Case Report

Strangulated Morgagni’s Hernia: A Rare Diagnosis and Management

Malav Modi, Amit Kumar Dey, Ajay Mate, and Samir Rege

Department of Surgery, Seth G.S. Medical College and KEM Hospital, Mumbai, India

Correspondence should be addressed to Amit Kumar Dey; amit5kem@gmail.com

Received 9 July 2016; Accepted 19 September 2016

Academic Editor: Boris Kirshtein

Copyright © 2016 Malav Modi et al. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Morgagni hernia is a rare type of congenital diaphragmatic hernia. It accounts for only 3% of all diaphragmatic hernias [1]. Though some are symptomatic, many remain asymptomatic for a long time and are discovered on X-ray incidentally [2]. Strangulated Morgagni’s hernia is very uncommon and only five such cases are reported in literature [4–6]. This paper is written with the objective of reporting a new case of strangulated Morgagni’s hernia along with its morphology and treatment in detail.

1. Introduction

Morgagni’s hernia is the herniation of the intra-abdominal organs through a congenital defect in the diaphragm immediately behind the sternum [1]. Its cases are very rare and make up 2–3% of all cases among the four types of congenital diaphragmatic hernias [2]. Though some are symptomatic, many remain asymptomatic for a long time and are discovered on X-ray incidentally [3]. Strangulated Morgagni’s hernia is very uncommon and only five such cases are reported in literature [4–6]. This paper is written with the objective of reporting a new case of strangulated Morgagni’s hernia along with its morphology and treatment in detail.

2. Case Presentation

A 40-year-old male patient presented to our emergency department with acute severe pain and distension of abdomen. Severe abdominal pain was associated with absolute constipation and multiple episodes of vomiting that started 2 days ago. He also complained of increasing breathlessness since then. Patient was a chronic alcoholic and an occasional smoker. There was no prior history of similar complaints in the past. On examination, patient was normotensive with persistent tachycardia and tachypnoea. Abdomen was distended, tender, guarded with no signs of peristalsis on auscultation. Air entry was reduced on right lower zone of chest on auscultation. Chest radiograph showed an elevated right hemidiaphragm (Figure 1). Scout film of the abdomen revealed multiple air fluid levels with elevated right hemidiaphragm. Ultrasonography of the abdomen was indicative of dilated small bowel loops with ascites with right pleural effusion. Computed tomogram (CT) of the chest and abdomen performed was suggestive of herniation of jejunoileal loops through a right diaphragmatic defect (Morgagni’s hernia) with ascites and right pleural effusion (Figure 2). After initial resuscitation, patient was explored through a midline laparotomy incision. There was a defect in the diaphragm on the right side anteriorly 10 × 6 cm in diameter between sternal and costal attachments of diaphragm with herniation of jejunoileal loops (Figures 3 and 4). Jejunoileal loops of around 50 cm were gangrenous and aperistaltic with poor turgor. The sac also contained about 400 mL of dark haemorrhagic fluid. The right lung was hypoplastic. Hernial contents were reduced and the ischemic segment of bowel was resected followed by an anastomosis. Peritoneal sac was dissected. Right intercostal drain was placed and pleura was sutured. The diaphragmatic defect was closed with an Ethibond 2-0. Postoperatively, patient was on ventilator support for 3 days. He was gradually weaned and extubated. Intercostal drain
Failure of fusion of the fibrotendinous elements of diaphragm, that is, sternal and costal attachments, leaves behind a muscle-free area known as the costosternal trigone or the space of Larrey or Morgagni’s foramen through which the hernia occurs eventually [1]. Because of extensive pericardial attachment on the left which does not allow herniation, the hernia is more common on the right side and is situated anteriorly [7]. Almost 90% of Morgagni’s hernias are reported to be on the right side, with 2% located on the left and 8% bilateral [8]. The sac lies between the pericardium and the right pleura. Comer and Clagett reported a series of 1135 cases of diaphragmatic hernia at Mayo’s clinic, of which only 50 were of Morgagni’s type with only one obstructed case [2]. Usually, the hernia sac contains the transverse colon followed by stomach, omentum, and small intestine but occasionally the liver may also protrude into the sac [9]. Diagnosis of Morgagni’s hernias is usually late because patients can be asymptomatic or present with vague respiratory (cough, expectoration, and dyspnea) or gastrointestinal (nausea, vomiting, subcostal pain, pain after food, or rarely acute intestinal obstruction) symptoms and signs [10]. Diagnosis is usually done by conventional radiography with occasionally missed cases. Computed tomography scan can be useful in diagnosing the contents of the hernia sac and is noninvasive and accurate. Magnetic resonance imaging can distinguish Morgagni’s hernia from other mediastinal masses and is noninvasive too [8]. Pleuropericardial cyst, lipoma, intrathoracic tumours, and eventration of diaphragm should be differentiated from Morgagni’s hernia while making a diagnosis [9]. Barium studies could be useful in supplementary investigation. In our case, diagnosis was made by physical examination and plain chest radiogram and confirmed by computed tomography. To avoid the risk of strangulation, some recommend repair even in asymptomatic patients [5] whereas some advocate conservative approach because Morgagni’s hernia remains asymptomatic for a long time [11].

