Infantile Bowel Obstruction in a Patient with Situs Inversus Totalis and Polysplenia: A Case Report

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Abstract: The infantile intestinal obstruction associated with situs inversus totalis and polysplenia is extremely rare, with only a few cases reported in the literature. Furthermore, the management of this association is complicated. We report a case of a 2-month-old boy with intestinal obstruction due to malrotation and volvulus with thin translucent omentum sac encasing the small intestine associated with situs inversus totalis, polysplenia, and pulmonary hypertension. To the best of our knowledge, this is the first case of situs inversus totalis with polysplenia, pulmonary hypertension, and intestinal obstruction due to malrotation and volvulus with thin translucent omentum sac encasing the small intestine.

Keywords: situs inversus, polysplenia, dextrocardia, malrotation, midgut volvulus

Introduction

Situs inversus is a rare condition in which body organs are reversed from their normal position.1

Situs inversus totalis is a rare genetic abnormality that has been estimated to happen 1 in 10,000 to 1 in 50,000. In situ inversus totalis rotation is clockwise, but the normal embryonic rotation of the midgut is counter-clockwise direction.6

The abdominal-thoracic viscera are mirrored images and are transposed in situs inversus totalis.2

It can be isolated or associated with other congenital abnormalities, the most common of which is intestinal malrotation, which affects 40–90% of patients.3

On the other hand, polysplenia is identified by the existence of multiple spleens. In approximately 20% of cases, polysplenia is associated with situs inversus.1

We report here a case of situs inversus totalis, polysplenia, and pulmonary hypertension in which intestinal obstruction due to malrotation and volvulus with thin translucent omentum sac encasing the small intestine was the initial presentation.

Case Presentation

A 2-month-old boy was referred to our pediatric surgery department with bilious vomiting and mild abdominal distention since birth.

The baby was born at term by cesarean section to a 25-year-old primigravida mother. On examination, the infant was afebrile and moderately dehydrated.

The abdomen was soft and distended, and the nasogastric aspirate was bilious. Plain radiographs revealed dextrocardia, liver opacity on the left side of his abdomen, and a stomach bubble on the right side suggestive of situs inversus totalis (Figure 1).

In addition, there were a few dilated bowel loops in his upper abdomen with an absence of distal gas shadows. Echocardiography demonstrated dextrocardia and pulmonary hypertension with a systolic pulmonary arterial pressure of 54 mmHg.
The patient was stabilized and resuscitated. After optimization, an abdominal exploration confirmed the situs inversus totalis with the liver and a C-loop of the duodenum on the left side. During the exploration, we also detected that our patient had twelve spleens of different sizes (Figure 2).

The entire small bowel was healthy but enclosed in a thin transparent sac containing jelly-like material in the left half of the abdomen (Figure 3).

Several Ladd's bands crossed the second part of the duodenum with a narrow-base mesentery.

The Ladd’s procedure was performed; the bowel was derotated after excising the thick Ladd’s bands. Subsequently, the sac was excised, and an inversion appendectomy was performed.

The child was referred to the newborn intensive care unit in an intubated state, but the next day, the child was extubated.

The immediate postoperative period was uneventful, and oral feeding started on day five after passing stool. Subsequently, hematemesis and rectal bleeding started for an unknown reason on the eighth postoperative day, and the patient died.
Situs inversus totalis (SIT) is a rare autosomal recessive congenital anomaly in which the abdominal organs are fully transposed. Therefore, there is a mirror-image transposition of all abdominal organs. The first known case in humans was reported by Fabricius. Situs inversus is estimated to occur in 1 in every 8000 to 25,000 live births. The condition is equally distributed among males and females, while boys predominate when there are concurrent splenic anomalies, similarly to the present case.

Polysplenia syndrome is characterized by the presence of two or more spleens, usually detected during childhood. It can be associated with congenital anomalies of the other organs. The position of spleens is variable, and their number can vary from 2 to 16. Approximately 20% of these cases have situs inversus, while more than 40% have cardiac anomalies.

Our patient had situs inversus totalis with dextrocardia and polysplenia; however, we did not detect any cardiac anomalies.

The midgut’s embryonic rotation is usually counterclockwise, but in situs inversus totalis, the rotation is clockwise. Due to this erroneous rotation, the visceral organs are inverted like a mirror image. In almost all cases, situs inversus totalis is diagnosed during a routine check-up or while performing imaging studies for another illness.

Abdominal and chest X-rays, computerized tomography (CT) scans, and barium X-rays can all be used to diagnose this condition. Our case was diagnosed after the referral for an erect abdominal x-ray and an upper gastrointestinal contrast study.

Differential diagnosis includes multiple organ malrotation syndrome. Intussusception is commonly present at age 6 and 36 months, and the male-to-female ratio is approximately 3:1. The most common type is ileocolic and others include ileo-ileal, jejuno-jejunal, jejuno-ileal, or colo-colic. The case has excluded intussusception according to age, radiologically and intra operative finding as reported (Mulita F, Tchabashvili L, Liolis E, Maroulis I).

Surgical management of neonatal intestinal obstruction associated with situs inversus totalis should follow the same surgical principles as performed for classical cases.

In addition, the direction of rotation should be examined since transposition of the viscera can lead to midgut volvulus. Ladd’s procedure is the standard corrective surgery for intestinal malrotation in children. Notably, Ladd’s procedure involves division of the peritoneal bands (ie, Ladd’s bands) traversing the posterior abdomen, reduction of volvulus, appendectomy, and functional positioning of the intestine with or without fixation.

To the best of our knowledge, there is only one published case of a thin translucent membrane sac encasing the small intestine in the literature. Mahalik et al presented a 2-year-old boy with situs inversus totalis, polysplenia, ventricular septal defect, and malrotation with volvulus. These authors reported that they removed the sac and thick Ladd’s bands, derotated the bowel and performed an inverted appendectomy.
This child recovered uneventfully and was discharged on the sixth postoperative day. During a two-year follow-up, he was asymptomatic and thriving well.

In our case, although a thin translucent omental sac enveloped the small intestine, the entire small bowel was healthy, and there were no cardiac anomalies.

Subsequently, the patient underwent Ladd’s procedure due to intestinal malrotation with volvulus. Unfortunately, on the eighth postoperative day, hematemesis and rectal hemorrhage began for an unclear reason, and the patient died.

To the best of our knowledge, this is the first case of situs inversus totalis with polysplenia, pulmonary hypertension, and intestinal obstruction due to malrotation and volvulus with thin translucent omentum sac encasing the small intestine.

**Conclusion**
All newborns should undergo a physical examination in the delivery room for the early detection of situs inversus totalis, and the possibility of encountering life-threatening complications should always be kept in mind.

In situs inversus, due to the transposition of viscera, midgut volvulus may occur in an anticlockwise direction, hence derotation is carried out clockwise.

**Ethics Approval**
Based on the regulations of the review board of the Mogadishu Somali Turkish Training and Research Hospital, it is not required institutional review board approval for the case reports.

**Consent for Publication**
Written informed consent had obtained by the patient’s parents to have the case details and any accompanying images published.

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**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.

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The authors declare no conflicts of interest in this work.

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