Tympanic membrane cholesteatoma: A case report in an adult and a child

Estefanía Miranda; Victoria García-Peces; Mar Martínez-Ruiz-Coello; Cristina García-García; Agustina Arbía; Guillermo Plaza

Abstract

Introduction and objective: Tympanic Membrane Cholesteatoma (TMC) is a rare pathology of unknown etiology which occurs more frequently in pediatric population, being the presentation in adults exceptional. Resection of the lesion is usually the surgical treatment through a transcanal approach.

Method: We report two patients, one adult and one pediatric, presenting TMC with no previous otologic history. We explain diagnosis and management, detailing complementary tests and final treatment.

Discussion: TMC is a rare pathology. Despite this, it should be considered when a whitish nodular lesion appears in the tympanic membrane in patients with no previous otologic history. Currently, there are few cases published in the literature.

Conclusions: TMC is a rare and infrequent entity. Due to the limited literature on the subject, the criteria to classify a TMC as congenital or acquired and its possible pathogenic mechanisms are still under discussion. The lesion is usually limited to the tympanic membrane and the treatment of choice is surgery.

Keywords: Tympanic membrane cholesteatoma; Intratympanic cholesteatoma; Tympanic membrane; Congenital cholesteatoma.

Abbreviations: TMC: Tympanic Membrane Cholesteatoma.

Introduction

Congenital cholesteatomas of the middle ear are infrequent, but even less frequent in the tympanic membrane [1,2]. Tympanic Membrane Cholesteatoma (TMC) is a rare pathology, as there are few cases reported in the literature. It is of unknown etiology, which occurs more frequently in pediatric population. However, cases have also been reported in adults. The typical presentation is characterized by presentation in patients with no previous otologic history, that is, chronic otitis media, trauma or otologic surgery [1]. In most cases, the lesion is limited to the tympanic membrane with the middle ear, ossicular chain, attic and mastoid cells being normal. In the few cases reported, resection of the lesion is usually the surgical treatment, being the transcanal approach the treatment of choice. We present the case of two patients: an adult and a child with a diagnosis of TMC. The aim of this article is to expose an uncommon pathology such as TMC.
Case presentations

Case 1

We present the case of a 66-year-old male patient who came to the otolaryngology office in our hospital for otalgia of the right ear with no other symptoms. He had no relevant pathologic history, denied any history of otitis, trauma or previous otologic surgeries. Physical examination by otomicroscope revealed a nodular lesion in the lower quadrants of the tympanic membrane (Figure 1). Audiometry and tympanogram were normal. A CT scan was performed, showing a lesion with soft tissue attenuation at the tympanic membrane level, with no alterations at the middle or inner ear level (Figure 2).

Surgical treatment was proposed with resection of the lesion through a transcanal approach. During the procedure, the edges of the lesion were detached, leaving the tympanic membrane intact (Video 1). The diagnosis of TMC was confirmed by histopathology. The patient was evaluated 6 months after with a normal tympanic membrane (Figure 3).

Case 2

A 1-year-old male patient was evaluated in the otolaryngology office referred by his primary care physician for suspicion of a foreign body in the left ear. The mother perceived adequate hearing. He had no previous otologic history. Physical examination revealed a nodular lesion in the inferior quadrants of the tympanic membrane (Figure 4). A CT scan was performed, showing otomastoid occupation without destruction of local structures (Figure 5). Therefore, a transcanal resection of the TMC was performed without violating the integrity of the tympanic membrane. Myringotomy was performed suctioning mucus. The tympanomeatal flap was lifted and the middle ear was explored without evidence of cholesteatoma. A transtympanic tube was placed. The diagnosis of TMC was confirmed by histopathology. After four years of follow-up tympanic membrane remained intact, with no recurrence of TMC (Figure 6).
tients with this pathology have no initial complaint of hearing loss, the lesion may go unnoticed for many years. Patients with TMC are more likely to present TMC. In our case, the two patients reported in the literature. 

The first description of a TMC was in 1863 by Hinton, who described it as a “sebaceous tumor” located on the external surface of the tympanic membrane [3]. In 1936, Teed found a total of five congenital cholesteatomas in the literature that were compatible with a TMC [4]. In 1997, Weber and Adkins described three other cases of TMC in children with no previous otologic history [5]. More recently, in 2016 and 2019, Predruzzi et al. and Cheon et al. respectively described the cases of patients with this pathologic, each of these cases had a small mass on the external surface of the TM [6,7].

The etiopathogenesis of TMC is still unknown and seems to be unrelated to congenital cholesteatoma of the middle ear. In cases of otitis previa, the origin of TMC can be explained by Sade et al. who observed that inflammation can lead to metaplasia of mesenchymal tissue within the tympanic membrane, which result in conversion to squamous epithelium [8]. On the other hand, Ruedi proposed that inflammation may develop proliferation of the basal cell layer, from the squamous epithelium of tympanic membrane into cones protruding in it to form a cholesteatoma in response to an inflammatory process [9]. Our patients differ from this theory because of the absence of prior middle ear pathology.

When cholesteatoma develops in subjects with no history of previous inflammatory processes of the external and middle ear, as in the cases we present, an embryologic origin is suspected. This could be explained by the persistence of an epidermoid formation, derived from the first branchial groove that normally regresses after the 33rd week of gestation [4,10,11]; a theory that would apply to our patients, since they do not have a previous otologic history.

The biggest series reported of TMC was done by Reddy et al. [12], including 15 cases ranging in age from 6 months to 34 years, most of them younger than 4 years. There were no sex differences and no quadrant of the tympanic membrane was expected. This could be explained by the persistence of an epidermoid formation, derived from the first branchial groove that may result in conversion to squamous epithelium [8].

Recurrence of TMC appears to be lower than cholesteatoma of the middle ear [1]. However, postoperative follow-up of these patients is necessary.

Conclusion

Tympagic membrane cholesteatoma is a rare and infrequent entity that should be taken into consideration both in pediatric age and in adults, always when a nodular lesion external to the tympanic membrane is observed. Due to the limited literature on the subject, the criteria to classify a TMC as congenital or acquired and its possible pathogenic mechanisms are currently under discussion. The lesion is usually limited to the tympanic membrane and the treatment of choice is surgery.

Declarations

The author(s) declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

References

6. Pedruzzi B, Mion M, Comacchio F. Congenital Intr tympanic


