Prenatal Diagnosis of Bronchogenic Cyst at the Lingual Base: Case Report

Asma K1*, Amenallah G1, Dorra K2, Aida M3 And Wiem Dk4

1Radiology resident, Radiology consultant, Department of pediatric medical imaging, Bechir Hamza Hospital, Beb Saadoun, Tunisia
2Internal medicine resident, Department of pediatric medical imaging, Bechir Hamza Hospital, Beb Saadoun, Tunisia
3Professor in embryology, Department of embryo-foetopathology, Maternity Center, Tunis, Tunisia
4Professor in radiology, Chief of pediatric medical imaging, Bechir Hamza Hospital, Beb Saadoun, Tunisia

Received: 04 Mar 2024
Accepted: 29 Apr 2024
Published: 04 May 2024

Copyright: ©2024 Asma K. This is an open access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and build upon your work non-commercially.


1. Clinical Case Report

A 39-year-old female woman with no previous history presented at 19 weeks of pregnancy for a routine fetal ultrasound. It was her third pregnancy with no notable event so far. No particular familial or personal medical history was revealed. The biometry corresponded to a term of 19 weeks and 4 days gestational age of a male fetus with no major malformation except a cervical cystic mass that was discovered near the floor of the mouth.

The cyst was located right on the midline, had a well-defined and thin peripheral wall with regular contours and pure anechoic material (Figure 1). It contained a single fine septum with no intrinsic signal on colour Doppler and measured 18 x 10 mm. All these findings supported a cervical lymphatic malformation, and a medical interruption of the pregnancy was proposed to the couple after interdisciplinary decision. A fetal pathological examination was led with parental consent. It confirmed the diagnosis of a cystic mass at the base of the tongue (Figures 2,3) containing structural elements of the airways including a ciliated epithelium, and thus concluding to an extra-thoracic bronchogenic cyst. The neck is a rare localisation of bronchogenic cysts, whereas the mediastinum and the thorax are the most frequent localisations [1,2]. We have not found many publications of similar localisations at the base of tongue through our brief literature review. Bronchogenic cysts originate from abnormal development of the primitive oesophagus and the tracheobronchial tree [3]. They are generally unilocular, filled with mucus and their wall is multilayered containing components of the airways: cartilage, smooth muscle, mucous glands and ciliated respiratory epithelium [3]. Prenatal ultrasound can easily detect a unilocular cyst located in the mediastinum, the thorax or the neck area, filled with anechoic fluid and surrounded by a well-defined thin wall [3]. However, research revealed the possibility of bronchogenic cysts presenting as hyperechoic lesions [4]. The principal threats to the fetus that can occur in cases of large cysts include heart compression and lung compression causing secondary parenchymal hypoplasia with the progressive bronchial obstruction, as reported by D. Levine et al. [5]. The evaluation of these risks has motivated the interdisciplinary decision of pregnancy interruption in our case. In the majority of cases, bronchogenic cysts are asymptomatic in the post-natal period. When they are symptomatic, they manifest through repeated bronchopulmonary infections and airway obstruction [6].
Figure 1: Midline sagittal view of fetal ultrasound showing a unilocular anechoic cyst surrounded by a thin regular wall at the base of the tongue.

Figure 2: Macroscopic examination of the fetus showing the cyst at the base of the tongue.

Figure 3: Macroscopic examination of the section of the cyst showing cartilage inclusions and mucosal compartments.

References


