Bochdalek Hernia and Partial Diaphragmatic Agenesis: Pedicled Intercostal Muscle Flap and Mesh Repair in a Young Adult with Sickle Cell Disease

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Abstract

Keywords

► congenital diaphragmatic hernia
► hemidiaphragm agenesis
► sickle cell disease
► intercostal muscle flap
► ectopic kidney
► vertebral fusion

Congenital Bochdalek hernia (BH) in an adult is rare and has an unusual presentation. They are confined to the pediatric age group with an incidence of 1:3,000 live births. It rarely persists asymptomatic until adulthood. Surgical repair by thoracic, abdominal, or thoraco-abdominal approach is the treatment of choice with diaphragmatic reconstruction in associated diaphragmatic agenesis. With only 10 cases of BH with partial diaphragmatic agenesis reported to date, we discuss the rarity, unusual presentation, and management of BH in a young adult with sickle cell disease that has not been reported in the literature.

Case Report

A 24-year-old female with a sickle cell disease presented with mild chest pain, an episode of vomiting, and no history of trauma or chronic illness; she was hemodynamically stable, afebrile with regular abdomen examination. Chest examination revealed decreased left air entry, chest X-ray showed left consolidation, and ultrasound thorax confirmed the presence of bowel loops and left kidney within the thoracic cavity. Contrast-enhanced computed tomography (CECT) of the thorax and abdomen showed colon, small bowel, left kidney in the thoracic cavity with scoliosis (Cobb angle of 24 degrees), fused 12th thoracic and 1st lumbar vertebra. Ultrasound of abdomen and kidneys revealed an ectopic left kidney. CT abdomen with contrast demonstrated absence of right kidney. Ultrasound of abdomen and chest confirmed the presence of left kidney, colon, and small bowel within the thoracic cavity. Chest CT showed a persistent right pleuroperitoneal canal. Thoracic CT showed hemidiaphragmatic agenesis on the left side and a persistent pleuroperitoneal canal. The patient underwent a combined thoracoabdominal approach for repair of the diaphragmatic defect and placement of mesh. The diaphragmatic defect was repaired using pedicled intercostal muscle flap and mesh. The patient had an uneventful postoperative course and was discharged on the 7th postoperative day.
lumbar vertebral bodies, and displaced left diaphragmatic crus and its defect (►Fig. 1A, 1B).

The patient received preoperative antibiotics, nebulization, and bowel preparation with normal blood investigation. Following left thoracotomy, the thoracic cavity was accessed through the fifth intercostal space, which contained a hypoplastic lung, the colon, small intestine up to the duodenal jejunal flexure, and left kidney with no hernia sac. The normal colonic mesentery was adherent to the aorta and pericardium (►Fig. 2A). Following vascular adhesiolysis and laparotomy, intrathoracic contents were transpositioned to normal anatomical positions. Primary partial diaphragmatic repair, pedicled intercostal muscle flap raised between the sixth and seventh ribs, followed by partial excision of the sixth rib obliterated posterolateral defect. Polypropylene mesh was placed over the primary repair and around the flap with Prolene 3-0 to provide diaphragmatic stability (►Fig. 2B). A single drain was placed in the chest and abdomen, followed by a closure. Mild splenomegaly was not an indication for splenectomy. Extubated patient was maintained nil by mouth for 24 hours. She received one unit (350 mL) of packed red blood cells and Ryles tube feeding with fluids on the first and second postoperative days followed by oral feeds and a soft diet from the third and fourth days. The postoperative course was uneventful, with chest and abdomen drains removed on the eighth and ninth days. CECT thorax and abdomen repeated after 2 weeks revealed minimal pleural effusion with an intact diaphragm that resolved with subsequent follow-up visits and no recurrence (►Fig. 3) with CECT obtained at the third and sixth months.

Discussion

Czech anatomist Vincent Alexander Bochdalek first described congenital adult diaphragmatic hernia in 1848.1 Due to varied clinical presentations, diagnosis is late.4 Volvulus, strangulation, organ perforation, and tension pneumothorax require emergency intervention.4 Contents include omentum (92%), splenic flexure of the colon (58%), stomach (25%), spleen and liver.3

Displaced diaphragmatic crus is explained by the maldevelopment of at least three of the four parts of the hemidiaphragm anlage, the unpaired ventral portion from the septum transversum, the unpaired dorsolateral portions from the pleuroperitoneal membranes, or the medial portion from persisting parts of the primary mesentery.2 Associated ectopic intrathoracic kidney accounts for 5% with a prevalence of <0.01% and the presence of vertebral fusion has been explained as a complex synchronous metameric defect of somite and intermediate mesoderm.1,6

Surgical repair is the treatment of choice, either by thoracoscopic/laparoscopic approach or by an open repair. We performed an open thoraco-abdominal approach as
repositioning of intrathoracic content into the abdominal cavity was not possible through the defect as mesentery of the colon was adherent to the descending thoracic aorta and pericardium as well as dense adhesions confined to the abdominal wall restricted manual reduction. Acute splenic sequestration crises are a complication of sickle cell disease. However, as there is a lack of evidence from the randomized control trials showing that splenectomy improves survival and decreases morbidity in patients with sickle cell disease, hence splenectomy was not considered.

Our case is unique in presentation. First, its association with intrathoracic ectopic kidney has been reported in less than 0.25% of the cases and further association of intrathoracic ectopic kidney with vertebral fusion has been reported only once in the literature. Second, with only 10 cases of partial diaphragmatic agenesis in adults reported to date, this case adds associated rarity in the form of asymptomatic presentation, absent sac with the adhesion of mesentry to the thoracic aorta, pericardium, and reconstruction with pedicled intercostal muscle flap in the patient with sickle cell disease, which has not been reported in the literature.

Conclusion

Adult congenital BH is rare, can have unusual presentation, and perioperative management in sickle cell disease is challenging, requiring multimodal management.

Author Contribution

Klein Dantis prepared and drafted the manuscript. All the coauthors edited, reviewed, and accepted the manuscript.

Conflict of Interest

None.

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Fig. 3 Postoperative contrast-enhanced computed tomography showing neo-diaphragm with minimal pleural effusion and no recurrence at 14 days.