Successful Expectant Management of the Anomalous Fetus with Sirenomelia in Twin Pregnancy: A Case Report and Literature Review

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Background: Sirenomelia is a rare congenital defect. Its management during pregnancy and after delivery is becoming a controversial issue because of its complex nature and management outcome. The possibility of expectant management in the sirenomelia twin drove us to write this case report.

Case Presentation: We report a case of successful expectant management in twin sirenomelia which was diagnosed in the second trimester. Prenatal counseling of the couple by a multi-disciplinary team regarding the diagnosis, treatment, and prognosis of the sirenomelia twin was done. The mother gave birth, at term, to one normal and one sirenomelia neonate by cesarean section.

Conclusion: Expectant management of sirenomelia one in twin pregnancy is advisable in a resource-limited setting.

Keywords: sirenomelia, twin pregnancy, expectant management

Introduction

Sirenomelia, which is also known as mermaid syndrome, is a multisystem congenital anomaly characterized by a partial or complete fusion of the lower limbs into a single lower limb, giving the appearance of a mermaid’s tail. 1-3 It is associated with the gastrointestinal tract (anorectal, esophageal), genitourinary tract (renal, genitalia, lower urinary tract), cardiovascular system, respiratory tract, musculoskeletal system, and central nervous system anomalies. 2,3

Sirenomelia is a very rare congenital malformation with an incidence of 0.98–4.2 per 100,000 live births. 3-5 Its incidence of twin pregnancy is extremely low. However, compared with a singleton pregnancy and dizygotic twins, the incidence of sirenomelia in the monozygotic twin is more than 100-fold. 2 Therefore, the diagnosis and management of sirenomelia in twin pregnancy is a worthy matter to be discussed. The exact etiology of sirenomelia is unclear. 4 Different works of literature associate it with extreme maternal age, 3 maternal diabetes, prenatal exposure to retinoic acid, cadmium, cyclophosphamide, 4 cocaine, landfills water, or lamotrigine. 6 Here we present successful expectant management of sirenomelia one in twin pregnancy which was diagnosed in the second trimester.

Case Presentation

This is a 20-year-old woman from Western Ethiopia who was married to a non-consanguineous –24–year–old man. She is a farmer and of low socioeconomic
status. She is a primigravida mother who does not remem-
ber her last normal menstrual period but claims to be
amenorrheic for the last 09 months. She presented to
Arjo primary hospital with the complaint of labor pain of
3 hours. There was no passage of liquor. She had no self or
family history of chronic medical problems like hypertension,
diabetes Mellitus, or bronchial asthma. She had no history of smoking, chewing khat, alcohol, or other drug
abuse. Folic acid supplementation was not given for the
mother during and/or before the pregnancy. She has no
history of exposure to hazardous chemicals. There was no
history of twinning or congenital anomaly in the family.
She had four visits of prenatal care follow up at Arjo
primary hospital. During the prenatal care visit, physical
examination, and essential laboratory tests including blood
group, blood glucose, hemoglobin, urinalysis, and serology
for human immunodeficiency virus (HIV), hepatitis
and syphilis were done. The laboratory results showed
hemoglobin = 13g/dl, blood sugar= 106mg/dl, hepatitis
B surface antigen (HBsAg) = negative, Venereal Disease
Research Laboratory Test (VDRL) = nonreactive, HIV
rapid test = negative and A negative blood group.
Second-trimester ultrasound examination, done at 3rd
antenatal care visit, revealed diaminotic twin intrauterine
pregnancy (one sirenomelia twin). The first twin was in
cephalic presentation with an average gestational age of 27
weeks while the second twin was in breech presentation
with an average gestational age of 26 weeks. There was
only one leg bone in the second twin fetus. There was
oligohydramnios in 2nd twin fetus. Placenta was fundal
and a dividing membrane was seen.
The 4th antenatal care visit was done one month after
the 3rd visit. It showed the normal fetus in good condition
but oligohydramnios in anomalous sirenomelia twin. The
couple was counseled by the multidisciplinary team
regarding diagnosis, treatment, prognosis, and possible
complications of sirenomelia. Finally, expectant manage-
ment was decided and the mother was appointed to come
after 4 weeks. Unfortunately, she came to the hospital in
labor.
Clinical examination during this time showed that she
is in labor pain. Her vital signs were blood pressure=
100/60 mmHg Pulse rate=86 beats per minute, respirator
rate=22 breaths per minute, and temperature=36.2°C. The
conjunctiva was pink. Abdominal examination showed
term-sized gravid uterus, multiple fetal poles felt, fetal
heart best heart at two places (143 and 130 beats
per minute), 3 contractions in ten minutes each staying
30 to 40 seconds. On pelvic examination cervix was 6cm
dilated, 100%effaced, vertex presentation, the station at
−1, membrane intact, and adequate pelvis. Intrapartum
ultrasound examination showed the first twin is a
cephalic presentation and the second twin in breech
presentation. Both fetuses were alive. With the impres-
sion of an active phase of labor and twin pregnancy, she
was admitted to the labor ward. After 4 hours, she was
reevaluated and the cervix was still 6cm. Thus, with the
final diagnosis of the arrest of cervical dilatation cesar-
ean section was done under spinal anesthesia to effect
delivery of first twin male alive neonate weighing
2500gms with APGAR scores of 8 and 9 at 1st and
5th minute respectively, second twin sirenomelic neonate
weighing 1800gms with APGAR scores of 7 and 9 at 1st
and 5th minute respectively. The single placenta was
delivered by cord traction. It is a monochorionic dia-
mniotic placenta. There were no identified gross placen-
tal and umbilical cord pathologies. Detailed physical
examination of the second twin showed the absence of
external genital organs, fused leg bones (both femur and
tibia), short neck (Figure 1), and imperforate anus. This
is a rudimentary limb, fused leg bones, no mobility, and
on foot. Though the prognosis is poor, there was an
attempt to refer the newborn to a better setting but the
families refused the medical advice. The newborn was
active and feeding well for the first two days of delivery.
On the third day, it got sick and passed away.
On the fourth operative day, the mother had a smooth
postoperative course and her first neonate was normal. They
were discharged with advice about postpartum care follow-up.

