An Unusual Case of Monozygotic Epigastric Heteropagus Twinning

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Epigastric heteropagus refers to unequal and asymmetric conjoined twins in which the dependent component (parasite) is smaller and usually attached to the epigastrium of the dominant component (autosite). Epigastric heteropagus is a very rare type of conjoined twins. It is generally assumed that conjoined twins represent a form of monozygotic twinning, but there has been no confirmation of this monozygotic hypothesis. Epigastric heteropagus twins differ in several ways from symmetrical conjoined twins. These include male preponderance, and no major connection of vessels, bowel, or bones. These significant characteristics support the authors that the fusion of two fertilized ova and the atrophy of one fetus results in this interesting malformation. However, a DNA study confirmed monozygotic origin in their case. The possible pathogenesis was ischemic atrophy of the body structure of the monozygotic conjoined twins at an early gestational age.

INDEX WORDS: Conjoined twins, epigastric heteropagus, monozygotic; DNA study.

Conjoined twins are one of the rarest congenital abnormalities. They occur once in every 50,000 to 100,000 births. Conjoined twins are classified as symmetrical or asymmetrical. The asymmetrical form is known as heteropagus. With heteropagus cases, the dependent portion (parasite) is smaller than the host (autosite). Epigastric heteropagus refers to the conjoined twin in which the parasite is attached to the epigastrium of the autosite. There are only nine similar cases in the English-language literature.

CASE REPORT

A male boy with a conjoined twin was referred to our hospital on the day of birth. His birth history consisted of a normal spontaneous vaginal delivery at 40 weeks of uncomplicated gestation. The mother was 34 years old (gravida 3, para 3). The first baby had syndactyly of one hand. No other congenital abnormality had been documented in the patient’s family history.

Clinical Presentation

The parasite was attached to the epigastrium of the autosite above an omphalocele. The autosite was active and appeared normal except for the omphalocele. The combined birth weight was 3,700 g. The parasite consisted of a lower trunk, a pelvic girdle, and two normal legs. The external genitalia of the parasite presented as a well-formed set of male genitalia with an empty scrotum. The perineum lacked an anal opening. The legs of the parasite did not respond to painful stimuli (Fig 1).

Radiological Examinations

A plain roentgenogram of the parasite showed an accessory bony pelvis with a well-developed set of lower limbs. No vertebral column was present in the parasite (Fig 2). A barium gastrointestinal study of the autosite showed a normal gastrointestinal tract and no crossover into the gastrointestinal tract of the parasite. Contrast-enhanced computed tomography showed a functioning kidney and a bladder in the parasite’s pelvis (Fig 3). Echocardiography of the autosite showed a ventricular septal defect (VSD), atrial septal defect (ASD), and patent ductus arteriosus (PDA).

Surgical Separation

Surgical separation was performed on the seventh day after birth. The parasite was attached to the autosite by skin, subcutaneous tissue, muscle, and fascial layers. The perirenal cavity of the parasite was connected to that of the autosite. The bowel of the parasite was herniated to the omphalocele of the autosite. Both ends of the parasite’s bowel were blind, without any connection to the autosite’s gut. The vascular pedicle of the parasite arose from the falciform ligament of the autosite. The autosite’s intraperitoneal organs were completely normal in appearance. The parasite had one kidney, which drained via one ureter into his own bladder. The parasite was resected from the autosite without difficulty. The abdominal wall defect and the omphalocele were closed without difficulty, and the postoperative course was satisfactory. Correction of the VSD, ASD, and PDA was performed at 8 months of age. The baby (autosite) is healthy, thriving, and normal (Fig 4).

Chromosome and DNA Studies

A chromosome study was performed with peripheral blood samples from the autosite and the parasite, using conventional methods. The karyotypes of both subjects were 46,XY with no evidence of chromosomal aberration. Discrimination of heteropagus zygosity was not possible from the results of chromosome study.

To investigate zygosity, DNA analysis of the heteropagus was performed using the polymerase chain reaction (PCR) technique. Total genomic DNA was prepared from leukocytes of peripheral blood from both the autosite and the parents. The genomic DNA of the parasite was extracted from a paraffin block of kidney tissue from the surgical specimen (parasite).

