Bronchial Mucoepidermoid Carcinoma in a 14-year-old Patient

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Abstract
A bronchial mucoepidermoid carcinoma is a rare malignant tumour. It usually appears in young subjects and most often involves the salivary glands. We report the case of a 14-year-old patient who presented with a pulmonary mucoepidermoid carcinoma that was discovered during the evaluation of a case presenting with recurrent pneumopathies. Recurring pneumopathies must lead to suspicion of various aetiologies and therefore require thorough evaluation so that rare tumours are not ignored, not missed, particularly in children.

Keywords
Mucoepidermoid carcinoma, Recurrent pneumopathies, Segmentectomy

Introduction
Bronchopulmonary tumours are rare among children. Bronchial mucoepidermoid carcinoma is a malignant tumour that accounts for 0.1% to 0.2% of pulmonary tumours [1]. They are usually observed in young subjects who are less than 30-years-old. These lesions are often found in salivary glands. They are composed of mucus-secreting cells, squamous cells and intermediate cells. To date, surgical resection remains as the first treatment. Its prognosis is good for low-grade lesions; high-grade lesions can result in local recurrences and metastases. We present the case of a 14-year-old girl patient who was treated for a pulmonary mucoepidermoid carcinoma that was discovered in a patient presenting with recurrent pulmonary infections.

Observation
A 14-year-old girl child presented with a history of recurrent pulmonary infections that was always involving the upper lobe of left side lung (Figure 1). In November 2010, she presented with the first acute pulmonary infection. The resolution was good after receiving empirical antibiotic therapy. During February to June 2011, she presented with recurrent pulmonary infections. The chest X-ray showed a left pulmonary opacity (Figure 1). The diagnosis of a pulmonary mucoepidermoid carcinoma was made after the histologic examination.
3 more attacks in the same pulmonary territory that resolved with antibiotic treatment. Chest X-rays taken after recovery were considered normal every time.

In April 2012, a chest CT scan was first performed to evaluate the acute pain in the left side chest in the patient (Figure 2). It demonstrated a 7 millimetre diameter tumour that was responsible for atelectasis of the anteroinferior segment of upper left lobe. Bronchial endoscopy confirmed the obstruction of the proximal lingual (Figure 3). A carcinoid tumour was first suspected from the endoscopic aspect. Several endoscopic biopsies of the obstructive lesion did not allow a pathological diagnosis. Surgical treatment, performed through transaxillary thoracotomy, consisted of a lingulectomy with lymph node dissection in areas 4, 5, 7, 10 and 11 L. Four lymph nodes were dissected. Pathological examination of the surgical resection demonstrated a 2 centimetre diameter low-grade malignant mucoepidermoid carcinoma. The diagnosis was confirmed by routine

![Figure 2: CT scan demonstrating atelectasis of the lingual (A) and an intrabronchic obstructive lesion (B).](image)

![Figure 3: Endobronchial lesion of the proximal lingula, a grey rosy nodular polyp on bronchial endoscopy (Arrow).](image)

![Figure 4: Low grade mucoepidermoid carcinoma A) The tumour surrounds the bronchial cartilage and is mainly of granular architecture, (25×); B) Higher magnification showing mucus cells (arrowhead) that are intermingled with intermediate cells (arrow 2) and epidermoid cells (arrow 1) (200×). (hematoxylin-phloxine-saffron stain).](image)
Discussion

Smetana, et al. first described the mucoepidermoid carcinoma in 1952 [2]. It occurs in all ages ranging from 4 to 80-years-old with prevalence in young adults between 20 and 30 years of age. It accounts for only 0.1% to 0.2% of primary bronchopulmonary cancers [3]. No supporting factor has found to explain the occurrence of this tumor. This tumor is classified as low-grade or high-grade according to histological criteria such as cellular atypia, mitotic activity, tumoural necroses, or local extension of the disease. This tumor mainly occurs in the lobar or segmental bronchi but rarely occurs in the most distal bronchi. In the same ages, the most common tumour types were carcinoid, inflammatory myofibroblastic tumour and pleura pulmonary blastoma. It is possible to find all types of bronchogenic carcinoma that are described in adults in the pediatric age group also. Carcinoid tumours accounted for 11% to 13% of all paediatric pulmonary tumours [4].

Among the two histological subtypes of mucoepidermoid carcinomas, low-grade lesions account for approximately 95% as compared to only 5% of high-grade lesions. Histologically, low-grade lesions primarily consist of a glandular structure, clear limits and a small few mitoses. There is no necrosis in low-grade lesions in contrast to high-grade lesions, the later present with a majority of squamous component [4]. Mitoses are numerous in high-grade lesions. For high-grade lesions, diagnosis is often difficult because they have very similar histological characteristics of adenocarcinomas and squamous carcinomas (Figure 4).

Low-grade lesions have a good prognosis with rare local recurrence, which is opposite in case of high-grade lesions that have a prognosis that seems poorer according to the small number of reported cases.

The clinical presentation of these tumours is not specific and initially consists of an irritation of one of the air routes, gradually progressing to complete obstruction. Thus, cough, haemoptysis, dyspnoea or recurrent pulmonary infections can be observed [3,5]. In this context, several diagnoses were possible in children like the ingestion of any object. Chest X-ray is usually not conclusive, but in the absence of a clear explanation repeated pneumopathy in the same territory of the lung should allow suspicion of the existence of a tumour. Bronchial endoscopy can then confirm the existence of a grey rosy nodular Polyp Endobronchial lesion; pathological examination of biopsies should allow the diagnosis.

In parallel, an injected CT scan typically shows oval or round well-defined lesions. Hyper vascularisation of the tumour can lead to misdiagnosis with carcinoid tumours [6,7]. Calcifications can also be noted.

The treatment of these tumours consists of complete surgical resection. Association with radiotherapy for lesions with a high rank has been proposed [8]. In our case, we performed a minimally invasive surgery, a transaxillary muscle-sparing thoracotomy. The aim was to preserve muscle and aesthetics for the young child.

Conclusion

Mucoepidermoid tumours with pulmonary locations are rare tumours that are difficult to diagnose because of variable clinical presentations. It is necessary to repeat routine examinations and bronchial endoscopy in order to understand the etiology of repeated respiratory episodes, particularly in young patients.

Regardless of the minimal frequency of these lesions, it is necessary to remember that mucoepidermoid tumours occur in young patients without particular risk factors and that the risk of metastases exists in the absence of treatment. Continuation of clinical and radiological monitoring seems necessary in order to detect local recurrences.

References

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