Sirenomelia: A Case Report of a Rare Congenital Anomaly and Review of Literature

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ABSTRACT

Sirenomelia, or the Mermaid Syndrome, is a very rare congenital anomaly. It is associated with varying degrees of fusion of lower limb bones, giving them the appearance of a “mermaid.” It is almost always associated with other birth defects, such as, renal abnormalities, genital anomalies, and cardiac anomalies. We report a case of sirenomelia associated with bilateral multicystic dysplastic kidneys along with bilateral hydronephrosis, severe oligohydramnios, single umbilical artery, absent anal opening, and absent genitals. To the best of our knowledge, this is the first case to be reported from our region.

Keywords: Congenital anomalies, Mermaid Syndrome, Sirenomelia.


Source of support: Nil

Conflict of interest: None

INTRODUCTION

Sirenomelia is an extremely rare and invariably fatal congenital anomaly, with an incidence of approximately 1 in 100,000 live births. Most of the cases are associated with stillbirth. Only 1% babies survive the first week of life. It is more common in monozygotic twins than in single births or fraternal twins. The incidence of male: female affection is 2.7:1. The most characteristic feature is the partial to complete fusion of lower limbs, resembling a mermaid. Associated anomalies make it difficult for the babies to survive; however, there are a few reports of children surviving with the help of operative procedures and treatment.

CASE REPORT

A 25-year-old female, G3P1L1A1, 33 weeks gestation by date, was brought to our hospital for complete placenta previa with severe oligohydramnios. She was not diabetic and had no other chronic pathologies. There was no history of any drug ingestion or any substance abuse. Antenatal USG in the second trimester was suggestive of severe oligohydramnios. A repeat USG in the third trimester showed complete placenta previa, with bilateral multicystic dysplastic kidneys, significant shortening of limbs with a foot not distinctly seen, and a complete cord around the neck. The patient was taken for LSCS in view of placenta praevia with severe oligohydramnios. A 1.103 kg baby was delivered through LSCS. Apgar score was 2 at 1 and 5 minutes. Resuscitation was given, but the child could not be revived and expired within an hour of birth. The parents were explained regarding the nature of the disease and death, and the baby was handed over to the parents. Postmortem examination was not done.

On examination, the baby was found to have multiple anomalies. There was complete fusion of both the lower limbs with a single foot and fused toes, absent genitals, absent urinary meatus, absent anal opening, single umbilical artery, and a vestigial tail-like structure at the sacrococcygeal area Figs 1 to 3. Other associated...
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syndromic features were microcephaly, hypotelorism, depressed nasal bridge, and low set ears.

DISCUSSION

Sirenomelia is a rare congenital anomaly involving fusion of varying degrees of the caudal part of the body. Other abnormalities like those of renal, cardiac, pulmonary, gastrointestinal, genital, and nervous system abnormalities are invariably associated with it. The survival depends on the degree of associated anomalies and adequate functioning. We could not ascertain the complete nature of anomalous abnormalities in our child given that post-mortem examination was not performed and no imaging was done postnatally.

ETIOLOGY

There is no single defined etiology, and various theories have been put forth.

Stevenson et al\textsuperscript{11} proposed a vascular steal phenomenon as the cause of defective development of the lower limbs. According to this theory, there is abnormal perfusion of the caudal portion of the body due to shunting of blood via an abnormal abdominal artery. Due to this, there is inadequate nutrition supply to the caudal part, leaving it susceptible to maldevelopment. Hence, all the structures present caudally are abnormally developed.

Another theory suggests that there is a defect in the later stages of gastrulation during the third week of gestation, which leads to defective development of the caudal parts of the body to varying degrees. This is referred to as the blastogenesis theory. According to this theory, sirenomelia is considered as an extreme form of caudal regression syndrome.\textsuperscript{12-15} However, recent views negate this, and sirenomelia is considered to be a different entity.

Advances in genetics have put forth a genetic possibility for this disease. According to a study by Garrido-Allepuz et al\textsuperscript{16} mice lacking Cyp26a1, an enzyme degrading retinoic acid, and reduced bone morphogenetic protein (Bmp) may lead to sirenomelia.

Teratogens, such as, diabetes, tobacco exposure, or drug abuse, also have been postulated to be a cause of sirenomelia.

CLASSIFICATION

Stocker and Heifetz\textsuperscript{17} have classified the entity into seven types as follows (Table 1):

<table>
<thead>
<tr>
<th>Type</th>
<th>Characteristic</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>All thigh and leg bones are present</td>
</tr>
<tr>
<td>II</td>
<td>Single fibula</td>
</tr>
<tr>
<td>III</td>
<td>Absent fibula</td>
</tr>
<tr>
<td>IV</td>
<td>Partially fused femurs, fused fibulae</td>
</tr>
<tr>
<td>V</td>
<td>Partially fused femurs</td>
</tr>
<tr>
<td>VI</td>
<td>Single femur, single tibia</td>
</tr>
<tr>
<td>VII</td>
<td>Single femur, absent tibia</td>
</tr>
</tbody>
</table>

Table 1: Classification by Stocker and Heifetz

![Fig. 3: Vestigial tail-like structure with absent anal opening](image)
In our case, no specific classification could be defined as radiographs were not taken.

DIAGNOSIS

Antenatal USGs are of utmost help in determining the presence of this disease. It may detect abnormal caudal development, such as, absent femurs, tibia or fibula, converging bones, abnormal renal development, such as, renal agenesis, hydronephrosis, absent bladder, and oligohydramnios, which may give a clue to the diagnosis. However, oligohydramnios may restrict the complete evaluation of the fetus. Since there is absence of any open anomaly, alpha fetoprotein evaluation is of seldom help. Previous affected conception may also help in diagnosing the disease. Post-delivery, clinical evaluation is useful for diagnosing this entity. Supportive evidence in the form of ultrasonography and X-rays may also help in diagnosing the extent of associated anomalies. A postmortem examination can also help in determining the nature and severity of the anomalies.

MANAGEMENT AND PROGNOSIS

Sirenomelia carries a very poor prognosis. The affected babies rarely survive beyond 5 days. Survival depends upon the severity of associated anomalies. If diagnosed early in the antenatal period, medical termination of pregnancy is advised depending upon the gestational age, extent of anomalies, and approval of parents. However, there are reports of a few babies surviving with the help of surgeries. However, financial constraints, the availability of expertise for treatment, familial motivation, and quality of life post the procedure are issues.

CONCLUSION

Sirenomelia is a rare fatal congenital anomaly associated with varying degrees of visceral defects. The etiology is still a matter of controversy and is not clearly defined. Antenatal diagnosis is possible with the help of radiology. A regular antenatal checkup, hence, is essential for all pregnant females.

REFERENCES