A subxiphoid preperitoneal approach has the benefit of a small incision [9]. In cases of certain diagnosis, abdominal approach (open or laparoscopic) is preferred over thoracic approach for surgery because of easier reduction of the hernia and because abdominal viscera within the hernia can be easily pulled down to their normal location [8]. In cases of unclear diagnosis, the recent trend is towards laparoscopy which has...
low morbidity [6]. Laparoscopy can be therapeutic, as well as diagnostic [12]. In our case, open abdominal approach was preferred because of ischemic bowel and general condition of patient. In conclusion, Morgagni’s hernia is a rare type of congenital diaphragmatic hernia and is more common on right side.

4. Conclusion

Most of the patients are asymptomatic and present late usually with complications. Diagnosis in suspected case can be confirmed by computed tomography scan. Surgery by abdominal approach (open or laparoscopic) is management of choice.

Consent

Written informed consent was obtained from the patient for publication of this paper and any accompanying images.

Competing Interests

All authors declare no conflict of interests.

Authors’ Contributions

Each of the authors was involved in preparation of the manuscript: Malav Modi, Amit Kumar Dey, Ajay Mate, and Samir Rege participated in writing the paper and revising the draft. Malav Modi and Ajay Mate took the photos. The manuscript has been read and approved by all the authors and represents honest work.

Acknowledgments

The authors would like to thank the patient for his written consent and permission to present this paper.

References

[1] T. P. F. Loong and H. M. Kocher, “Clinical presentation and operative repair of hernia of Morgagni,” Postgraduate Medical Journal, vol. 81, no. 951, pp. 41–44, 2005.
[2] T. P. Comer and O. T. Clagett, “Surgical treatment of hernia of the foramen of Morgagni,” Journal of Thoracic and Cardiovascular Surgery, vol. 52, no. 4, pp. 461–468, 1966.
[3] F. S. Rakas, K. G. Dayma, and D. B. Gabukamble, “Obstructed Morgagni hernia (A case report),” Indian Journal of Surgery, vol. 50, pp. 144–146, 1988.
[4] A. N. Gangopadhyay, V. D. Upadhya, D. K. Gupta, and S. P. Sharma, “Obstructed Morgagni’s hernia,” Indian Journal of Pediatrics, vol. 74, no. 12, pp. 1109–1110, 2007.
[5] M. D. Kelly, “Laparoscopic repair of strangulated Morgagni hernia,” World Journal of Emergency Surgery, vol. 2, no. 1, article 27, 2007.
[6] S. Arora, A. Haji, and P. Ng, “Adult Morgagni hernia: the need for clinical awareness, early diagnosis and prompt surgical intervention,” Annals of the Royal College of Surgeons of England, vol. 90, no. 8, pp. 694–695, 2008.
[7] R. P. Sakalkale, M. Sankhe, S. Nagral, and C. V. Patel, “Obstructed Morgagni’s hernia (a case report),” Journal of Postgraduate Medicine, vol. 37, no. 4, pp. 228–230, 1991.
[8] V. Papanikolaou, D. Giakoustidis, P. Margari et al., “Bilateral morgagni hernia: primary repair without a mesh,” Case Reports in Gastroenterology, vol. 2, no. 2, pp. 232–237, 2008.
[9] F. Paris, V. Tarazona, M. Casillas et al., “Hernia of morgagni,” Thorax, vol. 28, no. 5, pp. 631–636, 1973.
[10] S. T. Lin, D. M. Moss, and S. O. Henderson, “A case of Morgagni hernia presenting as pneumonia,” Journal of Emergency Medicine, vol. 15, no. 3, pp. 297–301, 1997.
[11] S. W. Harrington, “Various types of diaphragmatic hernia treated surgically; report of 430,” Surgery, Gynecology & Obstetrics, vol. 86, no. 6, pp. 735–755, 1948.
[12] R. Ackroyd and D. I. Watson, “Laparoscopic repair of a hernia of Morgagni using a suture technique,” Journal of the Royal College of Surgeons of Edinburgh, vol. 45, no. 6, pp. 400–402, 2000.