ABSTRACT

Bochdalek hernias (BHs) are produced in the posterolateral area of the diaphragm. They are generally congenital, appearing in childhood, but are also detected in asymptomatic adult patients seeking medical attention for other reasons. Computed tomography (CT) or magnetic resonance imaging (MRI) is used for the correct diagnosis of the hernia type and for its localization, facilitating its management and the choice of treatment. We describe three cases of Bochdalek hernia, two on the right side and one bilateral, which was larger on the right than left side. All of these hernias contained only omental fat. In one patient, the right kidney was adjacent to the diaphragmatic defect but remained within the abdomen. The patients showed no symptoms and were not surgically treated. Examination by multi-slice CT with the possibility of coronal and sagittal reconstruction should be considered the standard method for diagnosing this entity. MRI in T1 is highly valuable to evaluate fat-containing chest lesions. The incidental finding of BH in asymptomatic adults is increasing, thanks to the wider application of new imaging techniques.

Key words: Bochdalek hernia. Congenital diaphragmatic hernia. Computed tomography. Magnetic resonance.

INTRODUCTION

The presence of a weakened space or defect in the diaphragmatic muscle can allow some contents of the abdomen to enter into the thoracic cavity, forming a herniation. Diaphragmatic hernias (DHs) are most frequently produced in the esophageal (hiatal hernia) and paraesophageal hiatus (paraesophageal hernia) and posterolateral –Bochdalek hernia (BH)–, and anteromedial –Morgagni hernia (MH)– regions of the diaphragm (1-4).

The etiology can be congenital, due to alterations in diaphragm development, or acquired as the result of surgery, trauma, or infection. The causes of late-presenting hernias (i.e., in adults) include congenital herniation, penetrating or blunt trauma, physical effort (including sexual relationships), pregnancy, labor or delivery, sneezing or coughing episodes, and even the intake of a large amount of food (4-7). It has also been related to long-term complications of continuous ambulatory peritoneal dialysis (8).

Most congenital diaphragmatic hernias (CDHs) appear in the neonatal period with respiratory distress and can be life-threatening (6,9,10). However, cases have been reported outside this age group, usually in adults with non-specific or no symptoms (5,11-13). In Spain, the frequency (14) of a hernia or diaphragmatic agenesis diagnosis in the first three days of life was reported to be 2.15 per ten thousand in 2007 and has shown a linear tendency to decrease in subsequent years (2.80/000 in 1980-1985; 1.46/000 in 1986-2006). BHs are more frequent than MHs (2,9,15), to the extent that CDH and BH have become virtually synonymous (4). The estimated prevalence of BH ranges from 0.17 to 12.7 per cent (2,5,7,9,10,16), with the higher values being detected in CT-based studies. HBs are more frequent than MHs (2,9,15), to the extent that CDH and BH have become virtually synonymous (4). The estimated prevalence of BH ranges from 0.17 to 12.7 per cent (2,5,7,9,10,16), with the higher values being detected in CT-based studies. HBs are more frequent than MHs (2,9,15), to the extent that CDH and BH have become virtually synonymous (4). The estimated prevalence of BH ranges from 0.17 to 12.7 per cent (2,5,7,9,10,16), with the higher values being detected in CT-based studies. HBs are more frequent than MHs (2,9,15), to the extent that CDH and BH have become virtually synonymous (4). The estimated prevalence of BH ranges from 0.17 to 12.7 per cent (2,5,7,9,10,16), with the higher values being detected in CT-based studies. HBs are more frequent than MHs (2,9,15), to the extent that CDH and BH have become virtually synonymous (4). The estimated prevalence of BH ranges from 0.17 to 12.7 per cent (2,5,7,9,10,16), with the higher values being detected in CT-based studies. HBs are more frequent than MHs (2,9,15), to the extent that CDH and BH have become virtually synonymous (4). The estimated prevalence of BH ranges from 0.17 to 12.7 per cent (2,5,7,9,10,16), with the higher values being detected in CT-based studies. HBs are more frequent than MHs (2,9,15), to the extent that CDH and BH have become virtually synonymous (4). The estimated prevalence of BH ranges from 0.17 to 12.7 per cent (2,5,7,9,10,16), with the higher values being detected in CT-based studies. HBs are more frequent than MHs (2,9,15), to the extent that CDH and BH have become virtually synonymous (4). The estimated prevalence of BH ranges from 0.17 to 12.7 per cent (2,5,7,9,10,16), with the higher values being detected in CT-based studies. HBs are more frequent than MHs (2,9,15), to the extent that CDH and BH have become virtually synonymous (4). The estimated prevalence of BH ranges from 0.17 to 12.7 per cent (2,5,7,9,10,16), with the higher values being detected in CT-based studies. HBs are more frequent than MHs (2,9,15), to the extent that CDH and BH have become virtually synonymous (4). The estimated prevalence of BH ranges from 0.17 to 12.7 per cent (2,5,7,9,10,16), with the higher values being detected in CT-based studies. HBs are more frequent than MHs (2,9,15), to the extent that CDH and BH have become virtually synonymous (4). The estimated prevalence of BH ranges from 0.17 to 12.7 per cent (2,5,7,9,10,16), with the higher values being detected in CT-based studies. HBs are more frequent than MHs (2,9,15), to the extent that CDH and BH have become virtually synonymous (4). The estimated prevalence of BH ranges from 0.17 to 12.7 per cent (2,5,7,9,10,16), with the higher values being detected in CT-based studies. HBs are more frequent than MHs (2,9,15), to the extent that CDH and BH have become virtually synonymous (4).
(12). Although a diaphragmatic dysgenesis is widely considered to be the origin of CDH, only 10 per cent of CDH patients were found to have chromosomal anomalies (17).

