**Introduction**

Tracheal diverticulum is a rare entity consisting of an outpouching or sac arising from the wall of the trachea. We present here a case of TD diagnosed in a patient with blunt thoracic trauma.

**Case Report**

A 60 years old man reported a blunt thoracic trauma after falling from three meters high. He was conducted to the Emergency Department of our hospital and CT of the chest was carried out. A right scapula fracture with fragmentation and multiple ribs fracture (ninth, eleventh and twelfth right rib) were diagnosed. Moreover, an air collection of about $14 \times 8$ mm was found in the right paratracheal area, apparently in communication with the airway lumen (Figures 1, 2, 3).
A diagnostic bronchoscopy was scheduled the following day. Under conscious sedation, the bronchoscope was introduced through the nasal fossa. At the border of the upper third with the lower two thirds of the trachea, dorsally and slightly on the right of the median line, a little orifice was demonstrated (Figure 4).

The presence of this diverticulum was confirmed by the presence of air bubbles just below it (Figure 5).

A traumatic origin of this orifice was ruled out because of the integrity of the mucosa and the absence of signs of endoluminal bleeding. The diagnosis of tracheal diverticulum was clear. Aspiration of endobronchial fluids was carried out for routine microbiological and cytological analysis which later resulted within normal range.

The patient was treated conservatively because no complication developed as a consequence of the thoracic trauma. Indeed he was discharged in good health condition seven days after admission. A medical therapy based on pain killers was prescribed; a control chest X-ray and a follow-up visit in our outpatient clinic was scheduled ten days after discharge. No surgical treatment of the tracheal diverticulum was planned as the patient was asymptomatic.

**DISCUSSION**

Tracheal diverticulum (TD) consists of small air collection in the paratracheal area. It is a rare entity with an estimated incidence of 2.4%[1]. In most cases TD is located at the right posterolateral region of the trachea (97.1%) and rarely at the contralateral side (2.9%)[1]. It is lined by the same ciliated columnar epithelium of the trachea and...
its lumen is in communication, though often difficult to demonstrate, with that of the airways. These features enable to differentiate TD from other paratracheal air cysts (PTACs) such as tracheoceles, lymphoepithelial cysts and bronchogenic cysts. These entities which share the same radiological aspect of “air collection” have an altogether incidence of 0.75-8.1%[2-4].

**Tracheal diverticula can be congenital or acquired**

Congenital tracheal diverticula are thought to be the consequence of a defect in endodermal differentiation of the membranous posterior tracheal wall [5]. As a consequence, all the layers of the airways (respiratory epithelium, smooth muscle and cartilage) are involved and can be demonstrated within the wall of the diverticulum at pathological examination (true diverticulum). Congenital diverticula are reported to be more common in males, smaller than the acquired counterpart and generally located 4-5 cm below the vocal cords or just above the carina. Connection with the trachea is so narrow that it can be hardly demonstrated by bronchoscopy.

Acquired tracheal diverticula occur as a complication of surgical procedure or as a result of tracheomalacia; whatever the cause, an area of weakness in the tracheal wall develops. A long standing increased pressure inside the airways, such as in obstructive lung disease, emphysema and chronic cough, pushes the mucosa out of the defect. As a result, the acquired diverticulum is made of respiratory mucosal lining only, is wide mouthed and larger in size (pseudodiverticulum). Moreover it is usually located at the level of the thoracic inlet between the extrathoracic and intrathoracic area. Acquired diverticulum can be single or multiple and, when multiple, a tracheobronchomegaly or Mounier-Kuhn disease can be claimed[6-7].

Symptoms related to TD are chronic cough, dyspnea, stridor or recurrent tracheobronchitis[8]. Dysphagia, odynophagia, neck pain, hoarseness, hemoptysis, choking, recurrent episodes of hiccups and/ or burping have also been described[9]. Compression of the laryngeal nerves can lead to dysphonia. Several cases of paratracheal abscesses have been described in the literature following infection of a TD.

Nevertheless the great majority of tracheal diverticula, both congenital and acquired, are asymptomatic. Diagnosis is incidental following a chest CT. Multidetector computed tomography is considered the best method to demonstrate a TD. It is able to show the exact location, size and contour and wall thickness. If the CT scans are thinner than 1 mm, a communication between TD and trachea can be seen. Presence of such communication could not be equally obvious at bronchoscopy because TD can have a very narrow opening or it can be joined to the trachea by a fibrous tract only. Though communication of TD with the tracheal lumen enables differential diagnosis from other paratracheal air cysts which don’t have such connection.

The rarity of TD is that sheer and simple diagnosis of this entity deserves some mention. Yet the case we observed had some further peculiarities that makes it worth of consideration. First, diagnosis was based not only on chest CT which showed a paratracheal air collection with a likely communication with the airway lumen but, differently from usual, by bronchoscopy also. Indeed bronchoscopy was able to directly visualize the orifice of the TD. This enabled differential diagnosis from other PTACs.

Secondly, the context of blunt thoracic trauma of our case made diagnosis of TD challenging. Indeed the detection of a para tracheal air collection on CT images could have been misinterpreted as a traumatic rupture of the trachea leading to unjustified surgical treatment. Clinical judgment supported by the absence of other signs and symptoms of tracheobronchial injury, led to a more cautious treatment plan. The following bronchoscopy correctly diagnosed a TD.

**CONCLUSIONS**

Tracheal diverticulum is a rare entity to be distinguished by other paratracheal air cysts. Diagnosis is often difficult because of lack of specific symptomatology, and low sensibility of even invasive procedures such as bronchoscopy. The case of TD reported here has the peculiarity of being diagnosed by a rare combination of CT images and bronchoscopic direct visualization. Clinical judgment, guided by the above mentioned diagnostic tools, avoided a-first-instance, yet wrong, diagnosis of traumatic laceration of trachea and enabled the correct treatment of the patient.

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