Etiology of primary spontaneous pneumothorax
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ABSTRACT
With the advent of HRCT, primary spontaneous pneumothorax has come to be better understood and managed, because its etiology can now be identified in most cases. Primary spontaneous pneumothorax is mainly caused by the rupture of a small subpleural emphysematous vesicle (designated a bleb) or of a subpleural paraseptal emphysematous lesion (designated a bulla). The aim of this pictorial essay was to improve the understanding of primary spontaneous pneumothorax and to propose a description of the major anatomical lesions found during surgery.

Keywords: Pneumothorax; Pulmonary emphysema; Tomography, X-ray computed.

INTRODUCTION
Primary spontaneous pneumothorax is usually quantified and diagnosed in the ER by means of routine chest X-rays.¹,² However, it is not always easy to identify emphysematous bullae on routine chest X-rays. Emphysematous bullae are found in approximately 15% of cases and are primarily located on the apical margins of the affected lung lobes.³

Currently, HRCT is more sensitive than routine chest X-rays for the diagnosis of apical emphysematous lesions, which are found in approximately 80% of all patients with primary spontaneous pneumothorax.⁴,⁵ In contrast, the prevalence of paraseptal emphysema in such patients is only 3%.⁶

PATHOGENESIS
The pathogenesis of primary spontaneous pneumothorax can be best understood by understanding emphysematous lung lesions.

The anatomical lesion known as a bleb was first described in 1947 by Miller, who established an anatomical distinction between a bleb and a bulla (bulla emphysema).⁷

Later, in 1967, Reid divided bullous emphysema, which causes bullae, into three types: type I, a small amount of hyperinflated lung tissue that is narrow (pedunculated) and contains no lung parenchyma; type II, a relatively smaller amount of hyperinflated lung tissue that is broad (sessile) and usually contains vanishing lung; and type III, a large amount of hyperinflated lung tissue extending to the pulmonary hilum, with ill-defined margins and vanishing parenchyma in each bulla.⁸

Chest CT has greatly contributed to the description of lung disease.⁹

For a better description of the aforementioned emphysematous lung lesions, a glossary of terms for thoracic imaging is required.

Bleb
The word bleb is translated to Portuguese as vesícula enfisematosa subpleural (i.e., subpleural emphysematous vesicle). A bleb is caused by alveolar rupture, which allows air to travel through the interlobular septum that divides the secondary pulmonary lobules to the subpleural region. The subpleural region is displaced, and a subpleural emphysematous vesicle (i.e., a bleb) is thus formed. Unlike a bulla, which is a distal acinar (or paraseptal) emphysematous lung lesion, the displacement caused by subpleural interstitial emphysema is generally small in size, i.e., 1-2 cm in diameter (Figure 1).

On chest CT scans, a bleb appears as a thin-walled cystic air space contiguous with the pleura. However, the use of the term bleb by radiologists is discouraged because the distinction between a bleb and a bulla is arbitrary and of little clinical importance.

PULMONARY EMPHYSEMA
Pulmonary emphysema is characterized by irreversible enlargement of the airspaces distal to the terminal bronchiole, accompanied by alveolar wall destruction.

On CT scans, emphysema appears as focal areas or regions of low attenuation, usually without visible walls. In the case of panacinar emphysema, decreased attenuation is more diffuse. Emphysema is usually classified according to the part of the acinus that is predominantly affected, being didactically divided into centriacinar (centrilobular) emphysema, panacinar (panlobular) emphysema,
distal acinar (paraseptal) emphysema, and irregular emphysema associated with fibrosis.

**Centriacinar or centrilobular emphysema**

Proximal emphysema is designated centriacinar or centrilobular emphysema; however, it is more commonly termed simply emphysema.

Centrilobular emphysema is characterized by destroyed centrilobular alveolar walls and enlargement of respiratory bronchioles and associated alveoli. It is the most common form of emphysema in cigarette smokers.

In cases of centrilobular emphysema, spontaneous pneumothorax is usually more severe because it is associated with an underlying lung disease, being designated secondary spontaneous pneumothorax and possibly leading to severe respiratory failure.

CT findings are centrilobular areas of decreased attenuation, usually without visible walls, of nonuniform distribution and predominantly located in the lung apices.

**Panacinar or panlobular emphysema**

When emphysema affects the entire acinus, it is designated panacinar or, less commonly, panlobular emphysema.