BHs can affect both sides of the body, in the region where the lateral arcuate ligaments of the diaphragm curve, covering the quadratus lumborum muscles, and laterally fix on either side of the twelfth rib and medially to the transverse process of L1 (9,10,18).

CASE REPORTS

Case report 1

An 82-yr-old man was hospitalized for intense epigastric pain that did not respond to analgesic medication; he also reported persistent constipation with altered bowel habit. He suffered from senile heart disease, aortic sclerosis and venous insufficiency in lower limbs. He had a history of chronic obstructive pulmonary disease (COPD) and a major atrial fibrillation treated with actocortin in the emergency area, which has not recurred. He subsequently underwent surgery for bladder cancer (grade I papillary transitional carcinoma involving the corium).

Conventional posteroanterior chest X-ray (Fig. 1) showed no findings of interest. Chest-abdominal computer tomography (CT) study revealed some paraseptal bullae with pulmonary emphysema, increased density of residual appearance in the apex of the right lung, interstitial pattern with bibasal predominance, cardiomegaly (at the expense of both atria), mitral and aortic valve calcifications, small subcarinal lymph nodes, and possible right hilar lymph nodes. There were also mild pleural thickenings with right predominance and chronic appearance. Additionally, there was a small right BH with fatty content of around 25 mm (Figs. 2 A-C). In the abdominal region, a small vesical diverticulum (12 mm) was observed on the right lateral wall.

Case report 2

A 61-year-old male was hospitalized for epigastric pain and vomiting, which were interpreted as a biliary colic. However this diagnosis could not be confirmed by X-ray, which showed no findings of interest (Figs. 3 A and B). Cholangio-MRI (Fig. 4) and abdominal-pelvic MRI studies (Fig. 5) revealed a previously undiagnosed bilateral BH.

Three years earlier, he had been diagnosed with moderately differentiated rectal adenocarcinoma (pT3 pN0), which was treated with surgery, chemotherapy, and radiotherapy.

At his latest examination, he was disease-free with normal blood-count, although he showed elevated levels of total cholesterol, 278 mg/dL (upper limit: 200-239), triglycerides, 244 mg/dL (50-150), and LDL-cholesterol, 153 mg/dL (50-130).

