Pulmonary Sequestration: A 29 Patient Case Series and Review

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ABSTRACT
Introduction: Pulmonary sequestration also known as broncho-pulmonary sequestration is a rare disease, with very few case series reviewed in literature. In this study, we review the demographics, presentation, imaging and treatment of pulmonary sequestration in 29 patients from our institution, and provide comparison data from previously published series with an overview of the disease history.

Materials and Methods: Records reviewed for all patients evaluated and treated in our institution with a pathological proven diagnosis of pulmonary sequestration from January 2004 through December 2013. Collected data included demographics, clinical presentation, diagnostic imaging, location of the lesion, type of sequestration, and subsequent treatment.

INTRODUCTION
Pulmonary sequestration (PS) is a rare disease, of unknown aetiology, representing 0.1–6% of all structural lung diseases and developmental malformations. It is characterized by a mass of pulmonary tissue that becomes separated from the normal bronchopulmonary tree and supplied by one or more aberrant systemic arteries [1,2]. It is an entity for which no single embryonic hypothesis is clearly adapted, and no chromosomal abnormality yet identified. Many consider the disease part of an anomaly spectrum, with normal vessels supplying anomalous pulmonary tissue at one end, and aberrant vessels supplying normal lung tissue at the other [3,4] [Table/Fig-1].

Given its rarity, large institutional case series on PS are very few. In this study, we aim to analyse the demographics, presentation, diagnostic imaging, location, type, and treatment of pulmonary sequestration in our institution over a 9-year period. We also compare our findings to those reported in previously published series from different countries to assess regional differences in gender related PS type, and possible genetic and ethnic variations of the disease.

MATERIALS AND METHODS
We conducted a retrospective medical records review of all patients evaluated and treated in our institution with a pathological proven diagnosis of PS from January 2004 through December 2013. We searched the electronic medical records (EMR) using the key words: pulmonary sequestration, bronchopulmonary sequestration, extralobar and intralobar pulmonary sequestration. Reviewed EMR included office notes, surgical and interventional procedure notes, diagnostic imaging studies, and pathology reports. Our radiologists reviewed all patients’ imaging studies. We excluded patients who had imaging studies performed in outside facilities, which were unavailable for our review, and patients who had either an alternative or unproven diagnosis based on pathology reports. Collected data included demographics, clinical presentation, diagnostic imaging, location of the lesion, type of sequestration, and subsequent treatment.

RESULTS
Gender and Age
Of the 29 patients, 15 were males and 14 females, 8 (28%) were children 0-2 years, 1 adolescent age 17, and 20 (69%) adults 21-70 years, with a mean age of 42.

Presenting Symptoms
In adults, symptoms occurred in 18 patients (90%), with the most common presentation of cough associated with recurrent pulmonary infections in 13 patients (72%), haemoptysis in 3 patients (17%), and chest pain as the primary complaint in 2 patients (11%) [Table/Fig-2].

Imaging Studies
As the majority of the adult patients presented with respiratory symptoms mimicking pneumonia and asthma, chest X-rays were performed on all patients as the first imaging modality, this often...
showed nonspecific pulmonary opacities such as nodules, reticular infiltrates and mass-like consolidations. On CTA, lesions suspicious for pulmonary sequestration were identified, with a systemic arterial supply to the sequestered segment demonstrated in 25 patients (86%). In 4 patients (14%), where CTA failed to demonstrate the arterial supply, two were diagnosed with angiography and the other two with magnetic resonance angiography (MRA). Furthermore, a large aberrant artery was clearly identified on CTA in all three patients. In 19 patients (66%), the PS was located in the left lower lobe and who presented with haemoptysis (PPV=100%) [Table/Fig-3-5].

Age, Gender and Ethnic Group Related Sequestration Type: In the adult group (21 patients) intralobar sequestrations were found in the majority of cases (81%), while in children the extralobar type was more predominant (62%). Gender distribution of sequestration types revealed a higher prevalence of extralobar sequestrations among males 54% vs. females 46%. However, intralobar sequestrations remained the most common type in both sexes, found in 53% of males 54% vs. females 46%. Of the 29 patients the Whites ethnic group formed the majority (21/29) (73%), African Americans (5/29) 17%, and Hispanics (3/29) 10%. Ethnic related variations in sequestration types may not be representative of a true genetic influenced trend of the disease as the numbers of patients in the none-white groups are too small. However, we found a significant predominance of the extralobar type among the African American group of patients (4 of the 5 patients) 80% [Table/Fig-2].

Treatment
All children and symptomatic adults were treated surgically with resection of the sequestered segment (83%), and 2 of the 3 patients who initially presented with haemoptysis underwent preoperative embolization (7% of all patients). Surgical treatment was differed in two asymptomatic adults due to significant medical co-morbidities and high operative risk. These were later followed with repeated CT scans.
anomalies in association with intralobar sequestration, as opposed to the extralobar type [17]. Additionally, more than half of intralobar sequestrations are diagnosed in later childhood or adulthood, while the majority of extralobar types are diagnosed during infancy [18]. Previous studies have demonstrated a greater male prevalence for the extralobar sequestration, approximately 3:4 times more common in males than in females [19-22]. In our population, the extralobar type was only slightly more common in males 54% vs. females 46%. The main clinical manifestations of PS in adults are recurrent respiratory tract infections, and respiratory symptoms that often mimic other common pulmonary diseases, such as pneumonia, asthma, interstitial and obstructive pulmonary diseases [23]. In a recent study, the misdiagnosis rate of PS was reported as high as 87%, due to the wide array of nonspecific symptoms, delaying treatment and limiting therapeutic options [24].

