Pulmonary sequestrations: Experience of 31 patients

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ABSTRACT
Aims: Pulmonary sequestration (PS) is a rare congenital lung malformation. It is characterized by a nonfunctional pulmonary tissue that have no relation to the bronchial system. Case series are rare due to low incidence of pulmonary sequestration. In this study, we aim to analyze the presentation, demographics, diagnostic procedures, location, type and treatment of PS in our institution over 12-year period.

Methods: We retrospectively evaluated medical records of pathologically proved PS from January 2003 through December 2015 in our institution. Thirty-one patients were included in the study. Demographics, clinical presentation, diagnostic imaging, location, type of sequestration, type of surgery were collected.

Results: Of 31 patients, 3 patients (9.6%) were female and 28 patients were (90.3%) male. Average age was 24.6 years. Twenty-four (77%) patients had described symptoms. Chest X-ray scanning was performed to all patients as the first imaging modality. Of the 15 patients (48.4%) had intralobar sequestration and 16 patients (51.6%) had extralobar sequestration. Extralobar sequestrations were more common (55.1%). Lower lobectomy in 13 patients, segmentectomy in 2 intralobar sequestration patients, and simple mass excision in all extralobar sequestration patients (16 patients) were performed as surgical procedure. Six patients had postoperative complications: prolonged air leak in 4 patients, pneumonia in 2 patients.

Conclusions: It has similar appearance with many diseases of lung with traditional imaging methods and cause difficulties in diagnosis. Proving the presence of a feeding systemic artery with angiographic imaging is essential for definitive diagnosis. The recommended optimal treatment for pulmonary sequestration is resection of the sequestered tissue by segment or lobar resection.

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Introduction
Pulmonary sequestration (PS) is a rare congenital lung malformation with a rate of 0.15-6.4% (1). It is characterized by a nonfunctional pulmonary tissue that have no relation to the bronchial system. It provides arterial blood supply directly from thoracic or abdominal aorta or from one or several intercostal branches of the aorta (2). Pathogenesis of this rare entity has not clearly understood so far and has not been associated with a chromosomal abnormality yet. There is no consensus on one single embryonic hypothesis. The most common idea is that these lesions appear as accessory bud and move together with the developing esophagus and provides its blood supply from systemic circulation (3,4). This disease is considered in the disease group with normal or abnormal pulmonary tissue with normal or abnormal vascularization.

Two type of PS has been defined: in the lobar parenchyma as intralobar pulmonary sequestration (ILS) and outside of the lob as extralobar pulmonary sequestration (ELS). ILS is more common than ELS (86-75%) (5). Extralobar sequestrations are distinguished by having their own pleura.

Pulmonary sequestration in adult patients is usually diagnosed after a patient’s complaint of symptoms or incidentally when having a health check or thoracic scanning for other reasons.

Case series are rare due to low incidence of pulmonary sequestration. In this study, we aim to analyze the presentation, demographics, diagnostic procedures, location, type and treatment of PS in our institution over 12-year period.

Methods
We retrospectively evaluated medical records of pathologically proved PS from January 2003 through December 2015 in our institution. We reevaluated all patients with surgery notes and imaging studies. We excluded patients who had only radiologic diagnosis and refused surgical treatment. Thirty-one patients were included in the study. Demographics, clinical presentation, diagnostic imaging, location, type of sequestration, type of surgery were collected. Approval for the study was granted by the Ethics Committee of Gulhane Training and Research Hospital (Approval Date: March 26, 2019, Approval Number: 19/106).
Results

Of 31 patients, 3 patients (9.6%) were female and 28 patients were (90.3%) male. Average age of the patients was 24.6 years, ranging between 12 and 50 years. There were no children, two of patients were at adolescent age, 12 and 16 years. Twenty-nine adults were between 20-50 years (Table 1).

