Atypical Presentation of Tracheobronchopathia Osteochondroplastica: Is Chronic Inflammation a Perpetrator?

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Tracheobronchopathia osteochondroplastica · Airway inflammation · Bronchoscopy · Nodule · Ossification · Chronic cough

Introduction
Tracheobronchopathia osteochondroplastica (TO) is characterized by development of multiple osseous and cartilaginous nodules in the submucosa of the trachea and the main bronchus \cite{1, 2}. Patients usually present with cough, recurrent respiratory tract infections, and occasionally hemoptysis \cite{3, 4}. TO is not usually suspected until fiber-optic bronchoscopy is performed; the bronchoscopy views together with histopathological examination of the nodules confirm the diagnosis.

Case Report
A 59-year-old male smoker (30 pack-years) was investigated for productive cough of 1 month. He had a past history of pulmonary tuberculosis (25 years ago; unavailable medical records) and mentioned occasional episodes of dry cough every year for the last 3 years. His family and occupational history were not significant. One month earlier, he had been diagnosed with pulmonary tuberculosis at another institution, but his sputum smear and culture...
a better respiratory status. The patient was managed with bronchomegaly (to verify that his airway changes were not drug-induced). Within 1 week of discontinuation, his symptoms eased and he had only mild COPD symptoms. A provisional diagnosis of endobronchial tuberculosis was made and a combination regime and the symptomatic recovery after discontinuation of the drugs are an interesting association with lung cancer (adenocarcinoma in particular) has been suggested [9].

CT and bronchoscopy revealed features suggestive of TO, but acute deterioration is not typical of this disease [6]. Other differential diagnoses of TO include calcified fibroma, endobronchial sarcoidosis, polychondritis, and Wegener’s granulomatosis of the proximal airways. Isolation of Klebsiella ozaenae, both in atrophic rhinitis and TO, suggests a link between these disorders [8]. Isolation of Klebsiella ozaenae, both in atrophic rhinitis and TO, suggests a link between these disorders [8]. Isolation of Klebsiella ozaenae, both in atrophic rhinitis and TO, suggests a link between these disorders [8].

Discussion

TO is limited to the large airways and does not involve the lung or other organs [1]. Changes at the mucosal surface and altered clearance of secretions result in recurrent inflammation and infection [3]. These lesions typically spread over the anterior and lateral walls of the airways (but not the posterior wall). Studies suggest that only 51% of patients with TO are accurately diagnosed during their lifetime [5]. Our patient was reevaluated after 1 month, CT and bronchoscopy revealed features suggestive of TO, but acute deterioration is not typical of this disease [6].

Classic hypotheses include ecchondrosis and exostosis arising from the cartilaginous rings, or metaplasia of the submucosal elastic and connective tissue [1–3]. An association with lung cancer (adenocarcinoma in particular) has also been suggested [7]. Cystic fibrosis coupled with bacterial infection induces metaplastic bone replacement, as well as destruction and elimination of the bronchial cartilage. Degenerative changes in the cartilage and increased perichondrial fibrosis have been demonstrated in patients with chronic obstructive pulmonary disease and bronchial asthma. We assume that chronic inflammation of the large airway, in part due to recurrent infection, may have been the cause for our patient’s initial complaint of cough. Moreover, the acute changes in the airways observed 1 month after starting the antimycobacterial combination regime and the symptomatic recovery after discontinuation of the drugs are an interesting association that remains unexplained. Whether the antimycobacterial drugs played a synergistic role by accelerating the inflammatory process is debatable. There are reports suggesting coexistence of tracheobronchial amyloidosis and TO [8].

Isolation of Klebsiella ozaenae, both in atrophic rhinitis and TO, suggests a link between these disorders [9]. Other differential diagnoses of TO include calcified lesions secondary to tuberculosis, carcinoma, papilloma, fibroma, endobronchial sarcoidosis, polychondritis, and Wegener’s granulomatosis of the proximal airways. Immunohistochemical studies of TO lesions suggest a role for bone morphogenetic protein 2 [10]. The above-mentioned features were not evident in our patient, and the short history made understanding his case rather complex. Multiple factors are probably involved in the pathogenesis of TO; the cartilage ossification seen in our pa-

![Fig. 1. a Chest CT scan showing a ring (arrow) of multiple anterolateral calcified nodules surrounding the tracheal lumen but sparing the posterior walls. b, c Bronchoscopic view of airways after 1-month therapy, showing diffuse anterolateral distribution of nodules (arrows) in the typical cobblestone appearance of TO. The posterior wall is spared. d Histopathologic view of the nodules from the patient illustrates (arrows) submucosal calcification, ossification, and cartilage formation.](https://example.com/fig1.png)
tient may possibly be the result of an intense inflammatory reaction in the bronchial mucosa. More case reports and studies on the etiology of the condition will help to clarify this issue.

There is lack of consensus among clinicians on the optimum treatment, while conservative therapy aims at maintenance of airway humidity, control of infection, and avoidance of airway irritants, treatment modalities include bronchoscopy-guided excision of the nodule, laser ablation, surgical resection, and radiotherapy [1–4].

### Conclusion

This case report showed that TO should be considered in patients with cough not explained by noninvasive testing and not responsive to empiric medications. CT results may be suggestive, but bronchoscopy examination, followed by histopathological findings is diagnostic of TO. Interventional bronchoscopy has an important role in the symptomatic treatment of TO.

### References