A CASE OF BILATERAL PARASTERNAL DIAPHRAGMATIC HERNIA

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Amid the several recognized types of diaphragmatic hernia the incidence of the parasternal variety is sufficiently low to render an example of its bilateral occurrence worthy of record. This extremely rare condition was encountered during the recent routine examination of a dissecting-room cadaver, a female, aged 83, in the Department of Anatomy of St. Bartholomew's Hospital Medical College.

CLINICAL HISTORY

In life the patient had displayed no symptoms directly referable to the unsuspected diaphragmatic hernia. She had suffered from chronic bronchitis with alleged emphysema, but had complained of no digestive or other alimentary symptoms. The patient's obesity rendered it unlikely that any clear physical signs indicative of the condition could have been detected on clinical examination, and at no time of life did any suspicion arise of gross visceral displacement. She died of myocarditis and advanced senile change.

THE HERNIAE

General Account.—The abdomen being opened, a gross alteration of visceral topography was immediately apparent, a considerable portion of the bowel being found to occupy the right side of the chest. On this side a loop of gut, consisting of terminal ileum, caecum, and proximal colon, together with a portion of the great omentum, ascended anterior to the liver and entered the thorax through a parasternal diaphragmatic defect. On the left an additional small colonic loop ascended anterior to the left lobe of the liver and entered the thorax through a second parasternal diaphragmatic defect. In each instance the herniated organs lay within a complete peritoneal hernial sac. No other visceral displacement was present: the stomach and spleen were normally situated and the retroperitoneal organs were undisturbed.

The Diaphragmatic Defects.—The two diaphragmatic defects (Fig. 1) were oval, with long axes disposed transversely, and were bilaterally situated immediately behind the seventh costal cartilage, being separated by the xiphiform portion of the diaphragm. The right and larger defect measured 7 cm. by 4.5 cm. and the left defect 3.5 cm. by 2 cm. Their orifices were bounded anteriorly by the seventh costal cartilage, laterally by the costal fibres of the diaphragm, and posteriorly by the anterior leaf of the central diaphragmatic tendon. They were separated by three distinct muscular slips attaching the central tendon to the back of the xiphisternum, and these were disposed with a convexity to the left side. The larger dextral defect extended across the midline behind the xiphisternum. The superior epigastric vessels entered the rectus sheath through the anterior lateral margin of each defect.

The Hernial Sacs.—Each hernial sac (Fig. 2) was continuous with the peritoneum clothing the diaphragm and anterior abdominal wall. The sacs were complete and were discrete from the pleural and pericardial cavities. The large dextral sac invaginated the diaphragmatic pleura and extended as high as the second costal cartilage, the fundus being distant some 15 cm. from the neck. The smaller sinistral sac extended upwards and to the left for some 6 cm. into the extrapericardial fat of the anterior mediastinum and did not contact the pleura.

The ligamentum teres hepatis and the falciform ligament were incorporated in the wall of the large sac. These structures entered the sac on its lateral side, curving first upwards and backwards, and then medially and downwards to escape over the posterior...
BILateral parasternal diaphragmatic hernia

This situation. The ligamentum teres passed forwards and downwards over the posterior margin of the diaphragmatic defect and was attached to the liver in the floor of the trough.

The right lung (Fig. 2b) was small and deformed by the large dextral hernia. The interlobar fissures were markedly incomplete, but dissection of the bronchial tree revealed a normal pattern of broncho-pulmonary segments. The segmental bronchi, particularly those of the middle lobe, showed a distortion commensurate with the general compression of the lung. The lower lobe extended caudally behind the hernia into a costo-diaphragmatic recess of normal extent. The pulmonary vessels were of normal disposition.

The left lung (Fig. 2c) was larger than the right, although it was somewhat compressed by the cardiac displacement. It consisted of the customary two lobes with a complete oblique interlobar fissure. The anatomy of the bronchial tree and the pulmonary vessels showed no abnormality.

The heart and pericardium (Fig. 2a) were displaced to the left so that the right cardiac border lay in the median plane and the cardiac apex some 3 cm. lateral to the left mid-clavicular line. The lower part of its right ventricular region was displaced somewhat backwards by the small sinistral hernia. The outer wall of the right atrium was concave and the inferior vena cava entered the atrium from the right infero-lateral aspect.

**Reports in the Literature**

A brief review of the literature reveals the rarity of the parasternal variety, which would appear to account for less than 3% of all diaphragmatic hernias. No previous report of its bilateral occurrence has been found.