Finally, chest-abdominal CT revealed small subpleural pulmonary nodules, a small hiatal hernia, and bilateral BH with fatty content (Figs. 6 A-C), which was larger on the right side (around 48 mm). In the abdominal region, there were small cysts in the renal parenchyma and a metal suture in the rectum from the previous surgery.

Case report 3

An 81-year-old female with respiratory insufficiency was hospitalized for a respiratory infection. Analytical results showed: leukocytosis of 21,530 leukocytes per µL (normal range, 4,800-10,800), with a shift to the left due to the presence of 18,900 neutrophils per µL (1,900-8,000); 100 mg/dL glycemia (70-100); 51.7 mg/dL uremia (10-50); 3.2 mEq/L K (3.8-5.2); 14.30 mg/dL PCR (0-0.8); and 65 ng/mL myoglobin (25-58).

Posteroanterior (Fig. 7A) and lateral (Fig. 7B) chest X-rays showed alveolar-interstitial lung involvement, with predominance in inferior lobules, as well as right posterior diaphragmatic lobulation. The lateral X-ray showed dorsal kyphosis and wedge-shaped vertebrae. Chest CT (Figs. 8 A-D) confirmed the presence of an interstitial “honeycomb” pattern of micro- and macro-cystic lesions, with peripheral, apical, and basal predominance and the presence of traction bronchiectasis, indicating an advanced stage of pulmonary fibrosis. The scan also showed a sliding hiatal hernia and right BH with fatty content.

DISCUSSION

The diaphragm is formed between week 4 and 12 of gestation by four embryologic elements: septum transversum,
pleuroperitoneal membranes, mediastinal dorsal mesentery of the esophagus, and muscles of the body wall (10,19). It has been hypothesized that the liver usually obstructs herniation through a possible defect on the right side. Furthermore, the right hemidiaphragm is completely formed before the left, because of the earlier closing of the right pleuroperitoneal canal when the intestine returns to the peritoneum from its rotation in the yolk sac (11), hence 70-90 per cent of all BHs, better described as posterior diaphragmatic defects (19), are on the left side (6), although Mullins et al. (5), in 2001, observed that 68 per cent were located on the right side. They also found a 14 per cent possibility of bilaterality.

We report three cases of BH, two on the right side and one bilateral in which the largest hernia is on the right side. The size of a BH is usually highly variable, from a few millimeters to occupying most of the thorax, and it is not nec-

Fig. 2. Axial CT images of right unilateral BH with fatty content (arrows). A. Soft tissue window. B. Lung window. C. Soft tissue window of a more caudal slice, with small posteromedial defect of the right hemidiaphragm (arrow head).

Fig. 3. Posteroanterior (A) and lateral (B) chest X-rays. Absence of signs suggestive of BH. Slight medial and anterior elevation of the right hemidiaphragm (arrows) as a non-specific finding unrelated to BH. Catheter in superior vena cava (arrow heads).
Fig. 4. Axial cholangio-MRI image (117/4.6/70°) showing increased signal intensity behind both diaphragms (arrows) due to herniated fat.

Fig. 5. Axial T₁-weighted (1828/100) MRI image showing increased fat signal intensity in the bilateral BH (arrows).

Fig. 6. Axial CT images, with soft tissue window, showing bilateral BH (arrows) that contains fat alone: A. Cranial slice; B. Intermediate slice; and C. Caudal slice. We highlight the openings in both hemidiaphragms (arrow heads).

essarily related to the size of the diaphragmatic defect (5,11,20); 20 per cent of BHs are contained by a sac, and the remainder show direct communication between thorax and abdominal cavity (21). BHs typically contain omental fat, which can be accompanied by the stomach, spleen, colon, and even the small intestine (10,11,16,20,22,23). The involvement of the liver, gallbladder, pancreas, kidney, or retroperitoneal fat is rare (5), and there has been no report of a lung component in adult BH (10).