**Distal acinar or paraseptal emphysema**

When emphysema is distal, affecting primarily the margins of the lung lobes, it is designated distal acinar or paraseptal emphysema, which is the form of emphysema that is most closely related to primary spontaneous pneumothorax.

Although paraseptal emphysema is multiform and can affect various portions of the lung lobes, it primarily affects the lung apices. The major subtypes of paraseptal emphysema are described below.

**Bulla/bullae**

A bulla (i.e., a subpleural emphysematous bulla) is an airspace measuring more than 1 cm—usually several centimeters—in diameter, sharply demarcated by a thin wall that is no greater than 1 mm in thickness.

On CT scans, a bulla appears as a rounded focal lucency or area of decreased attenuation bounded by a thin wall, occurring predominantly in the lung apices (Figure 2).

**Bullous emphysema**

Multiple bullae can coalesce, characterizing bullous emphysema. Bullous destruction of the lung parenchyma is frequently associated with other signs of pulmonary emphysema (centrilobular or paraseptal emphysema).

**Irregular emphysema associated with fibrosis**

Absence of “obvious fibrosis” was historically regarded as an additional criterion for pulmonary emphysema. However, the validity of that criterion has been questioned because some interstitial fibrosis can be present in emphysema secondary to cigarette smoking.

**PATHOPHYSIOLOGY**

Distension of an emphysematous bulla leads to an extreme thinning of its structure (Figure 3).

The absence of pleural mesothelial cells on histological examination has been demonstrated by scanning electron microscopy, occurring primarily on the thin outer surface of type I bullae and in some areas on the surface of type II bullae.\(^{(10)}\) Therefore, distension of emphysematous bullae is due to a reduction in the surface tension of their walls, and this can be explained by Laplace’s law.\(^{(11)}\) In addition, there is evidence of a congenital etiology; however, the pathogenesis of bullae remains controversial.\(^{(12)}\)

**ANATOMY**

Bullae are found in approximately 85% of all patients with primary spontaneous pneumothorax undergoing surgery (Figure 4).\(^{(13,14)}\) However, there is no consensus in the literature regarding the anatomical classification of emphysematous lung lesions on the basis of surgical
The etiology of primary spontaneous pneumothorax findings during surgical exploration are as follows:

A. Normal lung
B. Apical lung scarring, primary spontaneous pneumothorax possibly being caused by a small bronchioloalveolar fistula surrounded by fibrotic tissue and approximately 1 mm in diameter
C. Single or multiple blebs smaller than 2 cm in diameter
D. Single or multiple bullae (forming a cluster) larger than 2 cm in diameter in a single lung segment
E. Giant bulla
F. Multiple bullae in one or more lung lobes
G. Multiple bilateral bullae
H. Lobar emphysema
I. Recurrent pneumothorax, which can be accompanied by multiseptated, multiform membranous pleuropulmonary adhesions in the region where the bullae are located
J. Multiloculated membranous chronic pneumothorax that can affect most of the pleural cavity
K. Spontaneous hemothorax related to rupture of cord-shaped vascularized pleuropulmonary adhesions during lung collapse (Figure 3)\(^\text{16}\)

**Surgery**

Currently, the vast majority of bullae are resected by video-assisted surgery with a surgical stapler (Figure 5).

**Figure 3.** Video-assisted thoracoscopic surgery image of a bulla showing extreme thinning in the left lung apex and accompanied by pleural adhesion.

**Figure 4.** Schematic illustration of the types of emphysematous lung lesions. A: normal lung; B: apical lung scarring; C: small bleb; D: bulla; E: giant bulla; F: multiple bullae; G: multiple bilateral bullae; H: lobar emphysema; I: recurrent pneumothorax with multiform membranous pleuropulmonary adhesions; J: multiloculated membranous chronic pneumothorax; and K: spontaneous hemothorax (the last three being accompanied by pleural adhesions).
The use of CT in the preoperative period allows the identification of the emphysematous bullae (Figure 5). Currently, the vast majority of subpleural emphysematous bullae is visualized and resected video-assisted thoracic surgery with a surgical stapler. The staple line should preferably be positioned in healthy parenchyma or in an area without bullous emphysema (Figure 6). Therefore, the possibility of air leak in the immediate postoperative period is minimized. However, new emphysematous lesions (as seen on CT scans) have been reported to appear at the staple line during late follow-up in 37.1% of cases, and it has been suggested that this is a risk factor for recurrent primary spontaneous pneumothorax in the late postoperative period.\(^{(17,18)}\)

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