Thoma (1882) describes four cases of diaphragmatic hernia of various types, one of which is
parasternal and closely resembles the dextral hernia of the present specimen, its sac containing the terminal ileum, caecum, and proximal colon. He reviews a total of 291 cases of diaphragmatic hernia, 26 of which he describes as "true herniae," i.e., possessing a peritoneal sac, and of these, eight are parasternal (six right-sided and two left-sided).

Keith (1910) reports on 34 museum specimens of diaphragmatic hernia, none of which is parasternal.

Kratzeisen (1921) describes a single example of left-sided parasternal hernia in an 83-year-old man. The peritoneal sac contained transverse colon and lay in the anterior mediastinum in front of the pericardium and thus closely resembled the condition encountered in the left side of the present subject.

Hume (1922) regards hernia through the foramen of Morgagni (space of Larrey) as a rare variety of diaphragmatic hernia, stating that only eight recorded cases were found in the literature, presumably those enumerated by Thoma.

Barrett and Wheaton (1934) refer to the rare class of diaphragmatic hernia through neural and vascular foramina and account parasternal hernia the least rare variety within this class, but say nothing further.

Dunhill (1935) reports 25 cases of diaphragmatic hernia, which include one case of parasternal hernia treated successfully by operation. In this patient, a man aged 29, the operative findings were as follows:

"There was an hiatus 4 in. long behind the xiphisternum, apparently central in situation with a tendency to lie towards the right side. The peritoneal sac contained the termination of the ileum, the caecum, appendix, ascending and transverse colon. The sac and its contents were in the anterior mediastinum resting against the pericardium on the left side and displacing the right pleura and lung backwards and to the right. The neck of the sac had spread the costo-sternal fibres of the diaphragm widely."

These findings show a considerable similarity to the condition in the right side of the present subject.

Harrington (1942) reviews 304 examples of diaphragmatic hernia treated by operation at the Mayo Clinic: five of these (about 2%) are described as "subcosto-sternal" (parasternal) in type. In one of these cases the hernial sac contained colon, caecum, appendix, terminal ileum, and great omentum, as in the present specimen. Of the others, one contained omentum and a loop of colon, while in the remainder the contents were omentum and fat.

Meyer (1950) reviews the subject of diaphragmatic hernia and describes "subcosto-sternal" hernia as the rarest type, giving, however, no statistics.

Discussion and Conclusions

Parasternal hernia is, then, one of the rarer forms of diaphragmatic hernia and no previous record of its bilateral occurrence has been discovered. It is clear that parasternal hernia is quite compatible with a normal life-span, as in the present example and in those described by Thoma and Kratzeisen, and, further, that it may be completely symptomless. When, however, symptoms occur, the hernia is susceptible of radiological diagnosis and surgical cure (Dunhill, 1935; Harrington, 1942).

Owing to its great rarity, little work has been done to elucidate the aetiology. The diaphragmatic defect occurs between the costal and xiphisternal origins of the diaphragm at the point where the superior epigastric vessels escape from the thorax to enter the rectus sheath, i.e., through the space of Larrey. This is also the site of the ventral part of the embryonal pleuro-peritoneal canal, and it thus seems reasonable to compare parasternal hernia with the more commonly occurring lumbo-costal hernia (through the pleuro-peritoneal hiatus of Bochdalek), which is acknowledged to be due to defective closure of the dorsal part of the canal. In this latter site the pleuro-peritoneal membrane may fail to close the canal completely with a resultant massive herniation of the abdominal viscera incompatible with life (Dunhill, 1935). On the other hand, the canal may be closed by the pleuro-peritoneal membrane, but subsequent failure of diaphragmatic muscle development may lead to herniation of abdominal viscera within a peritoneal sac, a condition compatible with life. The parasternal herniae described to date have all possessed a complete peritoneal sac, and Dunhill correctly concludes therefore that this hernia is not the result of a persistent embryonal pleuro-peritoneal hiatus. It appears (a point Dunhill omits) that parasternal hernia may be paralleled by the second type of lumbo-costal hernia and be due to defective muscular reinforcement of the pleuro-peritoneal membrane. The recording of parasternal hernia in adult subjects only does not imply the absence of a primary congenital defect, as such defect may become apparent only in later life. It would seem that the explanation of parasternal hernia lies primarily in the presence of a congenital structural defect.
that portion of the diaphragm which results from
the fusion of septum transversum and body-wall
components.

**SUMMARY**

A case of bilateral parasternal diaphragmatic
ehernia in an elderly woman is described.

No previous report of a similar condition has
been found in the literature.

Parasternal hernia is the rarest type of diaphragmatic hernia and probably accounts for less than 3% of all specimens.

The probable explanation of parasternal hernia is the presence of a congenital structural defect of that portion of the diaphragm which results from the fusion of septum transversum and body-wall components.

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**REFERENCES**