Haplotyping of the heteropagus and the parents was performed for microsatellites on chromosome 13 (D21S11) and chromosome 21 (D21S11), which contained tetranucleotide repeats polymorphisms. Primer sequences used to amplify each region were as previously described by Brandt et al and Shamra et al. Haplotyping for D21S11 (Fig 5) and Rb1.20 locus (data not shown) showed that the heteropagus had a monozygotic origin.

DISCUSSION

Conjoined twinning is one form of duplication or the “doubling anomaly.” It ranges from simple super-

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The parasitic twin was attached to the host's epigastrium, above an omphalocele. The parasite had a pelvis with two well-formed legs, a penis, and a scrotum. There was no anal opening in the parasite.

Twins can originate in two ways. When they originate from two zygotes, they are considered dizygotic twins. When they originate from one zygote, they are referred to as monozygotic twins. About two thirds of twins are dizygotic. Whereas the frequency of dizygotic twinning shows a marked racial difference, the incidence of monozygotic twinning is about the same for all races. The statistically significant tendency for dizygotic twins to repeat in a family leads us to believe that there is a hereditary link, but this is not so with monozygotic twinning. The tendency for binovular twinning appears to be determined by one or more recessive genes that affect the frequency of multiple ovulation. In certain populations, as in Nigerian blacks, the incidence of multiple ovulation and dizygotic twinning is very high.

It is generally assumed that conjoined twins derive from a single blastocyst and represent a form of monozygotic twinning rather than the fusion of dizygotic twins. All studies to date have indicated that conjoined twins are of the same sex, giving strong support to the theory that they are of monozygous derivation.

Through a review of the available literature on previous cases of epigastric heteropagus twins, we found several interesting and differing characteristics relating to epigastric heteropagus cases (Table 1).

1. Conjoined twins are predominantly female, and the cause of the female preponderance is not known.

2. Hereditary tendency is characteristic of epigastric heteropagus twins. In case 3 (Table 1), the mother has a twin and the father has twin cousins.

Fig 1. The parasitic twin was attached to the host's epigastrium, above an omphalocele. The parasite had a pelvis with two well-formed legs, a penis, and a scrotum. There was no anal opening in the parasite.

Fig 2. Plain x-ray shows the accessory bony pelvis and well-developed lower limb bones of the parasite. There was no bony connection between the autosite and the parasite. The parasite did not have a vertebral column.
The patient is a Nigerian black. This racial population has a higher frequency of twins; they account for 5% of all births. In case 7, the patient has older sisters who are twins. In the present case, the mother’s first child had syndactyly, which is a manifestation of the doubling anomaly.

<table>
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<tr>
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<th>Reference No.</th>
<th>Year</th>
<th>Sex</th>
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<th>Vascular Connection</th>
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Abbreviations: FHx, family history of conjoined twins; Me, Meckel's diverticulum.
AFMP

Fig 5. Result of haplotyping for tetranucleotide repeats polymorphism at the D21S11 locus of the patient's family shows the monozygotic origin of this heteropagus condition. A, father (F); M, mother (M); P, parasite.

3. There has been only one case of bowel connection between the autosite and the parasite (case 4); the parasite's bowel was connected to a Meckel's diverticulum of the autosite.

4. No reported case of epigastric heteropagus twins has involved a bony connection between the autosite and the parasite.

5. No reported case of epigastric heteropagus twins has involved a vascular connection between the autosite and the parasite.

In light of the above observations, we initially hypothesized that our epigastric heteropagus case was a dizygotic twin in which a body portion of an embryo fused parasitically to a complete embryo. Previous cytogenetic studies showed an identical, normal karyotype in symmetrically conjoined twins. However, an identical karyotype is not enough to confirm the monozygotic origin of conjoined twins. Therefore we performed karyotyping and a DNA study. Haplotyping in our case showed a monozygotic origin.

The female zygote is more likely to undergo conjoined twinning than the male, but once formed, male conjoined twins are much less viable and often are aborted. Among reports of stillborn fetuses of conjoined twins, male gender is predominant. A heteropagus twin could develop when one component is better placed such that it monopolizes the placental blood to the detriment of the other member (parasite). Epigastric heteropagus twins may result from ischemic atrophy of one part of a monozygotic conjoined twin early in gestational life. This case was the first in which the monozygotic origin of conjoined twins was confirmed by a DNA study. We hope that, in future studies, the surgical specimens of conjoined twins will be analyzed to further confirm the monozygotic origin.

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REFERENCES