. The expelled twines were fused at the cranium, thorax, and abdomen to the umbilicus. Twines had a shared placenta and umbilical cord. The faces were dissimilar and on opposite sides of the head, 90 degrees to the separate spines. There was one normal face with two eyes, one complete nose, and one mouth, and one set of low-set ears (Figure 1A), and the opposing face had incomplete facial features as one cyclopic eye and irregular fused nares and mouth orifices (Figure 1B). The arms and legs of each of the twins were of similar length and development and followed the orientation of the spine (Figure 1C). The twines’ sex was not obvious but more likely were females.

**Figure 1.** A. Normal face with two eyes, one complete nose and one mouth, and one set of low set ears; B. Incomplete facial features as one cyclopic eye and irregular fused nares and mouth orifice; C. The arms and legs of each of the twins were of similar length and development and followed the orientation of the spine

**Figure 2.** A. Longitudinal section of twins, mimicking singleton pregnancy at 11 weeks 6 days of pregnancy; B. Fused brains at 13-14 weeks of pregnancy; C. Two separate beating heart at 13-14 weeks of pregnancy; D. Longitudinal section of twins at 13-14 weeks of pregnancy.
Discussion

The worldwide use of the ultrasound in prenatal screening for Down syndrome provides the diagnosis of conjoined twinning in early pregnancy. Since all CTs are monochorionic monoamniotic, when only one yolk sac is seen alongside two embryos in very early pregnancy, the index of suspicion should be raised. Two of first-trimester sonographic indications for possible conjoined twinning are: 1) bifid appearance of a sole fetal pole in the first trimester seen before 10 weeks of gestation, and 2) visualization of two hearts or two stomachs (3). From 8 weeks of gestation, fetal activity increases progressively, which facilitates the differential diagnosis between conjoined and non-conjoined monoamniotic twins (1). The first trimester (11-13w+6days) ultrasound is the best method for early diagnosis of CTs (3).

The earliest prenatal diagnosis of cephalothoracopagus twins was reported by Lam et al. 1998 (11). The diagnosis was made by vaginal ultrasound examination at 8 weeks' gestation from the findings of a single fetal pole with an irregular body outline, a disproportional large head, and two separate cardiac pulsations and was confirmed after rescanning at 11 weeks (11). While early diagnosis of conjoined twins is feasible, it is not always easy in cephalothoracopagus twinning due to extensive conjoining; a single fused head and thorax, and limited fetal movement as was seen in our case.

Due to the monoamniocity of the conjoined twins, a single yolk sac alongside a single CRL and normal fetal heart were observed at 7-8 weeks of gestation in our case; mimicking a singleton pregnancy and substantial missing of the second fetal heart (Figure 2A). When the second ultrasound scan was performed at 11 weeks 4 days of LMP in our case for NT measurement, the operator reported a fetus with normal fetal heart, increased NT, non-visualized NB, and herniated mass at the ventral abdominal wall, but yet to notice the second fetal heart and fused brains. Following the detection of the aforementioned anomalies, a thorough evaluation of the fetal thorax and cranium via scanning the entire uterus both in a longitudinal and transverse approach could be helpful in revealing two separate hearts, fused brains, and duplicated extremities, leading to the accurate diagnosis of CTs. Increased NT, cystic hygroma, and abdominal wall defects are the most common findings in conjoined twins (12). Currently, certain congenital fetal anomalies are diagnosed even before the twentieth week of gestation with the use of high-resolution ultrasound equipment, but without following a standardized approach to fetal ultrasound and in case of lack of suspicion to any abnormal condition, misdiagnosis may happen (13, 14).

The final ultrasound scan was performed at 13-14 weeks of GA in our reported case. The diagnosis of cephalothoracopagus CTs was made by finding one shared placenta and umbilical cord, two fused brains without separating bony skull, two deformed spinal columns, one common thoracic cage with two separate beating hearts and one stomach, two separate sets of lower and upper extremities, and two separate pelvises. One of the twins had cystic hygroma in the posterior neck. The estimated GA based on both twines' approximate CRL was 13 weeks 6 days (Figures 2B, 2C and 2D).

The next step after diagnosis of conjoined twinning is a prognostic assessment to propose timely medical termination of pregnancy in case of non-viability and prevent complicated surgical delivery. In cephalothoracopagus CTs, due to the fusion of the heads and abnormal arrangement of the superior central nervous system, no attempt at surgical separation should be considered and the prognosis is very poor (15). We arranged a medical pregnancy termination accordingly. The recurrence risk for conjoined twins appears to be negligible (5).

Conclusion

In Cephalothoracopagus twining a single yolk sac alongside a single CRL and normal fetal heart at 7-10 weeks of gestation may mimic a singleton pregnancy and lead to missing the second fetal heart and fused brains without applying a standardized scanning of the entire uterus both in a longitudinal and transverse approach. A high level of concerns may raise for conjoined cephalothoracopagus twinning in case of finding a single fetal pole with an irregular body outline and a disproportional large head in the presence of two separate fetal hearts in early pregnancy. Two fused heads with two brains and two sets of lower and upper extremities do confirm the diagnosis.

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Conflict of Interest

Authors declared no conflict of interests.

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