A Case of Tracheobronchial Amyloidosis Treated with Endoscopic Debunking and External Beam Radiation Therapy

Therapie einer tracheobronchialen Amyloidose mit endoskopischer Rekanalisation und Bestrahlung

Authors
I. Firlinger1, U. Setinek1, H. Koller1, P. Feurstein3, H. Prosch4, O. C. Burghuber1, A. Valipour1

Institutions are listed at the end of article.

Introduction

Amyloidosis refers to a variety of conditions in which amyloid proteins are abnormally deposited in organs and/or tissues. Tracheobronchial Amyloidosis is an orphan disease, thus there are no uniform treatment recommendations.

Case Report

We report on a 55-year-old man of Turkish descent who was admitted to hospital because of recurrent pneumonia. Besides a smoking history of 30 pack-years his past medical history was unremarkable.

Physical examination was entirely unremarkable. Spirometry demonstrated an (postbronchodilator) airway obstruction with a FEV1 of 1.27 L (38% predicted), FVC 2.83 L (71% predicted), and FEV1/FVC ratio of 54%. Bodyplethysmography disclosed normal total lung capacity (6.03L, 107% predicted) and a residual volume of 3.2 L. Laboratory results were normal. A chest CT scan revealed extensive irregular thickening of the walls of the right and left main bronchi, as well as the lobar and segmental bronchi on both sides with some calcifications. The thickening of the walls of the bronchi led to a significant narrowing of the right upper lobe and the lower lobe bronchi on both sides.

Videobronchoscopy demonstrated polypoid, toric-shaped changes of the distal tracheal wall spreading into both the left and right bronchial system up to the 4th generation of bronchi. The observed bronchial abnormalities were leading to multilevel obstruction of the central airways bilaterally (Fig. 1a, 1b). Biopsies were taken. The microscopic work-up showed amorphous eosinophilic deposits in the bronchial walls. Congo-red staining revealed the characteristic apple-green birefringence of the amyloid deposits by
Fig. 1a, b Endobronchial polypoid, toric-shaped changes of the distal tracheal wall spreading into the left and right bronchial system.

polarizing microscopy. The overlying mucosa showed squamous cell metaplasia. Between the amyloid deposits osseous metaplasia was found. Immunhistochemical analysis revealed evidence of Amyloid A. Further diagnostic testing was performed to rule out systemic involvement of amyloidosis. Due to the extent of endobronchial manifestations we performed endoscopic resection of the amyloid tumor masses under combined rigid and flexible bronchoscopy and general anesthesia in 3 subsequent sessions within a period of 6 months. Treatments were uneventful. Besides a clinical improvement of our patient, there was evidence of functional improvements in oxygenation and spirometric testing (Table 1).

At a follow-up visit after 12 months computed tomography and bronchoscopy revealed progression of the tracheobronchial lesions with recurrence of bilateral airways obstruction. Due to persistent endobronchial obstruction external beam radiation therapy (EBRT) was initiated. The dose administered was 20Gy in 10 fractions of 2Gy each. The volume encompassed the entire trachea beginning inferior to the vocal cords down to and including both mainstem bronchi. The radial margin was 1.5cm; the dose was administered in 4-field-technique. The patient tolerated the irradiation well. Six months later, our patient underwent a repeat bronchoscopy, the endobronchial situation revealed stable disease. During a final telephone interview 24 months after radiation therapy the patient reported a good general state without further progression of the disease.

Table 1  Pulmonary Function Testing (PFT) before and after endoscopic resection.

<table>
<thead>
<tr>
<th></th>
<th>PFT before endoscopic resection</th>
<th>PFT after endoscopic resection</th>
</tr>
</thead>
<tbody>
<tr>
<td>VC</td>
<td>2,831 (71% pred.)</td>
<td>3,351 (85% pred.)</td>
</tr>
<tr>
<td>FEV1</td>
<td>1,271 (38% pred.)</td>
<td>2,261 (69% pred.)</td>
</tr>
<tr>
<td>FEV1/FVC</td>
<td>0,45</td>
<td>0,67</td>
</tr>
<tr>
<td>TLC</td>
<td>6,031 (107% pred.)</td>
<td>6,001 (107% pred.)</td>
</tr>
<tr>
<td>RV</td>
<td>3,21</td>
<td>2,651</td>
</tr>
<tr>
<td>DLCO</td>
<td>77%</td>
<td>85%</td>
</tr>
</tbody>
</table>

Discussion

Localized pulmonary amyloidosis is rather rare and considered a primary form of amyloidosis. The mean age of patients with tracheobronchial amyloidosis ranges between 54 and 62 years [1 − 5]. While Rubinow et al [4] reported an increased prevalence of tracheobronchial amyloidosis in men, Capizzi et al [5] did not observe gender-related differences. There is some evidence of a relationship with smoking history [5]. Patients may have symptoms associated with endobronchial obstruction, such as recurrent lower respiratory tract infections, cough, rhonchi and wheezing on auscultation, hemoptysis, hoarseness, and/or stridor [1, 2, 4]. Chest-imaging (X-ray, computed tomography) usually shows narrowing of the distal trachea and mainstem bronchi, with some postobstructive evidence of atelectasis and/or pneumonia [1, 2]. CT imaging provides quantitative assessment of airway narrowing and mural thickening, two major consequences of amyloid infiltration. [6] The presence of mural calcifications sparing the posterior tracheal membrane may be misclassified as tracheobronchopathy osteochondroplastica [6]. Endobronchial visualization of the pathology reveals submucosal shiny plaques which may result in airway obstruction. Endoscopic biopsies are needed to obtain the diagnosis [2]. The disease can progress without treatment and life threatening complications may occur [5], however, spontaneous regression has also been documented [7]. Thirty percent of patients die within 7 − 12 years after diagnosis [8]. Systemic medications such as colchicine, melphalan, and corticosteroids have been attempted with little or no benefit. [9]. Local treatment is indicated in the presence of symptoms due to airway obstruction. Bronchoscopic resection with forceps and laser therapy has been used in this indication [5]. The recurrence of lesions and the inaccessibility of some lesions to bronchoscopic intervention, however, is an important limitation to this technique. Repeated bronchoscopic resections are therefore often necessary [5]. Neben-Wittich and co-workers reported a series of seven patients who were treated with EBRT as the only intervention. Objective improvement was observed in 4 out of 7 patients with a 57% success rate within 1 year when using FEV1 as a surrogate. It appears that radiation therapy may change the progressive course of tracheobronchial amyloidosis [10]. The mechanism by which EBRT exerts its effects on the amyloid deposits is unclear. It is possible that radiation therapy has an effect on local plasma cells as shown in patients with multiple myeloma [11]. EBRT may further result in a local inflammatory re-
sponse which may stop the progression of the disease, however, there is no particular evidence to support this theory.

In conclusion, we recommend considering a combination of bronchoscopic resection and consecutive external beam radiation therapy in patients with tracheobronchial amyloidosis.

Conflict of interest

The authors have no conflict of interest.

Institutions

1 Department of Respiratory and Critical Care Medicine, Ludwig-Boltzmann-Institute for COPD and Respiratory Epidemiology, Otto Wagner Hospital, Vienna, Austria
2 Institute of Pathology and Bacteriology, Otto Wagner Hospital, Vienna, Austria
3 Institute of Radiooncology, Wilhelminenspital, Vienna, Austria
4 Department of Radiology, Otto Wagner Hospital, Vienna, Austria

References