In the largest PS case series, Yong et al., retrospectively evaluated 2625 patients from China, where the majority had presented with respiratory symptoms of cough, sputum production, fever, haemoptysis, and chest pain, while 13% of patients presented with incidental imaging findings without respiratory symptoms. The main CT findings in this large group of patients included a mass lesions (49%), cystic lesions (29%), cavity lesions (12%), and pneumonic lesions (8%) [25]. A comparison review of previously published series from different countries [Table/Fig-8] [6,18,25-27], demonstrates a universal male predominance of the disease among the adult population, with the majority of cases being intralobar, and more than 80% of patients typically presenting with respiratory symptoms. Surgical resection of the sequestered lesion was performed in approximately 90% of cases without significant postoperative complications.

Surgical resection remains the standard of care for all patients with PS, as it serves to establish the diagnosis, differentiate anomalous pulmonary tissue from malignancies, and prevent recurrent infectious or haemorrhagic complications [27]. Imaging techniques that can accurately identify the arterial supply and venous drainage of the sequestered segment are considered the cornerstone of preoperative planning. To prevent intraoperative morbidity and mortality caused by an aberrant vascular haemorrhage in these patients, imaging studies are applied as an essential guide for the surgical procedure.

Pulmonary angiography is thought to be the gold standard for diagnosis; however, in recent years, several studies have shown that less invasive imaging techniques such as CTA and MRA may be equally effective and safer alternatives [28-31]. In a study by Yue et al., the treatments planned with CTA showed high accuracy, and sensitivity (both >90%) in both patient and artery-based evaluations, suggesting the potential utility of CTA as the primary tool in pre-treatment planning for most PS patients [32].

In 1861, Rokitansky introduced the first description of the disease, and proposed that PS occurs when pulmonary tissue separates from the lung during embryogenesis; this was later known as “the fraction theory” [6] [Table/Fig-6]. Since then, many theories were introduced to explain the possible underlying pathophysiological mechanisms producing the malformation [Table/Fig-7] [7-15].

In 1946, Pryce was the first to use the term “pulmonary sequestration”, and classified the disease into intralobar and extralobar types. An intralobar sequestration is the most common form of the disease where the lung malformation remains within the visceral pleura of its lobe, whereas the extralobar type corresponds to a true accessory lung, with a separate pleural envelope and an aberrant venous drainage [7].

As the ectopic accessory bud arise from the primitive foregut, during the fourth week of gestation, a lung-resembling structures develops without communicating with the main bronchial tree, it becomes enveloped by its separate visceral pleura and derives its arterial supply from the systemic circulation. An extralobar sequestration can be sub-classified depending on the level at which the accessory bud branches off the primitive foregut into intrathoracic type (supradiaphragmatic), and abdominal type (subdiaphragmatic) [16].

Intralobar sequestration theories debate in defining the disease as either a congenital or acquired condition. Theories of the acquired aetiology state that chronic pulmonary infections of a lung tissue disconnected from the normal bronchial tree can cause hypertrophy of a regional systemic artery, hence the aberrant arterial supply. Despite being separated from the bronchopulmonary tree, infections may spread to the sequestered segment from adjacent aerated lung tissue via accessory inter-alveolar or broncho-alveolar connections, e.g. pores of Kohn, and canals of Lambert. These theories are supported by the low incidence of other congenital anomalies in association with intralobar sequestration, as opposed to the extralobar type [17].

In a recent study, the misdiagnosis rate of PS was reported as high as 87%, due to the wide array of nonspecific symptoms, delaying treatment and limiting therapeutic options [24].

<table>
<thead>
<tr>
<th>Reference</th>
<th>Theory</th>
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<tbody>
<tr>
<td>1861 Rokitansky</td>
<td>Fraction theory: separation of normally- developed lung.</td>
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<tr>
<td>1875 Ruge</td>
<td>Excess theory: the malformation arises separately from the normal lung as the development of a rudimentary third lung.</td>
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<tr>
<td>1946 Pryce</td>
<td>Blood supply theory: insufficient arterial blood supply results in persistence of embryologic thoracic aortic arteries to the lung segment, with systemic blood pressure induced cystic degeneration of lung tissue.</td>
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<tr>
<td>1958 Boyden</td>
<td>No causal relationship between nonfunctioning lung and systemic artery.</td>
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<tr>
<td>1959 Gebauer and Mason</td>
<td>Theory of acquired anomalies: PS results from recurrent localized infectious process.</td>
</tr>
<tr>
<td>1965 Smith</td>
<td>Failure of embryologic growth factor signaling.</td>
</tr>
<tr>
<td>1968 Genovese and Warner</td>
<td>Congenital bronchopulmonary foregut malformation theory: Accessory lung bud develops in the embryo and becomes either incorporated in the normal lung (Intralobar) or remains separated (extralobar).</td>
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<tr>
<td>1968 Morscacci and Wylie</td>
<td>Intalobar sequestration is a collection of bronchogenic cysts associated with a systemic artery.</td>
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<tr>
<td>1984 Stocker and Malczak</td>
<td>An acquired malformation with arterial supply derived from hypertrophy of pulmonary ligament arteries.</td>
</tr>
<tr>
<td>1987 Clements and Warner</td>
<td>Malinosculation theory: a disruption the normal bronchopulmonary connection caused by an insult.</td>
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[Table/Fig-7]: Theories in the pathogenesis of Pulmonary Sequestration [7-15]
In our series, CTA had a diagnostic sensitivity and specificity of 86% and 100%, respectively, and served as the primary diagnostic and preoperative planning tool.

**CONCLUSION**

Pulmonary sequestration is a rare disease with multiple theoretical aetiologies, often misdiagnosed due to its variable and nonspecific presentations. CTA and other imaging modalities, which can accurately identify the arterial supply and venous drainage of the sequestered segment, are essential for diagnosis and preoperative planning.

**REFERENCES**


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