Table 1. Demographic distribution of pulmonary sequestration patients (n=31)

<table>
<thead>
<tr>
<th>Gender</th>
<th>n</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>28</td>
<td>90.3</td>
</tr>
<tr>
<td>Female</td>
<td>3</td>
<td>9.7</td>
</tr>
</tbody>
</table>

Age

<table>
<thead>
<tr>
<th>Age</th>
<th>n</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Children</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Adolescent</td>
<td>2</td>
<td>6.5</td>
</tr>
<tr>
<td>Adults</td>
<td>29</td>
<td>93.5</td>
</tr>
</tbody>
</table>

Presentation

<table>
<thead>
<tr>
<th>Presentation</th>
<th>n</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Symptomatic</td>
<td>24</td>
<td>77</td>
</tr>
<tr>
<td>- Dyspnea</td>
<td>10</td>
<td>42</td>
</tr>
<tr>
<td>- Recurrent episodes of pneumonia</td>
<td>7</td>
<td>29</td>
</tr>
<tr>
<td>- Chest pain</td>
<td>5</td>
<td>21</td>
</tr>
<tr>
<td>- Hemoptysis</td>
<td>2</td>
<td>8</td>
</tr>
<tr>
<td>Asymptomatic</td>
<td>7</td>
<td>23</td>
</tr>
</tbody>
</table>

Twenty-four (77%) patients had described symptoms: shortness of breath in 10 patients, recurrent episodes of pneumonia in 7 patients, chest pain in 5 patients, hemoptysis in 2 patients (Table 1).

Chest X-ray scanning was performed to all patients as the first imaging modality, which often showed nonspecific opacities such as mass like density. In 6 patients (19.3%), Computed Tomography Angiography (CTA) identified the suspicious lesions for pulmonary sequestration by demonstrating a systemic arterial supply to the sequestrated segment. In one patient (3.2%), arterial blood supply was diagnosed with an Magnetic Resonance Angiography and in one patient (3.2%), arterial blood supply was diagnosed with conventional angiography. CTA revealed aberrant arteries in two patients which were presented with hemoptysis. Bronchoscopy was performed to determine the suspected area of hemoptysis in symptomatic patients.

Of the 15 patients (48.4%) had ILS and 16 patients (51.6%) had ELS. There were 15 male (93.7%) patients and one female (6.3%) patients in ELS group, and there were 2 female (13.3%) patients and 13 male (86.7%) patients in ILS group (Table 2).

Table 2. Gender related types of PS

<table>
<thead>
<tr>
<th></th>
<th>Female</th>
<th>Male</th>
</tr>
</thead>
<tbody>
<tr>
<td>ELS</td>
<td>1</td>
<td>15</td>
</tr>
<tr>
<td>ILS</td>
<td>2</td>
<td>13</td>
</tr>
<tr>
<td>Total</td>
<td>3</td>
<td>28</td>
</tr>
</tbody>
</table>

Although the rates were similar in adult group (29 patients), extralobar sequestrations were more common (%55.1). Two adolescent patients’ sequestrations were both intralobar type, and located in left lower lobe. In all female patients, sequestrations were located in left lower lobe and two of them were ELS but one was ILS. The most common type in women was ILS, but ELS in men.

In 21 patients (67.8%), the PS were located in the left lower lobe and in 8 patients (25.8%) in the right lower lobe. One PS (3.2%) was located in the right middle lobe and one PS (3.2%) was located in the left upper lobe.

Half of the sequestrations in the left lower lobe (22 patients, 70.9%) were ELS and the other half were ELS. One patient had ILS-type sequestration in the right middle lobe. Of the 8 patients (25.8%) in which the sequestration was located in the right lower lobe, 3 of them were (37.5%) ILS, and 5 of them (63.5%) were ELS (Table 3).

Table 3. Type related localization of PS

<table>
<thead>
<tr>
<th></th>
<th>ELS</th>
<th>ILS</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Right</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Upper lobe</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Middle lobe</td>
<td>0</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Lower lobe</td>
<td>5</td>
<td>3</td>
<td>8</td>
</tr>
<tr>
<td>Left</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Upper lobe</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Lower lobe</td>
<td>11</td>
<td>11</td>
<td>22</td>
</tr>
<tr>
<td>Total</td>
<td>16</td>
<td>15</td>
<td>31</td>
</tr>
</tbody>
</table>

Lower lobectomy in 13 ILS patients, segmentectomy in 2 ILS patients, and simple mass excision in all ELS patients (16 patients) were performed as surgical procedure. Twenty-nine of the resections were performed through posterolateral thoracotomy and two of them were performed by video-assisted thoracoscopic surgery. There were intraoperatively detected pleural adhesions around the lesions in all patients. In 2 patients a concomitant bronchogenic cyst was detected and resected in the same session. In one patient, there was solitary pulmonary nodule in the right upper lobe which sized in diameter as 1x1.5 cm and have diffuse calcification. The postoperative pathological result was hamartoma.

