Treatment of seborrheic keratosis in bilateral external auditory canal using fiber CO₂ laser

Fiber CO₂ lazer kullanarak iki taraflı dış kulak kanalında seboreik keratoz tedavisi

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ABSTRACT

In this article, we report a 33-year-old female patient with lesions of bilateral external auditory canals who was treated using fiber CO₂ laser without canaloplasty. Histopathological examination confirmed the diagnosis of irritated type seborrheic keratosis, which is rare in external auditory canal. Twelve months after the treatment, the patient showed no signs of recurrence.

Keywords: External auditory canal; laser; seborrheic keratosis.

ÖZ

Bu makalede, iki taraflı dış kulak kanallarında lezyonlar olan ve kanaloplasti olmadan fiber CO₂ lazer kullanılarak tedavi edilen 33 yaşındaki bir kadın hasta sunuldu. Dış kulak kanalında nadir görülen irrite tip seboreik keratoz tanısı histopatolojik incelemede doğrulandı. Tedaviden 12 ay sonra hastada herhangi bir nüks belirtisi görülmedi.

Anahtar Sözcükler: Dış kulak kanalı; lazer; seboreik keratoz.

Seborrheic keratosis (SK) is one of the most common, benign skin lesions in older patients that can involve any part of the body except the palms and soles. The most frequently affected sites are the chest, back, head (near the temples), and neck. The presence of SK in the external auditory canal (EAC) is rarely seen. To date, only a few reports of SK in the EAC have been published. Generally, in all these cases, SK occurred unilaterally and was treated by complete surgical excision with or without skin graft. Here, we describe bilateral EAC SK that were removed using fiber CO₂ laser.

CASE REPORT

A 33-year-old woman presented with one-month history of lesions in the right and left EAC with bilateral conductive hearing loss. On otoendoscopy, skin-colored, verrucous lesions were found originating from all walls of the EACs and totally occluding them bilaterally (Figure 1, for left EAC). The review of history revealed that...
two years before, she presented with lesions in both EACs which were histopathologically diagnosed as SK by incisional biopsy and subsequently removed by electrocoagulation. Here, we chose flexible fiber delivered CO₂ laser (Omniguide Inc., Cambridge, MA, USA) to excise the recurrent lesions without performing incisional biopsy or radiological examination. After excision, both EACs seemed to be smooth and both tympanic membranes were intact. (Figure 2, for left EAC) Histopathological examination confirmed the diagnosis of irritated type SK (Figure 3a, b). There was no recurrence of disease on 12-month follow-up. Written informed consent was obtained from the patient.

DISCUSSION

Seborrheic keratosis is a common, benign epithelial skin tumor that is also known as verruca senilis, senile wart, seborrhic wart, verruca seborrhoica, basal cell acanthoma, benign acanthokeratoma, and basal cell papilloma. The prevalence of this benign tumor increases particularly in older patients without predilection for either sex. Clinically, lesions appear as sharply demarcated and round or oval shaped, usually elevated and stuck on the skin with a verrucous, uneven surface. Flat forms may exist and often have a smooth surface. The sizes and colors of lesions are variable. They may be skin-colored, yellowish, gray-brown, or black. Although SK is usually asymptomatic, they may cause itching, pain, bleeding and redness. They can be generally observed on the chest, back, head and neck. The EAC is a rare affected site of disease that can cause conductive hearing loss. In our patient, the lesions were bilateral as distinct from other cases of EAC SK in the literature and there was no symptom except bilateral hearing loss.
The etiology and pathogenesis of SK are not well understood. However, it was believed that SK is more prevalent among Caucasians. In recent years, human papillomavirus infections with risk factors including older age and exposure to ultraviolet light were mentioned in pathogenesis. Genetic predisposition as an autosomal dominant pattern has also been suggested although there is a lack of validated data. In addition, FGFR3 gene and PIK3CA oncogene mutations were described in the molecular pathogenesis of SK.

Histopathologically, SKs have characterized findings including acanthosis, papillomatosis, hyperkeratosis, horn cysts and horn pseudocysts. There are seven different histological subtypes: acanthotic, hyperkeratotic (also verrucous), adenoid, clonal bowenoid, irritated, and melanocanthoma. The acanthotic subtype is believed to be the most frequently seen in EAC. Kim et al. reported seven patients with SK in the ear and six of them had the acanthotic subtype. In our case, the irritated subtype of SK was confirmed on histopathological examination, which is rare in the EAC. To best of our knowledge, there is only one report of irritated SK in EAC in the literature. The irritated subtype is characterized by onion-skin like aggregations of eosinophilic squamous epithelial cells (squamous eddies), and inflammatory cell infiltration with partly lichenoid appearances in the dermis. Acantholysis, dyskeratosis and spongiosis may also be present. In our patient, hyperkeratosis, horn cysts, small squamous eddies, proliferation of basaloid cells, and moderate inflammatory infiltrate were observed in the epidermis. In the differential diagnosis of SK, benign lesions such as senile keratosis, papillomas, keratoacanthomas, fibroma and malignant lesions such as malignant melanoma, basal or squamous cell carcinoma should be considered.

Removal of the SK may be required to exclude the presence of malignancy if clinical appearance is equivocal, or to end the symptoms, inflammation, and cosmetic problems. Treatment options include curettage, complete excision and ablative procedures (laser therapy and cryotherapy). Treatment using topical and systemic drug therapies have also been reported, but have not yet been proven. Erbium YAG or CO₂ laser may be used for removal of SK. In addition, the use of a 532 nm diode laser in dermatosis papulosa nigra, a clinical variant of SK, has been reported. On literature review, there was only one case of SK in the EAC treated by CO₂ laser, which was presented as a poster and here, we describe the second case. Although ablative procedures are known to be inconvenient for obtaining histologic analysis and incisional biopsy can be necessary before surgery, we managed to obtain adequate specimens for diagnosis without performing biopsy before laser treatment. Our knowledge about the previous diagnosis of the lesions and the clinical presentation of the patient consistent with SK led us to choose laser surgery. An advantage of this procedure was having no need for canaloplasty after excision. No recurrence was observed in our patient 12-months post-ablation. However, this tumor always requires clinicians to be alert about potential local recurrence.

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