In all our cases, the content was omental fat alone. However, in one of them, the right kidney was adjacent to the diaphragmatic defect but inside the abdomen (Fig. 8B).
In newborns, the protrusion of the liver inside the right hemithorax is typically due to a CDH. In these cases, the liver will cause a mass effect and displace the mediastinum towards the left. However, if the mediastinum is not displaced, other less common causes for intrathoracic liver should be considered, such as primary right pulmonary hypoplasia (e.g., scimitar syndrome) or agenesis and hepatic-pulmonary fusion (24).

Most congenital BHs are associated with a respiratory insufficiency (21) that becomes evident during the first weeks of life, and they are among the most frequent causes of respiratory distress in neonates. In adults, most BHs are usually asymptomatic (16) and their detection is incidental, as in our three patients. Symptoms, if any, are typically imprecise. Patients usually report chest pain or gastrointestinal symptoms (5, 7, 13). A case was reported of a 23-yr-old woman in the second trimester of pregnancy with symptoms of retching, oliguria, and shortness of breath that were initially attributed to the pregnancy itself but finally proved to be caused by a left BH with acute gastric volvulus (25). One of our patients was hospitalized for intense epigastric pain, but its relationship with the BH could not be demonstrated. There have been rare reports of late presenting BH with gastric volvulus, spleen rupture, obstruction, and gastrointestinal perforation (23, 25, 26).

Congenital BHs can be diagnosed by ultrasound, even in the prenatal period, as a displacement of the mediastinum and a mass in the thorax of the fetus (2). Chest-abdominal X-rays of a neonate with BH and respiratory distress show a soft tissue mass in the thorax that reveals gas or hydro-air levels when the neonate swallows air. X-rays with radiopaque contrast may be useful before surgery to assess the herniated intestinal loops and their possible malrotation (10, 22).

Occasionally, a BH can appear as a lesion with soft tissue density in the posterior area of the pulmonary base in lateral chest X-rays (6, 11), which may be confused with a pulmonary lymph node adjacent to the diaphragm (27) or, as in one of our cases, with a posterior diaphragmatic lobulation (Figs. 7 A and B). Using CT, however, a BH can be readily differentiated from a diaphragmatic everted or lobulation (“bosselation”) by examining the continuity of the diaphragm muscle itself. In BH, the muscle always shows a break or defect, whereas it is always whole, though thinned, in the other two conditions (11).

In two of our patients, with known malignant disease, the predominant clinical indication for CT was to rule out metastatic disease, as reported by other authors (5).

Examination by multi-slice CT, with the possibility of coronal and sagittal reconstruction, should be considered the standard method to diagnose BH (5). Some authors recommended replacing conventional axial images with coronal reformatted images, because there are fewer images and less time is required for interpretation (28). However, it has been verified that the lack of familiarity of observers with coronal visualization may contribute to detection errors and the missing of defects (28).

CT and MRI studies are highly valuable for the assessment of fat-containing thoracic lesions (6), detecting, localizing, and identifying lesions and markedly reducing the time required for the differential diagnosis (Figs. 4 and 5). CT or MRI diagnosis can be definitive. In BHs, sagittal and coronal reformatted images can reveal the diaphragmatic defects and hernia contents (Figs. 8 C and D).

The treatment of incidental BH remains controversial (5). All DH types are assumed to have a congenital origin, but the clinical presentation, management, complications, and prognosis markedly differ between neonate and adult.
cases. The prognosis for late-presenting BH is usually more favorable (13,16), as in the present patients, who have shown no clinical or imaging changes after a follow-up of up to six years. Despite the generally symptom-free nature of incidental BH in adults, some authors recommend surgery, including hernia reduction and defect closure, arguing that the risk of tissue strangulation and even death warrants an aggressive approach (7).

The incidental finding of BH in asymptomatic adults appears to be increasing largely due to the application of new imaging techniques. The use of CT or MRI offers greater precision in the diagnosis, localization and characterization of the hernia, facilitating its management and the choice of treatment.

**REFERENCES**