One patient (3.2%) with ELS had systemic blood supply originating from left subclavian artery (Figure 1), and the rest of the patients (96.8%) had systemic blood supply originating from the aorta (Figure 2). One patient (3.2%) had abdominal aortic branch as systemic blood supply. Three patients (9.7%) had between 2 and 5 aortic branches and one patient had over 5 aortic branches as systemic blood supply.

Figure 1. CT scan of the left ELS (a) and angiography of systemic blood supply originating from left subclavian artery (b).

Six patients had postoperative complications: prolonged air leak in 4 patients, pneumonia in 2 patients. Pneumonia treated with appropriate antibiotic treatment. During the follow up peri-
tumor and lung cancer are prominent (13). This large differen-

denoid malformation, congenital lobar emphysema, neurogenic

Figure 2. CTA scan of systemic blood supply originated from the aorta (a) and the intraoperative view of the systemic artery (b).

od ranging between 4 month and 13 years (mean 28.4 month),

there was only one mortality because of myocardial infarction

at the age of 54 after 4 years from the surgery.

Discussion

Pulmonary sequestration may remain silent in patients and
can be diagnosed at an adult age (6). Adult patients usually
manifest upper airway syndrome, hemoptysis, or repeated lung
infections. Diagnosis is usually confirmed by imaging results
based on the above symptoms. Surgical resection of the dis-
eased lung, either lobar or sublobar resection, has been estab-
lished as the definitive treatment for eliminating active symp-
toms and preventing the progression of airway complications

The etiology of PS is still not clear but some theories have
been suggested so far. The commonly accepted hypothesis is
suggested by Corbett and Humphrey. Accordingly, a separate
lung bud is formed during intrauterine growth and this bud is
blooded from the systemic circulation (4). It is widely accepted
that ELS has congenital origin, but the origin of ILS is contro-
versial. The low incidence of congenital anomalies in ILS sup-
ports the idea of acquired etiology (8). According to this view,
small systemic arteries in the pulmonary ligament invade the
infected tissue as a result of continuous infection attacks (7).

ILS is seen about three-quarters of all cases (10,11). In our
series, number of the ELS and ILS cases were almost equal in
numerical order. It has been reported that ELS is more common
in men. ILS does not have such a frequency (1,4). In our series,
15 of 16 ELS patients were male.

Extralobar sequestrations are separated from the ILS with
their own visceral pleura. Extralobar sequestration is most
commonly seen (63%) between the diaphragm and the lower
lobe. Intralobar sequestration is located in lower lobes with rate
of 98% (4). Bilateral cases are found very rarely (12). Frequent
occurrence of the PS in the lower lobes should be stimulant
for clinicians. In our series of 31 patients, in 30 patients’
sequestrations were located in the lower lobes and there were
no bilateral cases. Atypical placements for ELS have been
reported and congenital anomaly association is more frequent
than ILS (11). The most common anomaly associations are;
diaphragmatic hernia and defects, cardiopulmonary anomalies
and foregut connection. In our series, we determined that ELS
is located in the middle lobe in one patient. Two patients with
bronchogenic cysts had ELS-type sequestration.

Wei et al. showed that 58.3% of cases were misdiagnosed
in the preoperative diagnosis of sequestration in their series of
2625 patients (12). In differential diagnosis, malignancies such
as abscess, pneumonia, hypoplasia, pericardial cyst, cystic ad-
encoid malformation, congenital lobar emphysema, neurogenic
tumor and lung cancer are prominent (13). This large differen-
tial diagnosis list demonstrates the importance of preoperative
rigorous evaluation.