Figure 1 Anomalous fetus with sirenomelia in twin delivery at Arjo Primary
Hospital, 2020, Western Ethiopia.
Discussion

Sirenomelia is a severe congenital malformation involving multiple organ systems.\textsuperscript{1–3,7–9} The fusion of the lower extremities, called symelia, is the pathognomonic finding in sirenomelia.\textsuperscript{9,10} The visceral anomalies occur uniformly and significantly affect survival.\textsuperscript{9,10} Gross examinations of our case showed completely fused lower limbs, absence of external genitalia, and imperforated anus. But we did not have an abdominal ultrasound examination and x-ray of the sirenomelia neonate to comment on visceral anomaly and detail skeletal defect.

The etiology and pathogenesis of sirenomelia remain unknown.\textsuperscript{1,10} However, there are two proposed hypotheses. These are the defective blastogenesis hypothesis and the vascular steal hypothesis.\textsuperscript{9} Experimental analysis in mice shows genetic origin in sirenomelia.\textsuperscript{9} But in humans, sirenomelia is still considered sporadic.\textsuperscript{1} In our case, there is no risk factor and chronic medical condition detected. But the mother was not supplemented with folic acid during and before pregnancy.

Though sirenomelia is a very rare condition,\textsuperscript{1} its incidence in twin pregnancies is more common when compared to singleton pregnancies.\textsuperscript{2} There are approximately 300 cases reported in the literature, 15% of which are associated with twinning, most often monozygotic.\textsuperscript{7,11} In twin pregnancies, in addition to the above-mentioned mechanisms, mechanical defects resulting from lateral compression by amniotic folds can end up in sirenomelia.\textsuperscript{3} Therefore, the diagnosis and management of sirenomelia in twin pregnancy need attention.

Early diagnosis of sirenomelia is very crucial for timely prenatal counseling and decision of its management. High-resolution or color Doppler ultrasound during prenatal care can safely make the diagnosis of sirenomelia.\textsuperscript{12} During the first trimester, the amniotic fluid volume is usually normal, unrelated to fetal urine production. In the second and third trimesters, oligohydramnios from renal agenesis is a common finding in sirenomelia.\textsuperscript{13} This makes ultrasound examination difficult. Therefore, it advisable if to have an ultrasound examination in the late first and early second trimester.\textsuperscript{1} In our case, the diagnosis was made in the late second trimester.

In the presence of oligohydramnios and after 20 weeks of pregnancy, fetal Magnetic Resonance Imaging (MRI) is the preferred option. It demonstrates various anomalies in greater detail than a fetal sonogram.\textsuperscript{12} Sirenomelia is a fatal malformation.\textsuperscript{1} About 50% of cases are stillborn because of bilateral renal agenesis and associated visceral anomalies.\textsuperscript{11} Therefore, early diagnosis and decision are important. In singleton pregnancies, the decision to terminate or continue the pregnancy depends on factors like the extent of visceral anomalies, family-related issues, and the presence of a multidisciplinary medical team.\textsuperscript{10,14,15} The prognosis worsens especially in association with VACTERL and VACTERL-H syndrome.\textsuperscript{16}

In a twin pregnancy, expectant management is possible as far as there is no unique complication of twin pregnancies and other obstetric problems necessitating termination of pregnancy.\textsuperscript{13} In a better setting and when the diagnosis is made in early pregnancy, voluntary selective termination of sirenomelia one in twin pregnancy may be advised.\textsuperscript{2} Our case was diagnosed in the late 2nd trimester of pregnancy. After having appropriate prenatal counseling, expectant management was decided. Prenatal care was successful. The mother delivered by cesarean section at term.

Sirenomelia neonates usually die within a few days of delivery because of associated anomalies and poor medical care.\textsuperscript{1,2,10} In our case, the neonate died on the 3rd day of delivery. However, because of advanced medical care and multidisciplinary approach, sirenomelia is no longer considered a lethal condition.\textsuperscript{14} Survival depends on visceral associated anomalies, particularly kidneys, rather than sirenomelia itself.\textsuperscript{10} So far, there are six cases of sirenomelia that survived by multidisciplinary treatment.\textsuperscript{14,17} In our case, though an effort was made to refer the neonate to a better setting, it was not possible. The family ignored the neonate due to cultural and religious reasons. Thus, it did not have any imaging, genetic testing, or a post mortem examination.

Abbreviations

HIV, human immunodeficiency virus; HBsAg, hepatitis B surface antigen; MRI, magnetic resonance imaging; VDRL, Venereal Disease Research Laboratory Test.

Data Sharing Statement

The datasets used during the current study are available from the corresponding author on reasonable request.

Ethics Approval and Consent to Participate

The Ethics Committee of Wollega University has approved the publication of this case.
Consent for Publication
Written informed consent was obtained from the couple for publication of this case report and any accompanying images. A copy of the written consent is available for review by the Editor-in-Chief of this journal.

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We thank the couple for allowing the publication of this case report.

Author Contributions
All authors made a significant contribution to the work reported, whether that is in the conception, study design, execution, acquisition of data, analysis, and interpretation, or in all these areas; took part in drafting, revising, or critically reviewing the article; gave final approval of the version to be published; have agreed on the journal to which the article has been submitted; and agree to be accountable for all aspects of the work.

Disclosure
The authors report no competing interests in this work.

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