Chest X-ray and computed tomography (CT) scan shows
nonspecific images like mass, cystic, cavity, or pneumatic
lesions. For the diagnosis of PS, the presence of a feeding
systemic artery must be proved (9). Imaging examination can
clearly reveal aberrant feeding arteries, providing key informa-
tion for preoperative diagnosis. Enhanced chest CT can clearly
show aberrant feeding arteries in a noninvasive manner and
has thus become the preferred examination for PS; it can also
reveal the abnormal drainage veins. Magnetic resonance im-
ageing can detect the relationship between the lesion and the
abnormal feeding arteries of the systemic circulation without
the use of contrast agents; however, it is less useful than enhanced
CT in revealing the pathologic features of PS (14, 15). In our
series, six of the patients were evaluated with CTA, one patient
was evaluated with MRA in the preoperative period. It is stat-
ed in literature that anomalous systemic arterial supplies came
from the descending thoracic aorta (72%); the abdominal aorta,
celiac axis, or splenic artery (21%); the intercostal artery (3%);
and rarely the subclavian, left gastric, superior mesenteric,
phrenic, and renal arteries (and their branches) or via the peri-
cardium and coronary artery (14,16). In our series, as it is very
rare in the literature, one patient (3.2%) with ELS had systemic
blood supply originating from left subclavian artery (Figure 1),
and the rest of the patients (96.8%) had systemic blood supply
originated from the descending aorta (Figure 2). One patient
(3.2%) had abdominal aortic branch as systemic blood supply.
Three patients (9.7%) had between 2 and 5 aortic branches
and one patient had over 5 aortic branches as systemic blood
supply.

All patients selected for this study were operated in our
clinic and their diagnosis was confirmed perioperatively and
confirmed by pathology. In two cases, we observed the bronchial
relationship of PS to the main bronchi, intraoperatively, but
we could not prove this radiologically because the imaging
studies could not be reached due to technical problems. If this
connection can be proved in the future studies, this connection
may create a new subtype of congenital malformation.

Clinical findings are miscible with many other lung diseases
and are not specific in the diagnosis of PS. Cough, sputum,
fever, hemoptysis and chest pain are the most common
symptoms (12). The most common symptom in our series
was dyspnea (n=10). Five patients had chest pain. Pulmonary
sequestration may be asymptomatic or accidentally diagnosed.
Seven of our cases were asymptomatic. Although patients can
be diagnosed at any age, they are most frequently diagnosed
in the first two decades (12). In our series, the mean age at
diagnosis was 24.6 years.

Surgical resection of the diseased lung is recommended for
symptomatic patients and should also be performed for some
asymptomatic patients to control and prevent recurrent
pulmonary infection by stopping the inflammatory process and
to lower the incidence of airway complications (7). Medical
treatment is effective only in the treatment of accompanying
infections. In our series, sequestered mass was removed by
cutting the systemic artery in all ELS cases. Thirteen of the
ILS cases underwent lobectomy and two of them underwent
gonectomy. During the surgery, especially trans diaphragmatic
abdominal aortic branches may cause dangerous situations. While dissection, rupture of an abdominal
aortic branch may drop in to abdomen which can cause massive bleeding which needs urgent laparotomy.

The main purpose of surgery is to prevent complications. Because, fungal infection, tuberculosis, fatal hemoptysis, massive hemothorax and malignancy have been reported in untreated cases (17). In our series, 2 patients presented with hemoptysis. Embolization may be considered in certain cases that do not accept surgery or not suitable for medical condition (18).

**Conclusion**

Pulmonary sequestration is a rare disease in which some claims about etiology have been suggested. It has similar appearance with many diseases of lung with traditional imaging methods and cause difficulties in diagnosis. Proving the presence of a feeding systemic artery with angiographic imaging is essential for definitive diagnosis. Differential diagnosis is clinically important because untreated cases are open to complications. The recommended optimal treatment for pulmonary sequestration is resection of the sequestered tissue by segment or lobar resection.

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**Conflict of Interest**

The authors declared they do not have anything to disclose regarding conflict of interest with respect to this manuscript.

**References**